
An animal’s long-term survival requires that its organs adapt to unpredictable environmental changes. Yet how do mature organs sense external change and deploy an appropriate, reversible response? We seek the cellular and molecular basis of adult organ plasticity—i.e., the remarkable facility with which organs adaptively switch between states of renewal, remodeling, and repair. Our focus is the adult Drosophila gut, whose deep playbook of adaptive responses is executed by a multi-fated collective of stem and terminal daughter cells.

As the cell is the fundamental unit of an organ, we posit that the cell’s lifecycle—from birth to differentiation, maturity, and finally death—is the fundamental unit of an organ state. Using new techniques we devised to image functioning guts in live animals, we are resolving these surprisingly diverse lifecycles in real time. We have demonstrated the basis of conservative lifecycle kinetics that maintain the status quo, such as when the gut sustains steady-state turnover by ‘closing the loop’ between apoptosis and death through the intimate relay of a transient, spatially delimited EGF. When ingested toxins raze large swaths of differentiated cells, the damaged cells unleash cytokines that restore the status quo by concerted acceleration of stem cell divisions and daughter cell differentiation. We also have discovered non-conservative lifecycles that adapt the status quo to dietary change. For instance, when food is withdrawn, the gut quickly shrinks as mature cells abort their lifecycles and forego apoptosis to eject live from the tissue. Altogether, our findings illustrate how organ plasticity flows from extrinsically triggered tuning of single-cell lifecycles.

10 Regulating host phospholipid metabolism to fight infection Michelle Bland University of Virginia

Infection and subsequent immune signaling drive synthesis and secretion of immune effectors and mobilization of immune cells, processes that are supported and directed by host metabolism. In the Drosophila larval fat body, the metabolic challenge of fighting infection must be balanced with the energetically-demanding processes of growth and nutrient storage that this organ coordinates. We find that Toll signaling in larval fat body disrupts whole-animal growth and directs a lipid metabolic switch from fat storage to phospholipid synthesis. Although reduced triglyceride storage in larvae with active Toll signaling impairs the response to stress later in life, elevated phospholipid synthesis in fat body supports humoral immune function. Toll signaling in larval fat body directs increased production of membrane phospholipids that coincides with expansion of the endoplasmic reticulum. Phospholipid synthesis in larval fat body supports antimicrobial peptide secretion and promotes bacterial clearance. Our results define a metabolic role for the pleiotropic Toll signaling pathway, and they demonstrate that the demands of fighting infection are met with trade-offs in growth and energy storage to sustain immune function.

11 The genetics basis of inviability in hybrids between Drosophila melanogaster and D. santomea Daniel Matute University of North Carolina

Evolved changes within species lead to the inevitable loss of viability in hybrids. Inviability is also a convenient phenotype to genetically map and validate functionally divergent genes and pathways differentiating closely related species. Here we identify the Drosophila melanogaster form of the highly conserved essential gap gene giant (gt) as a key genetic determinant of hybrid inviability in crosses with D. santomea. We show that the coding region of this allele in D. melanogaster/D. santomea hybrids is sufficient to cause embryonic inviability not seen in either pure species. Further genetic analysis indicates that tailless (tll), another gap gene, is also involved in the hybrid defects. giant and tll are both members of the gap gene network of transcription factors that participate in establishing anterior-posterior specification of the dipteran embryo, a highly conserved developmental process. Genes whose outputs in this process are functionally conserved nevertheless evolve over short timescales to cause inviability in hybrids.

12 Tissue Biology of Chromosomal Instability Marco Milan IRB Barcelona

Chromosomal Instability (CIN), defined as an increased rate of changes in chromosome structure and number, is a feature of most, if not all, solid tumors. While CIN promotes the gain of oncogene-carrying chromosomes and the loss of tumor-suppressor-gene-carrying chromosomes in certain cancers, its impact on the biology of the cell and on the homeostasis of the tissue, as well as its role in tumorigenesis, are far from being fully elucidated. Our lab has generated an epithelial model of CIN in Drosophila, where the generation of highly aneuploid karyotypes drives cell delamination and cell death. Upon apoptosis inhibition, aberrant karyotypes promote a cell autonomous epithelial to mesenchymal (EMT)-like cell fate transition associated with a highly invasive behavior and the entry into a senescence-like state. I will present our most recent data on the genetic identification and functional characterization of the molecular mechanisms underlying these cellular behaviors and the leading role of aneuploidy in driving these behaviors.

13 Collective Action for Institutional Transformation Shaila Kotadia Stanford University School of Medicine

Within higher education institutions and the sciences is a lack of support, care, and safety for marginalized communities due to a history of systemic oppression. In order to change this, we need to take justice-based approaches for our diversity, equity, and inclusion work, including bringing people together to reenvision culture rather than hundreds of
individual efforts. In decentralized institutions, which are common in higher education, we often do not connect across local (for example, departments, divisions, or units) efforts nor connect local to central administration where decision making occurs. By coming together and breaking down silos, we can gain power in a grassroots manner to lead to culture change across the organization. I will speak on how we can use collective action for institutional transformation using my efforts at Stanford University’s School of Medicine Human Resources Group as examples, specifically the efforts of the School of Medicine Staff JEDI (Justice, Equity, Diversity, and Inclusion) Collective. This model demonstrates one method as to how we can move from individual allyship to group co-conspiratorial work to challenge the institution for transformative change. I will also share my personal and professional journey to JEDI work, including transitioning away from Drosophila research to administrative staff JEDI positions.

14 Strategies at UCSF for addressing barriers in science that disproportionately affect people from marginalized groups. Todd Nystul UC San Francisco

A major challenge for the scientific community is that structural inequities in our society create barriers to success in science that disproportionately affect people from currently and historically marginalized groups. These inequities include institutional policies and practices that favor nonmarginalized groups, socioeconomic factors, and societal norms that perpetuate historical biases. They are pervasive, encompassing individuals’ experiences both inside and outside of the scientific community, and are often invisible to those who align with or benefit from these systems. The result is that our processes for identifying and promoting individuals based on merit fall short, and thus the scientific community cannot reach its full potential. Because these inequities have multiple causes, a sustained and multifaceted set of strategies are needed to address them. UCSF has a long history of valuing diversity and pursuing efforts to promote and advance justice, equity, and inclusion. I will discuss our recent efforts to increase the diversity of our graduate student population, and to cultivate and sustain a learning and training environment in which students from all backgrounds are able to thrive. Specifically, I will describe the establishment of our new postbaccalaureate program, PROPEL, which aims to provide trainees from historically marginalized backgrounds with the research experience and career mentorship needed to be competitive for top-tier biomedical science PhD and MD/PhD programs; the holistic review process we use for graduate program admissions; and new initiatives to strengthen mentor-mentee relationships in a diverse community. In addition, I will discuss the process we are using to engage students, faculty, and administrators in the development of these initiatives as well as the methods we are using to assess their efficacy. We know that creative solutions to address structural inequities are being developed at many different institutions around the country, and we are eager to share ideas and learn from one another.

15 NINDS Strategies for Enhancing the Diversity of Neuroscience Researchers Marguerite Matthews National Institute of Neurological Disorders and Stroke

The National Institute of Neurological Disorders and Stroke (NINDS) is committed to the development of a biomedical research workforce that is representative of the diversity in American society. NINDS seeks to promote diversity in all of its training and research programs and to increase the participation of individuals from underrepresented populations. The Office of Programs to Enhance Neuroscience Workforce Diversity coordinates NINDS’s diversity activities, spanning the training pipeline from innovative neuroscience education outreach (grades K-12) to funding opportunities and important mentoring networks across critical career transition points (from undergraduate up to junior faculty). Some of the funding mechanisms that will be discussed in this session include the following: fellowships for graduate students and postdoctoral trainees, diversity supplements that add funds to an existing grant for mentoring and training individuals from high school to faculty stage, career development awards to assist transition to independent research careers, institutional research education awards to enhance professional development and career advancement of diverse researchers. The strategy to enhance the diversity of neuroscience researchers at NINDS includes interventions that connect programs across critical transition points and activities to assist trainees with the professional development skills and scientific networks to achieve their goals.

16 A conserved anoctamin regulates olfactory neuron firing in Drosophila Pratyajit Mohapatra, Karen Menuz University of Connecticut

Insects are vectors of several diseases affecting millions of people globally. They interact with their environment primarily by recognizing and distinguishing a plethora of odors. These odors are recognized by odor receptors in the dendrites of olfactory neurons. Insect odor receptors are classified into 2 broad families: Odorant receptors (Or) and Ionotropic receptors (Ir). These 2 families are widely studied and well characterized. However, most individual receptors are poorly conserved across insect species, making them unsuitable candidates for the development of insect repellents. In a previous bioinformatics study, we identified many evolutionarily conserved genes whose expression is highly enriched in the Drosophila antenna relative to other tissues. Here, we investigate the role of one of these genes, the anoctamin AnoA. The anoctamin family contains many transmembrane proteins, some of which serve as calcium-gated chloride channels. Our work using a transgenic reporter line indicates that AnoA is broadly expressed in all classes of olfactory neurons; it is also found in hygrosensory and thermosensory neurons. Using epitope tagging, we find that
AnoA protein is specifically localized to the olfactory dendrites. Investigation of AnoA mutants with in vivo extracellular electrophysiological recordings reveals that it limits the neuronal firing response to odorants. Together, our data indicate that AnoA plays a critical role in odorant signaling in olfactory neurons, and its expression in the antenna of other insects suggests its function is likely to be evolutionarily conserved.

**17 Dissection of the BMP-activated synaptic gene network identifies dichotomous BMP-responsive elements regulating synaptic functions** Robin Vuilleumier\(^1\), Mo Miao\(^1\), Sonia Giro\(^1\), Clara-Maria Eli\(^1\), Stephane Fibotte\(^1\), Tianshun Lian\(^1\), Annie Collins\(^1\), Sophia Ly\(^1\), George Pyrowolakis\(^2\), Pejmun Haghighi\(^3\), Douglas Allan\(^1\) 1) Department of Cellular and Physiological Sciences, University of British Columbia, Vancouver, British Columbia, Canada.; 2) BIOS, Centre for Biological Signaling Studies and Institute for Biology I Faculty of Biology, Albert-Ludwigs University of Freiburg, Freiburg, Germany.; 3) Buck Institute for Research on Aging, Novato, CA 94945, USA.

Members of the Transforming growth factor beta (TGF-β)/Bone morphogenetic protein (BMP) family play critical roles in the regulation of synaptic growth, stability, neurotransmission, homeostasis and plasticity of several neuronal subtypes; yet how TGF-β/BMP signaling controls and coordinates these functions remains mostly unknown. Here, we uncovered a neuronal gene network, enriched for essential neurotransmission genes, that is directly controlled by retrograde BMP signaling in Drosophila motor neurons, by combining Drosophila genetics with differential RNA-seq, computational discovery of Smad-binding cis-regulatory motifs (BMP-activating elements, BMP-AE and BMP-silencer elements, BMP-SE), and reporter analysis of enhancer activity. Surprisingly, we discovered that the BMP-SE motif mediates BMP-dependent upregulation of neuronal gene transcription, similar to BMP-AEs. Exploring the underlying mechanism for this atypical activity, we found that absence in motor neurons of Shnurri (shn), a Smad transcriptional corepressor, switches the repressive function of the BMP-SE motif to an activator. This mechanism is underpinned by a lack of brinker (brk) repression by retrograde BMP signaling in motor neurons; thus brk-dependent de-repression is not a mechanism for BMP gene activation in these neurons. Finally, genome editing of identified BMP-SE and BMP-AE motifs in the loci of bruchpilot (brp) and a novel gene, without maturity (witty), showed that BMP signaling operates through these motifs to regulate synaptic maturation. Taken together, our data broaden the concept of BMP enhancer and silencer motifs, and demonstrate how the retrograde BMP signaling pathway regulates synaptic function.

**18 γ-secretase promotes postsynaptic development through the cleavage of a Wnt receptor** Lucas Restrepo, Alison DePew, Elizabeth Moese, Stephen Tymanskyj, Michael Parisi, Michael Aimino, Juan Duhart, Hong Fei, Timothy Mosca Department of Neuroscience, Thomas Jefferson University, Philadelphia, PA USA

Developing synapses mature through the recruitment of specific proteins that stabilize pre- and post-synaptic structure and function. Wnt ligands signaling via Frizzled (Fz) receptors play many crucial roles in neuronal and synaptic development but whether and how Wnt and Fz influence synaptic maturation is incompletely understood. We demonstrate that Fz2 receptor cleavage via the γ-secretase complex is required for postsynaptic development and maturation. In the absence of γ-secretase, Drosophila neuromuscular synapses fail to recruit postsynaptic scaffolding and cytoskeletal proteins, leading to behavioral deficits. Strikingly, introducing γ-secretase mutations linked to familial early-onset Alzheimer’s Disease into flies leads to synaptic maturation phenotypes that are identical to those seen in null alleles. This conserved role for γ-secretase in synaptic maturation and postsynaptic development highlights the importance of Fz2 cleavage, identifies its mechanism, and suggests that receptor processing by proteins linked to neurodegeneration may be a shared mechanism affecting aspects of synaptic development.

**19 Copia, a Drosophila retrotransposable element, regulates structural synaptic plasticity at the larval neuromuscular junction** Peter M’Angale, Adrienne Lemieux, Alfred Simkin, Cong Xiao, Vivian Budnik, Travis Thomson University of Massachusetts Chan Medical School, Worcester, MA

Despite advances in elucidating the genome, the function, of much of the genetic material, the so called “junk DNA”, remains largely unknown. A large portion of junk DNA is comprised of genomic parasites, known as transposable element (TEs). New evidence suggests that a domesticated TE, activity-regulated cytoskeleton-associated protein 1 (Arc1), serves as a mechanism to transport RNAs across the synapse. Here we show that a TE, Copia is enriched at the Drosophila NMJ and Copia is transported across synapses in extracellular vesicles (EVs). We observe Copia\(^\text{act}\) acting antagonistically to dArc1. Strikingly, downregulation of Copia\(^\text{act}\) at the Drosophila NMJ results in abnormal NMJ development and increased plasticity. From RNA-sequencing data coupled with digital PCR, we determined that copia is encoded by ~40 different loci each of which is found at different abundances in the Drosophila larval brain or body wall muscles. To our knowledge, this is the first report documenting a physiological role of a TE at synapses, lending further weight to recent arguments and data suggesting that TEs and potentially other types of “junk DNA” are not junk after all.

**20 Chromatin regulatory networks underlying coordinated synaptic gene expression** James Kentro, Erica Larschan, Kate O’Connor-Giles Brown University, Providence, RI

The cellular diversity of the nervous system underlies its ability to respond to stimuli and produce complex behaviors.
Neurons form intricate circuits that transmit information by passing signals through synaptic connections. We have found that hundreds of genetically encoded proteins, both common (pan-neuronal) and neuron subtype-specific, that form the structural and functional components of synapses share a temporal pattern of expression that suggests coordinated regulation. However, no mechanism coordinating the expression of these distinct classes of synaptic genes within and across neuronal subtypes is known. Using motif discovery and analysis of chromatin binding data, we have identified a candidate gene regulatory network that may coordinate expression of synaptic genes. DEAF1, linked to intellectual disability and seizures in humans, and pioneer factors GAF and CLAMP are enriched for binding at synaptic genes and are known to interact with chromatin remodelers and other co-repressor or co-activator proteins. In the Drosophila nervous system, we observe a correlation between the timing of initial synaptic gene transcription and substantial increases in chromatin accessibility at synaptic gene promoters that supports a mechanism of coordination through control of the chromatin environment. In clamp null mutants, we find an upregulation of synaptic gene transcription. RNAi-mediated knockdown of DEAF1, driven by pros-Gal4 in immature neurons, results in increased synaptic bouton number at neuromuscular junctions. These data indicate repressive roles for CLAMP and DEAF1, which have been found to physically interact. The pioneer factor GAF has been found to co-regulate many genes in opposition to CLAMP or DEAF1 in other tissues, and may perform a similar role to increase expression of synaptic genes. We predict this gene regulatory network coordinates expression of pan-neuronal and subtype-specific synaptic genes upstream of terminal selectors by controlling chromatin accessibility at synaptic genes to allow the development of shared traits across distinct neuronal fates.

21 Fatty acid flux through triacylglycerol regulates neuroblast proliferation during oxidative stress Eva Islimye, Victor Girard, Andrew Bailey, Alex P. Gould The Francis Crick Institute, London, UK

During development, multipotent neural stem cells proliferate within a specialised local microenvironment called the niche. Work in Drosophila and mammals has shown that lipid metabolism is important for regulating the divisions of neural stem cells. During Drosophila development, the proliferation of stem/progenitor cells is more protected from hypoxia and oxidative stress in the central nervous system (CNS) than in other tissues. We previously found that the biosynthesis of lipid droplets (LDs) in glia of the neural stem cell niche functions to protect against lipid peroxidation — not only in the glia themselves but also in neighbouring neural stem cells (neuroblasts). Using labelled fatty acids and cell type-specific genetic manipulations, we now investigate the origin of the lipid cargo in the triacylglycerol (TAG) core of glial LDs. We present evidence that oxidative stress triggers neurons to release fatty acids via a process that requires functional mitochondria. Extracellular fatty acids are then taken up by glia and directed towards the TAG core of LDs. Diacylglycerol acyltransferase 1 (Dgat1/Mdy) and adipose triglyceride lipase (Atgl/Bmm) manipulations demonstrate that functional mitochondria. Extracellular fatty acids are then taken up by glia and directed towards the TAG core of LDs. Diacylglycerol acyltransferase 1 (Dgat1/Mdy) and adipose triglyceride lipase (Atgl/Bmm) manipulations demonstrate that fatty acid flux through the TAG compartment of glia sustains neuroblast proliferation during oxidative stress. In contrast, fatty acid flux through the TAG compartment of neuroblasts inhibits proliferation during oxidative stress. The contrasting functions of TAG metabolism in glia and neuroblasts are consistent with a model whereby oxidative stress triggers neurons to release potentially toxic fatty acids that are then preferentially captured and processed in niche glia in order to safeguard neuroblast proliferation. Together, our findings reveal how lipid metabolic crosstalk between the three major cell types of the developing CNS functions to spare neural proliferation during oxidative stress.

22 Neuronal activity induces Glucosylceramide that is extruded via exosomes upon glial BMP signals for lysosomal degradation in glia Liping Wang1,4, Guang Lin2,4, Zhongyuan Zuo2,4, Hugo Bellen1,2,3,4 1) Program in Developmental Biology, Baylor College of Medicine, Houston, TX 77030, USA.; 2) Department of Molecular and Human Genetics, Baylor College of Medicine, Houston, TX 77030, USA.; 3) Department of Neuroscience, Baylor College of Medicine, Houston, TX 77030, USA.; 4) Jan and Dan Duncan Neurological Research Institute, Texas Children’s Hospital, Houston, TX 77030, USA.

Autosomal recessive variants in GBA1 cause Gaucher disease (GD), the most prevalent form of lysosome storage disease. GBA1 encodes a lysosomal enzyme that hydrolyses glucosylceramide (GlcCer) into glucose and ceramide. Loss of GBA1 leads to increased levels of GlcCer which causes lysosomal dysfunction. We generated a gba1b null allele (gba1bΔ22-Gal4) using CRISPR technology. We show that Gba1b is specifically expressed in glia and not in neurons at all stages of development as well as in adults. Loss of Gba1b causes a significant reduction in lifespan and a progressive activity-dependent loss of synaptic activity of neurons, which is rescued by glial specific expression of human GBA1. Moreover, glial but not neuronal specific knockdown of Gba1b recapitulates the defects found in Gba1b mutants, indicating that Gba1b is necessary and sufficient in glia to support neuronal function. We show that GlcCer is synthesized upon neuronal activity and that it is transported from neurons to glia through exosomes. In the absence of Gba1b, the glial cells accumulate GlcCer which in turn causes vacuolization and the demise of glia and is followed by neuronal death. The transfer of GlcCer from neurons to glia is induced by a factor secreted by glia. We identified this factor as a BMP ligand, TGF-β, for vertebrate cells. Finally, we show that the White protein, an ABCG transporter, plays an important role in pigment glia and that it modulates GlcCer trafficking to the glial lysosome for degradation by Gba1b through the endolysosomal pathway. Based on our observations the White protein is involved in the degradation of GlcCer in parallel with Gba1b.
23 Sequential addition of neuronal temporal cohorts generates a stimulus on-set detection circuit Ellie Heckscher1,2,3, Yi-wen Wang1,2,3, Chris Wreden1,2,3, Maayan Levy1,2,3,4, Zarion Marshall1 1) University of Chicago, Chicago, IL, 60637; 2) Institute of Neuroscience; 3) Department of Molecular Genetics and Cell Biology; 4) Department of Neurobiology

Neuronal stem cell lineages are fundamental developmental units of complex brains and neuronal circuits are the fundamental functional units. Understanding how circuits self-assemble starting from neuronal stem cells stem cells is still a largely un-answered developmental question with implications for neuroscience, evolution, and medicine. We use the Drosophila larval nerve cord as a model to study motor circuit assembly. Motor circuits are responsible for processing somatosensory stimuli and generating movement. In prior work, we found that neurons in a single lineage can be organized into groups called “temporal cohorts”. A temporal cohort is a set of neurons that are born during a small time window from a single stem cell and that have shared circuit level function (Wreden et al., 2017, Meng, et al., 2019, Meng, et al., 2020). In this study, we ask how do neurons from different lineages wire with each other?

As a model, in segment A1 of the Drosophila larval nerve cord, we studied the NB3-3 lineage and the Even-skipped Lateral (EL) interneurons it produces. First, we used Twin-spot MARCM to determine the birth order and morphology of each EL. Then, we mined the Drosophila larval connectome to identify all major direct synaptic inputs onto ELS. Using graph theory, we found that early-born and late-born ELS get significantly different inputs. This supports the idea that early-born and late-born ELS are two temporal cohorts, as was proposed by Wreden et al., 2017. We next characterized the development and function of the circuit containing early-born ELS. early-born ELS get a majority of their input from just four classes of neurons: Chordotonal sensory neurons, Basin interneurons, Ladder interneurons, and EL interneurons segments other than A1. Anatomically, early-born ELS are embedded in a feed-forward motif, getting and functionally they act as on-set detectors for vibrational stimuli. Finally, we determined the developmental origins (e.g., stem cell parent and birth timing) of Ladders and Basins. Basins are a mid-late temporal cohort from the NB3-5 neuroblast, and Ladders are the only temporal cohort from the MNB neuroblast. Basins and Ladders synapse onto early-born ELS, and are born after early-born ELS. Together these data show: 1. Additional lineages are organized into temporal cohorts. 2. There is selective wiring among temporal cohorts. 3. Post-synaptic interneurons are born after their pre-synaptic interneuron partners. These data support a model in which a somatosensory stimulus on-set circuit develops by sequentially assembling a small subset of genetically pre-specified of temporal cohort units.

24 Partial overlap between inversions and genomic islands of divergence during early stages of ecological speciation in Drosophila yakuba Erina A. Ferreira1,2, Cathy C. Moore1, David Ogereau1, Arnaud Suwalski1, Héloïse Bastide1, Jean R. David1, Rebekah L. Rogers1, Amir Yassin1,2 1) Laboratoire Évolution, Génomes, Comportement et Écologie, CNRS, IRD, Université Paris-Saclay, Gif-sur-Yvette, France; 2) Institut Systématique Evolution Biodiversité (ISYEB) CNRS, MNHN, Sorbonne Université, EPHE, Paris, France; 3) Department of Bioinformatics and Genomics, University of North Carolina, Charlotte, NC

During the early stages of speciation, genetic differences tend to accumulate more rapidly at certain regions of the genome leading to the formation of genomic islands of divergence (GIDs). However, this pattern that has been observed in a wide range of organisms may either be due to selection on beneficial alleles contained within the GIDs and/or difference in the rate of recombination due to structural variation (e.g., inversions, heterochromatic content, etc.). Here, we investigate the possible causes of GIDs in Drosophila yakuba mayottensis, a subspecies of D. yakuba specializing on toxic noni (Morinda citrifolia) fruits on the island of Mayotte. We have previously described this specialization based on a population from a single locality collected in 2013, and identified multiple GIDs through genome comparison with mainland generalist D. y. yakuba populations. In 2017, we recollected flies from Mayotte and found D. y. mayottensis to be present in three localities, always on noni fruits, reconfirming its strong association with the toxic plant. We sequenced the genomes of the three populations in pools as well as from individual isofemale lines. Comparing these sequences to mainland genomes, we identified 7 GIDs. We assembled a new genome for D. y. mayottensis using a combination of illumina short-read and Oxford Nanopore long-read sequencing methods. We identified 10 genomic regions with major chromosomal rearrangements. Three of those inversions (on the X, 2L and 2R chromosomal arms) overlapped with the GIDs. Experimental evolution using four replicate populations issued from crosses between D. y. mayottensis and D. y. yakuba showed that the three inversions were likely involved in adaptation on noni, suggesting a partial role of the genomic architecture in host shift. Interestingly, no chromosomal inversions have been detected in Drosophila sechellia, the only other Drosophila species known to be specialist on noni in the Seychelles archipelago. Despite some similarities at the genic level that we have previously described between the two noni-specialist species, our results indicate that distinct genomic architectures underlie convergent response to common selective pressures in independent lineages.

25 Chromatin Architecture Constrains Where Inversion Breakpoints Occur Over a Short-Time Scale in D. pseudoobscura Dynisty Wright, Stephen Schaeffer The Pennsylvania State University

DNA in the nucleus is nonrandomly organized into chromosomal territories, compartments, and topological associated domains or TADs. Chromosomal rearrangements have the potential to reorganize this structure, which could potentially
lead to altered gene expression. In humans, genome rearrangements can lead to altered phenotypes, genetic diseases, and cancer due to the misregulation of genes. Unlike in humans, paracentric inversions in *Drosophila* are quite frequent because they do not have the negative effects associated with the formation of unbalanced gametes because males do not recombine, and in females, aberrant meiotic products are lost in polar bodies. As a result, *Drosophila* populations can harbor extensive gene arrangement variation in populations. However, it is not clear whether natural selection acts to remove genome rearrangements because breakpoints alter gene expression because of changes in chromatin architecture. Liao et al. (2021) found that TAD structure is conserved over long-evolutionary time between *D. melanogaster* and *D. pseudoobscura*. We used the recent inversion events within *D. pseudoobscura* to determine if chromatin architecture (TAD structure) constrains where breakpoints occur over short-evolutionary time. *D. pseudoobscura* has an abundance of different gene arrangements on the third chromosome. Several hypotheses have been proposed for how new mutations are established in populations including the position effect hypothesis that suggests that inversion breakpoints produce variation for selection to act. We test whether the third chromosome breakpoints alter TAD structure in *D. pseudoobscura* and whether these disruptions altered gene expression. To determine whether breakpoints are constrained, we used seven pairs of inversion breakpoints in *D. pseudoobscura* to determine whether they disrupt TADs or fall between adjacent TADs using the TAD maps of Liao et al. (2021). Five of the 13 breakpoints occur at TAD boundaries, which is unlikely due to chance, while eight breakpoints disrupt TADs. We used RNA-Seq data from Fuller et al. (2016) to determine if there are differentially expressed genes in disrupted TADs. Of the eight disrupted TADs, seven had differentially expressed genes. Some breakpoints alter gene expression when TADs are disrupted supporting the hypothesis that position effects may contribute to inversion establishment by generating variation for selection to act.

### 26 Cis-regulatory Changes at the Fatty Acid Elongase eloF Underlie the Evolution of Sex-specific Pheromone Profiles in *Drosophila Prolongata*  
**Yige Luo**, Ayumi Takau, Takashi Matsuo, Santiago Ramirez, Artyom Kopp  
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Binary communication systems involving signaling and signal perception play a key role in sexual selection and in the evolution of sexually dimorphic traits. The selective forces and genetic changes underlying such traits can be studied in systems where sex-specific signaling and perception have emerged recently and show signs of coevolution. A promising model is found in *Drosophila prolongata*, which exhibits a species-specific increase in the number of male gustatory bristles. We find that this transition coincides with a change in cuticular hydrocarbon (CHC) profiles. *D. prolongata* males show increased amounts of the long-chained 9-pentacosene (9P) and 9-heptacosene (9H), and a coincident decrease in the shorter-chained 9-tricosene (9T), compared to females. This is in contrast with the closest relatives of *D. prolongata*, in which these CHCs are sexually monomorphic. Perfuming *D. prolongata* females with 9P or 9H reduces copulation success, suggesting that they act as sex pheromones. To unravel the genetic mechanism responsible for evolutionary transition from sexually monomorphic to dimorphic production of these mono-alkenes, we sequenced the oenocyte transcriptomes from males and females of *D. prolongata* and its relative *D. carrolli*. The top candidate gene that emerged from this analysis is *elongaseF (eloF)*, which encodes an enzyme responsible for fatty acid elongation. *eloF* shows elevated expression in *D. prolongata* males, consistent with the male-specific enrichment of long-chain CHCs in this species. Male-biased *eloF* expression in *D. prolongata* appears to be associated with a species-specific insertion of a transposable element (TE) at this locus. We are currently testing this hypothesis using chimeric reporter constructs that carry a combination of *D. prolongata* and *D. carrolli* sequences. Our results suggest that pheromone synthesis may have coevolved with chemo-sensory perception and suggest a possible molecular mechanism for the evolution of male-female communication.

### 27 Dissecting the genetic changes underlying the adaptation of the carbon dioxide receptor in the *D. suzukii* species complex  
**Alice Gadau**, Xin Yu Zhu Jiang, Sylvia Durkin, Nicolas Svetec, Li Zhao  
1) The Rockefeller University; 2) CUNY-Hunter College

Adaptation is fundamental to the survival and reproduction of living organisms. However, the genetic and neural basis underlying adaptation to new environments remains largely unknown. Novel traits allow species to invade new ecological niches and habitats. Pest species therefore provide a unique opportunity to study adaptation as they exploit new environments. One pest species, *Drosophila suzukii*, has evolved a preference to oviposit in ripe fruits instead of rotten fruit, unlike most *Drosophilids*. The *Drosophila* sensory and central nervous system may contribute to adaptive behavioral shifts. To determine candidate genes that contribute to the adaptive shift in *D. suzukii*, we identified sensory genes under directional selection in *D. suzukii* and showed differential expression between *D. suzukii* and *D. biarmipes*, *D. subpulchrella*, and *D. melanogaster* using population genetic analysis and RNA-seq analysis. From this screen, 15 genes evolved under positive selection and were significantly differentially expressed between species. One candidate, the carbon dioxide receptor *Gr63a*, is of particular interest as its sequence has diverged in *D. suzukii* yet is expressed significantly more in *D. subpulchrella*. We hypothesize that CO$_2$ is playing a role in decision-making in the *D.suzukii* species complex. By conducting a two-choice preference trial for high and low CO$_2$ concentration, we found that *D. suzukii* and *D. subpulchrella* exhibited the same indifference to CO$_2$, which is significantly different from *D. melanogaster’s* aversion. In
addition, we found that the CO2 sensing neuron is more sensitive to CO2 in D. suzukii and D. subpulchrella compared to D. melanogaster, using single sensillum electrophysiology. These data suggest that the behavior is conserved between the two species and that the evolution of Gr63a may have occurred at the regulatory region in D. subpulchrella yet the gene sequence in D. suzukii. To test this hypothesis, we use a GAL4-UAS system to dissect the role of regulatory and protein changes of D. subpulchrella and D. suzukii. Our results provide novel insight into the link between novel genotypes and phenotypes, and the complexity of the evolution of novel behavior.

28 Faster-X: Evolution of Drosophila melanogaster and Drosophila simulans Sex-biased Expression and Associated Chromatin Adalena Nanni1,2, Natalie Martinez3, Rita Graze4, Sarah Signor1, Srna Vlaho5, Sergey Nuzhdin5, Rolf Renne1,2, Lauren McIntyre1,2 1) Department of Molecular Genetics and Microbiology, University of Florida, Gainesville, FL; 2) University of Florida Genetics Institute, University of Florida, Gainesville, FL; 3) Department of Biological Sciences, Auburn University, Auburn, AL; 4) Department of Biological Sciences, North Dakota State University, Fargo, ND; 5) Department of Biological Sciences, University of Southern California, Los Angeles, CA

Antagonistic coevolution between the sexes should drive the faster evolution of reproductive-related characteristics, including sexually dimorphic morphology and behavior. Interestingly, the closely related human commensal species, D. melanogaster and D. simulans, have diverged in sexual dimorphic behaviors. These behaviors are potential targets of sex antagonism. While trends of sex-biased expression tend to be tissue-specific in Drosophila, there has been more male-biased expression observed in head and brain tissues, with an enrichment of male-biased genes on the X chromosome. However, the relationship between sex-biased expression and chromatin in the context of sex dimorphism and evolution is not well understood. Here we show evidence for faster evolution of the X for both chromatin and gene expression in D. melanogaster and D. simulans, with underlying sex-biased genes conserved between the species. We observe more sex-biased expression, specifically male-biased expression, on the X compared to the autosomes in both species. Additionally, male open chromatin marks are enriched on the X. Open chromatin marks in each of the sexes are associated (70% concordant) with sex-biased expression in the corresponding sex. These results indicate chromatin plays a role in sex dimorphism of expression and evolution.

29 Phage-derived DNAases are novel innate immune cell effectors in animals Kirsten Verster1, Gyongyi Cinege2, Zoltan Lipinszki2, Lilla Magyar2, Rebecca Tarnopol1, Eva Kurucz2, Istvan Ando2, Noah Whiteman1 1) University of California - Berkeley, Berkeley, CA; 2) Szeged Biological Research Centre, Szeged, Hungary

Symbiotic mutualisms with microbes have played an important role in the evolution of phenotypic novelties in animals. One of the most striking examples is a protective mutualism of the Acyrthosiphon pisum-Hamiltonella defensa cytolethal distending toxin B (cdtB) genes encoding CdtB, a widespread DNAase I and virulence factor, were horizontally transferred at least five times from APSE phages to insect nuclear genomes. In Drosophila ananassae, in addition to cdtB, we found two copies of cdtB fused to another apoptosis-inducing toxin gene, apoptosis inducing protein 56 (aip56). We found that expression of these phage-transferred insect cdtB and cdtB::aip56 fusion genes is upregulated in insect fat body and immune cells (hemocytes) after wasp parasitism. Loss-of-function cdtB and cdtB::aip56 mutant lines of D. ananassae were more susceptible to wasp attack than wild-type flies and show phenotypic differences in sexual maturity and development time. Our results show how potent effectors can be co-opted through symbioses to become integral components of animal immune systems.

30 Widespread introgression across a phylogeny of 155 Drosophila genomes Anton Suvorov1, Bernard Kim2, Jeremy Wang1, Ellie Armstrong2, David Peede3, Emmanuel D’Agostino4, Donald Price4, Peter Wadell5, Michael Lang6, Virginie Courtier-Orgogozo7, Jean David7,8, Dmitri Petrov7, Daniel Matute3, Daniel Schrider2, Aaron A. Comeault 1) Department of Genetics, University of North Carolina, Chapel Hill, North Carolina. USA; 2) Department of Biology, Stanford University, Stanford, CA, USA; 3) Biology Department, University of North Carolina, Chapel Hill, North Carolina. USA; 4) School of Life Sciences, University of Nevada, Las Vegas. USA; 5) School of Fundamental Sciences, Massey University, Palmerston North. New Zealand; 6) CNRS, Institut Jacques Monod, Université de Paris. France; 7) Laboratoire Evolution, Génomes, Comportement, Ecologie (EGCE) CNRS, IRD, Univ. Paris-sud, Université Paris-Saclay, Gif sur Yvette, France; 8) Institut de Systématique, Evolution, Biodiversité, CNRS, MNHN, UPMC, EPHE, Muséum National d’Histoire Naturelle, Sorbonne Universités, Paris, France; 9) Molecular Ecology & Evolution Group, School of Natural Sciences, Bangor University, Bangor, Gwynedd LL57 2DG, UK

Genome-scale sequence data have invigorated the study of hybridization and introgression, 28 particularly in animals. However, outside of a few notable cases, we lack systematic tests for 29 introgression at a larger phylogenetic scale across entire clades. Here we leverage 155 genome 30 assemblies, from 149 species, to generate a fossil-calibrated phylogeny and conduct multilocus 31 tests for introgression across nine monophyletic radiations within the
An odorant binding protein is required for mating plug formation and male fertility in *Drosophila* Nora Brown¹, Benjamin Gordon¹, Snigdha Misra¹, Caitlin McDonough-Goldstein², Andrew Clark¹, Geoffrey Findlay³, Mariana Wolfner¹ 1) Cornell University, Ithaca, NY; 2) University of Vienna, Vienna, Austria; 3) College of the Holy Cross, Worcester, MA

In *Drosophila melanogaster* and other insects, the seminal fluid proteins (SFPs) and male sex pheromones that enter the female with sperm during mating are essential for fertility and induce profound post-mating effects on female physiology and behavior. Genes encoding SFPs include some of the fastest evolving in the genome and display remarkable levels of turnover between species, likely the consequence of sperm competition and/or sexual conflict. The SFP suite in *D. melanogaster* includes several members of large gene families, including the odorant binding protein (Obp) family. Previous work in *Drosophila* has shown that some Obp genes are highly expressed in the antennae and can mediate behavioral responses to odorants, potentially by binding and carrying these molecules to odorant receptors. These observations have led to the hypothesis that the seminal Obps might act as molecular carriers for pheromones or other compounds important for male fertility in the ejaculate. Interestingly, Obps are found in the seminal fluid of several arthropod species, though the reproductive functions of these proteins in any species remains uncharacterized. By analyzing comparative RNAseq data from the male reproductive tract of multiple *Drosophila* species, we found that one seminal Obp, Obp56g, shows high male reproductive tract expression only in a subset of species in the *melanogaster* and *obscura* groups, suggesting co-option of this protein for reproductive function over evolutionary time. We used RNAi and CRISPR/Cas9 generated mutants to test the role of Obp56g, as well as the other six seminal Obps, in *D. melanogaster* fertility. Whereas male flies lacking six of the seven seminal Obp genes tested so far were fully fertile, males lacking Obp56g failed to induce post-mating responses in their mates. We found that Obp56g is expressed in the male’s ejaculatory bulb, an important tissue in the reproductive tract that synthesizes components of the mating plug. Indeed, we found males lacking Obp56g fail to form a mating plug in the mated female’s reproductive tract, which is needed to induce post-mating responses and sperm storage. We then used tissue specific RNAi experiments to show that ejaculatory bulb/duct expression of Obp56g is necessary for both mating plug formation and male fertility. Together, this work uncovers a novel role for Obp56g in reproduction and enhances our understanding of seminal fluid evolution.

A novel role for CRTC linking age-related cardiac dysfunction and fibrosis to metabolic syndrome Cristiana Dondi, Stan Walls, Anais Kervadec, Sean Zeng, Cecilia Hurtado, Karen Ocorr Sanford Burnham Prebys Medical Discovery Institute

Heart disease (HD) is a leading cause of mortality for both men and women, currently accounting for approximately 1 of every 4 deaths in the US. Obesity and diabetes greatly increase the risk of HD, although the underlying genetic and metabolic mechanisms remain unclear. CRTC, a CREB (cAMP-responsive element binding protein) -regulated transcriptional co-activator, has been identified as a “nutrient sensor” and a key regulator of metabolism in mammals and is conserved in *Drosophila*. In the liver, CRTC mediates signaling from glucagon and insulin, however, a cardiac-specific role for CRTC has yet to be identified. Using our *Drosophila* heart model, we found that CRTC null mutants (CRTC⁺/⁻) exhibit severe cardiac restriction with reduced stroke volume and tachycardia, accompanied by cardiac fibrosis, a hallmark of heart disease. Cardiac-specific knockdown (KD) of CRTC, or its coactivator CREB, mimics the heart defects of CRTC⁻/⁻. In the liver, CRTC is activated by calcineurin (CN), and in vertebrate hearts CN is known to cause cardiac hypertrophy via activation of NFAT. In the fly, overexpression of CRTC or activated CN, also known as Pp2B, also cause hypertrophy. However, in the fly cardiac KD of NFAT has no effect on function and CRTC knockout (KO) blocks all hypertrophic effect of cardiac-specific overexpression (OE) of Pp2B suggesting that, in the heart, CRTC mediates the downstream effects of CN. We also show that CRTC3 is highly expressed in zebrafish hearts and KD causes cardiac restriction, as in flies. In both fly and fish, CRTC localizes in the Z-bands as well as in myocardial cell nuclei. CRTC 1, 2 & 3 are also expressed in human iPSC-derived cardiomyocytes (hiPSC-CM) and KD of both CRTC 2&3 caused broadened action potentials, further supporting a fundamental role of CRTC in heart function. Comparative analysis of cardiac gene expression revealed that pathways involved in glucose, fatty acid, and amino acid metabolism were contra-regulated between cardiac CRTC KD and OE flies, suggesting that CRTC acts as a metabolic switch in the heart in regulation of lipid, carbohydrate, and protein metabolism. In summary, we have identified CRTC gene and its orthologues as a novel signaling pathway that maintains heart function with likely relevance linking nutrient sensing to heart disease.

glial GBA links neural lipid metabolism and proteostasis with sleep John Vaughen1, Emma Theisen2, Ina Anreiter3, Irma Magaly Rivas-Serna3, Vera Mazurak3, Thomas Clandinin3, Tom Clandinin2 1) Stanford University, Department of Developmental Biology; 2) Stanford University, Department of Neurobiology; 3) University of Alberta, Department of Agriculture, Food, and Nutritional Science

#Dros22 | 8
Lifelong brain health is sustained by macromolecule recycling through autophagy-lysosomal and proteasomal degradation pathways. How brains balance membrane recycling and intensive neurotransmission remains mysterious. To better understand the cellular underpinnings of brain membrane homeostasis, we investigated glucocerebrosidase (GBA), a critical lysosomal protein which hydrolyzes the sphingolipid glucosylceramide (GlcCer) and is commonly mutated in neurological diseases, including Parkinson’s. We show that Drosophila gba1b brains harbor widespread and heterogeneous proteostasis defects: engorged neural lysosomes appear during pupal development and persist throughout life, whereas ubiquitinated protein aggregates progressively afflict neurons and glia. Using cell-type specific Gba1b manipulations, we demonstrate that glial Gba1b is necessary and sufficient for regulating brain proteostasis.

Throughward life, whereas ubiquitinated protein aggregates progressively afflict neurons and glia. Using cell-type specific Gba1b manipulations, we demonstrate that glial Gba1b is necessary and sufficient for regulating brain proteostasis. Which glia mediate GBA’s wide-ranging effects? Depleting Gba1b in individual glia subsets caused no detectable changes, whereas ubiquitous protein aggregates progressively afflict neurons and glia. Using cell-type specific Gba1b manipulations, we demonstrate that glia mediate GBA’s wide-ranging effects? Depleting Gba1b in individual glia subsets caused no detectable changes, whereas ubiquitous protein aggregates progressively afflict neurons and glia. Using cell-type specific Gba1b manipulations, we demonstrate that glial Gba1b is necessary and sufficient for regulating brain proteostasis. Which glia mediate GBA’s wide-ranging effects? Depleting Gba1b in individual glia subsets caused no detectable changes, whereas ubiquitous protein aggregates progressively afflict neurons and glia. Using cell-type specific Gba1b manipulations, we demonstrate that glial Gba1b is necessary and sufficient for regulating brain proteostasis.

Given these complex phenotypes, we hypothesized that differential lipid accumulation occurred across time and cell-type. We conducted targeted lipidomic across multiple ages and timepoints from gba1b brains. GlcCer species were immediately elevated in young mutant brains, and ectopic GlcCer staining colocalized with neural lysosomes. Additionally and unexpectedly, younger gba1b brains harbored certain GlcCer species only at a specific zeitgeber, while these species were always detectable in older brains. We thus re-examined young gba1b brains for aggregates across time and found transient aggregates prominently in glia processes in the optic lobe chiasms. We are exploring if this aggregate clearance cycle is under circadian clock control and/or driven by light and neural activity. Do gba1b mutants show other circadian-related phenotypes? Using activity assays, we demonstrate that gba1b mutants as well as glial gba1b knockout flies are hyperactive and sleep less but still retain circadian rhythms when housed in constant darkness. Intriguingly, other short sleeping mutants also harbor ectopic aggregates in optic chiasm glia, hinting at a connection between sleep and glia proteostasis. These studies reveal compartmentalization of sphingolipid lysosomal enzymes in glia and support the hypothesis that membrane recycling via glia is a molecular function of sleep.

34 Macrophages-derived Pvf2 modulates developmental transition by ecdysone synthesis regulation Sergio Juarez-Carreño, Frederic Geissmann Immunology program, Sloan Kettering Institute, Memorial Sloan Kettering Cancer Center, NY, NY, USA.

Disturbances during development are sensed by monitoring hormones and signals released from the impaired tissues. All growing tissues and organs have the ability to release and receive information about the growth status of themselves or the different parts of the body. This communication mechanism involves hormones, peptides, and metabolites. Recent evidence suggests that macrophages act as a multi-organ rheostat that constantly senses and monitors homeostasis. Moreover, macrophages have the ability to relay signals, as growth factors, to mitigate perturbations and restore homeostasis during development. Recent studies from my host lab demonstrated that flies lacking macrophages disrupt the developmental transition between juvenile (larva) to puberty (pupa) stages. Developmental timing is regulated by sterols hormones synthesis in endocrine glands. However, how macrophages sense perturbations (such as nutrition) and communicate with the neuroendocrine system to control developmental timing remains unsolved. In this presentation, we will discuss how macrophages communicate with the neuroendocrine system. First, we proved that the lack of macrophages during development regulates the sterol hormone synthesis (ecdysone) in the prothoracic gland by reducing the Halloween genes transcription. To further understand how macrophages regulate ecdysone synthesis, we carried out a knockdown genetic screen of growth factors secreted by macrophages during development. We obtained Pvf2 as a factor required in macrophages to control developmental timing. Reduction of pvf2 expression in macrophages induces developmental delay by reducing Halloween genes transcription and ecdysone synthesis. Moreover, the lack of Pvf2 receptor, pvr, in the prothoracic gland induces developmental delay by ecdysone production inhibition. In addition, the lack of pvf2 in macrophages reduces their total number in larva stages, suggesting a mechanism where Pvr monitors the total number of macrophages in the body through Pvf2 before ecdysone pulse turn up for developmental transition in L3-instar larvae. The pvf2-depleted macrophages phenotypes are recapitulated in the pvr mutant. The developmental delay and ecdysone synthesis inhibition in pvf2 null mutant were rescued by overexpression of pvf2 in macrophages or pvr in the prothoracic gland. Therefore, our data suggest a communicative process between macrophages and the neuroendocrine system involving Pvf2/Pvr to fine-tune ecdysone synthesis in the larva to pupa transition.

35 The Drosophila enzyme L-2-hydroxyglutarate dehydrogenase is required in the renal system for recovery from hypoxic stress Nader Mahmoudzadeh1, Yasaman Heidarian1, Katherine Beebe2, Alexander Fitt1, Aylin Rodan2, Jason Tennesen1 1) Indiana University; 2) University of Utah

The oncometabolite L-2-hydroxyglutarate (L-2HG) is considered a waste-product of central carbon metabolism that is capable of disrupting chromatin architecture, mitochondrial metabolism, and cellular differentiation. As a result, ectopic L-2HG accumulation in humans is toxic and promotes the growth of renal cell carcinomas, however, few studies have
examined how this molecule functions in vivo. The fruit fly Drosophila melanogaster has emerged as a powerful model to study L-2HG. Not only are the metabolic mechanisms that regulate L-2HG accumulation conserved between flies and humans but L-2HG levels undergo predictable fluctuations during the fly lifecycle that allow for mechanistic studies. Here we exploit this system to understand how inappropriate L-2HG accumulation affects the metabolism and physiology of adult flies. Using CRISPR/Cas9, we generated mutations in the sole Drosophila L-2-hydroxyglutarate dehydrogenase (L2HGDH) ortholog, which is required to degrade L-2HG. Although L2hgdh mutant adults accumulate L-2HG levels that are 50-times higher than controls, these mutants are viable and fertile under normal conditions and exhibit no apparent phenotypes. However, L2hgdh mutant adults are extremely sensitive to hypoxic stress and die following exposure to 1% O₂. Subsequent metabolomic studies indicate that hypoxia exposed L2hgdh mutants become locked in a glycolytic state and are unable to restart mitochondrial metabolism upon reoxygenation. Finally, we find that the hypoxia-sensitive phenotype of L2hgdh mutants stem, in part, from defects in the Malpighian tubules (MTs). Not only do L2hgdh mutant MTs display mitochondrial defects, but L2hgdh mutants that express a rescuing transgene in only the principal cells of MTs exhibit normal viability following hypoxia exposure. Considering that renal cell carcinomas are the only cancer where L2hgdh is known to function as a tumor suppressor, our findings suggest that L-2HG is capable of uniquely disrupting renal cell function and establish the fly as a powerful disease model for studying this oncometabolite.

36 Differential regulation of glycogen homeostasis by TGFβ/Activin ligands Heidi Bretscher, Michael O’Connor University of Minnesota - Twin Cities

Maintaining carbohydrate homeostasis is essential for organismal health. Insufficient carbohydrates stores results in an inadequate amount of energy to fuel cellular processes. Conversely, excess carbohydrates leads to metabolic diseases such as type II diabetes. We have found that the TGFβ/Activin signaling pathway is required for homeostasis of stored carbohydrates known as glycogen. In Drosophila the TGFβ/Activin signaling family consists of three ligands: Activinβ, Dawdle and Myoglianin. All ligands signal through a single type I receptor, Baboon, however Baboon has three separate splice isoforms that differ in the ligand binding domain. Each ligand signals through a single splice isoform and tissues express a specific splice isoform. Thus, each ligand results in activation of the TGFβ/Activin signaling pathway in a specific subset of tissues. Interestingly, Activinβ is a positive regulator of glycogen storage, whereas Dawdle negatively regulates glycogen stores. This suggests the effect of TGFβ/Activin signaling on glycogen homeostasis depends on the tissue in which the signal is received.

We have found that motorneuron derived Activinβ signals directly to the muscle to positively regulates glycogen storage. Thus, activinβ mutants have low levels of glycogen in muscle. Glycogen levels in activinβ mutants can be partially rescued by inhibiting glycogen phosphorylase (GlyP), which catalyzes the breakdown of glycogen. Despite increased breakdown of glycogen by GlyP, loss of Activinβ does not result in increased glucose levels suggesting that byproducts of glycogen breakdown are being used to fuel another process.

In addition to serving as a positive regulator of glycogen storage in muscle, Activinβ is also a positive regulator of fat body glycogen levels. Intriguingly, Dawdle negatively regulates fat body glycogen storage, highlighting the tissue specific role of TGFβ/Activin signaling in maintaining glycogen homeostasis. Additionally, loss of Activinβ and Dawdle results in differing responses to a high sugar diet. Loss of Activinβ protects animals from carbohydrate accumulation on a high sugar diet, while Dawdle is required for survival on a high sugar diet.

37 Spenito, m⁴A RNA modification and the establishment of metabolic sexual dimorphism in larvae Arely V. Diaz, Kelsey Hazegh, Tânia Reis University of Colorado, Anschutz Medical Campus

Metabolism in males and females is fundamentally different. We found dimorphic expression of sex-determining genes in metabolic tissues (the fat body) and investigated the underlying molecular mechanisms. We previously found that Spenito (Nito), an RNA-binding protein and a subunit of the N6-methyladenosine (m⁴A) methyltransferase complex, is required for proper fat storage. Through its m⁴A RNA modification role, Nito is essential for splicing of sex determination genes and consequently for proper sex establishment. We propose a similar role for Nito in the establishment of metabolic sexual dimorphisms. Fat body-specific Nito knockdown recapitulated the lean phenotype and abolishes fat differences between males and females. To address the requirement of m⁴A, we knocked down other members of the complex. Fat body-specific knockdown of three other members of the m⁴A complex also made larvae lean, but the differences between the sexes were mostly preserved. At the molecular level, Nito was also required to maintain sex-specific fat body identity: Nito depletion caused mis-splicing of Sxl, a master regulator of the sex determination cascade, masculinizing the expression profile of female fat bodies. Altogether, our findings support a model in which Nito establishes differential expression of target genes in males versus females, leading to sexual dimorphisms in fat storage.

38 Acetyl-CoA mediated autoacetylation of fatty acid synthase as a metabolic switch for de novo lipogenesis in developing Drosophila Ting Miao, Hua Bai Iowa State university, Ames, IA

De novo lipogenesis (DNL) is tightly regulated during animal growth and development. It is well known that under conditions like excess nutrition, obesity, and cancer, DNL is activated through transcriptional regulation of lipogenic genes, including fatty acid synthase (FASN). Surprisingly, we show that the levels of FASN protein remain unchanged...
During *Drosophila* larval development, while its acetylation increases in fast-growing larvae and is positively correlated to the FASN enzymatic activity. This finding suggests that the regulation of FASN activity and developmental DNL is not at the transcription and translation steps; rather, it is regulated through post-translational modifications (PTMs), especially lysine acetylation. Through mass-spectrometry analysis, we identified two evolutionarily conserved lysine residues that are acetylated in both fly and human FASN proteins. One of them, K813 in fly, is located nearby the active site of the malonyl/acyltransferase (MAT) domain. Acetylation of K813 is predicted to enlarge the binding pocket of MAT domain to allow fast substrate loading. Indeed, flies with lysine to arginine substitution (K813R, acetylation deficiency mutants) show decreased lipogenesis and FASN activity, reduced body weight, and delayed pupariation. To further understand how acetylation of lysine residue K813 is regulated, we screen all major lysine acetyltransferases (KATs) and deacetylases (KDACs and Sirtuins). We identified deacetylase Sirt1 as the key negative regulator of FASN acetylation. However, none of the KATs that we examined seems to play a role in promoting the acetylation of K813. Intriguingly, our genetics and biochemical studies show that acetylation of lysine residue K813 does not require the participation of KATs, while it is mediated by acetyl-CoA levels. Through site-specific mutagenesis, we identified three residues downstream of K813 that consist of a new functional motif (R/Q-xx-G-x-G/A) of FASN, which is responsible for acetyl-CoA recognition and binding. Mutations of these residues block the autoacetylation of K813 *in vitro*. Taken together, our results uncover a novel regulation in metabolic homeostasis. It is suggested that the site-specific autoacetylation at lysine residue K813 is crucial in sensing the fluctuated fuels (acetyl-CoA) and fine-tuning FASN enzymatic activity. Thus, our findings discover a novel self-regulatory module that links signaling metabolite acetyl-CoA, lysine acetylation, and DNL.

### 39 Mechanical activation of mitochondrial energy metabolism during cell differentiation

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Mitochondria affords eukaryotes great metabolic flexibility to balance energy metabolism and cellular homeostasis. Unicellular organisms can adjust metabolic programs response to the availability of nutrients in environment. In multicellular organisms, a metabolic shift from glycolysis to oxidative phosphorylation (OXPHOS) is often observed during the differentiation of various types of stem cells and progenitors. To support proliferation, cells emphasize on glycolysis to preserve carbon sources for biosynthetic pathways. Post-mitotic cells utilize OXPHOS, a more effective way producing ATP, to power cellular activities. However, the cellular processes orchestrate this metabolic transition are largely unknown. We previously demonstrated that mitochondrial biogenesis and OXPHOS are activated in the differentiating ovarian cyst in *Drosophila*. Here, we report that mechanical forces generated by the surrounding somatic cells trigger JNK signaling and mitochondrial energy metabolism in differentiating germ cells. Somatic cells compress the differentiating cyst. Using fluorescence lifetime imaging microscopy (FLIM), we found that compression increases the membrane tension of germ cells. Abolishing somatic cells’ engulfment by genetically inhibiting Notch signaling, relieves the membrane tension on germ cells and blocks the activation of OXPHO activity as well. We carried out candidate RNAi screen on genes involved in mechanosensation and identified that the stretch-activated ion channel (SAC), Tmc, is required for OXPHOS activation. In compressed differentiating cysts, gating of Tmc maintains cytosolic Ca\(^2+\) levels, which induces the transcriptional activation of OXPHOS through a CaMKI-Fray-JNK signaling relay. Moreover, our preliminary data show that SACs mediate contraction-induced OXPHOS during the maturation of human induced pluripotent stem cell-derived cardiomyocytes (hiPSC-CMs), suggesting that mechanical activation of mitochondrial energy metabolism is evolutionarily conserved among animals.

### 48 The septate junction protein Bark beetle (Bark) is required for *Drosophila* intestinal barrier function and homeostasis

Rachel Hodge\(^1\), Martin Resnik-Docampo\(^2\), Emma Edmond\(^3\), Fernando de la Torre\(^3\), Cecilia D’Alterio\(^1\), D. Leanne Jones\(^1,2\)

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In the intestine, the epithelial barrier is maintained by tight junctions (TJs) in mammals and septate junctions (SJs) in insects. The intestinal barrier allows para-cellular flow of water, ions and nutrients across the epithelium, while maintaining food matter and microbes inside the intestinal lumen. Age-related loss of intestinal barrier function has been found across multiple species, including *Drosophila melanogaster* and humans. The age-related causes of barrier dysfunction remain unknown. The tricellular junction (TCJ) is a specialized region of the SJ where three adjacent cells meet. Previous studies by our lab indicated that mis-localization of the TCJ protein Gliotactin (Gli) is correlated with aging. Depletion of Gli in young flies leads to loss of intestinal homeostasis, including increased intestinal stem cell (ISC) proliferation, a hallmark of aging. In the embryonic epithelium, the TCJ protein Bark is required to recruit Gli to the TCJ. Therefore, we hypothesized that Bark would be required for maintenance of intestinal homeostasis and barrier function, similar to Gli. Indeed, depletion of Bark from the TCJ of enterocytes in a young fly posterior midgut (PMG) led to an increase in ISC proliferation, accelerated age-associated intestinal barrier loss, and shortened lifespan, suggesting Bark is required for maintenance of intestinal homeostasis. Antibody staining for Gli and Bark shows a decrease in intensity at the TCJ in intestines from aged flies, with a modest, but significant, increase in Bark staining at the BCJ and in the cytoplasm. Previous RNAseq data indicated that expression of SJ protein genes, including Bark, is not decreased with age.
and overexpression of Bark does not rescue age-associated loss of barrier function. Unlike in the embryonic epithelium, depletion of Gli disrupted proper localization of Bark to the TCJ in the PMG. In summary, the TCJ protein Bark is required at the TCJ to maintain intestinal homeostasis and barrier integrity in *Drosophila*. Our work on the mechanisms leading to loss of the intestinal barrier will provide insight into strategies to treat age-related gastrointestinal diseases, such as cancer.

49  **Role of Intramembrane Spastic Paraplegia Proteins in Organization of Axonal ER and ER-mitochondria Contacts in Drosophila**  
*Zeynep Ozturk*, Holy Robertson, Joe Stone, Shuler Xu, Cahir Okane University of Cambridge, Department of Genetics, Cambridge, UK

The Hereditary Spastic Paraplegias (HSPs) are a group of rare, clinically and genetically heterogeneous, inherited neurodegenerative and neurodevelopmental diseases characterised by spasticity and lower limb weaknesses. More than 80 causative genes are known, and some of them imply the importance of endoplasmic reticulum (ER)-a neuron within a neuron-function and morphogenesis in HSPs. These HSP proteins are Spastin (SPG4), Atlastin (SPG3A), Receptor Expression Enhancing Protein 1 (REEP1/SPG31) and Reticulon (SPG12), which share a common feature of one or two intramembrane hairpin domains that can recognise or drive curvature of ER membrane. Proteins of the REEP and reticulon families appear to be responsible for forming most peripheral ER tubules in yeast. In *Drosophila*, removing these families leads to fewer ER tubules in axons, of wider diameter, although there is no widespread absence of tubules. Therefore, other proteins must be involved in shaping the tubular ER network in *Drosophila* axons. Another HSP protein with predicted hairpin domains is C19orf12; this is therefore another candidate protein for contributing to shaping the axon ER network. Mutations in this gene are found in patients with autosomal recessive HSP and Neurodegeneration with Brain Iron Accumulation (NBIA). C19orf12 protein colocalises with mitochondria and ER, and with ER-mitochondria contacts. Most C19orf12 mutations are found in predicted transmembrane (TM) regions. To investigate possible roles of C19orf12 in ER and mitochondria structure and function, we have generated loss-of-function mutants of the widely expressed *Drosophila* ortholog of C19orf12, CG3740, using P element excision and CRISPR/cas9. These mutants are homozygous viable, as are quadruple mutants lacking CG3740 and all the widely expressed reticulon and REEP proteins, suggesting that these 4 proteins together are not sufficient for tubular ER formation. Testing of ER and mitochondria morphology in these mutants is in progress. For this purpose, we have also generated flies carrying a split-GFP reporter for ER-mitochondrial sites. To identify additional candidate proteins for ER shaping, we have also performed bioinformatic analyses of proteins shown by proteomic analysis to be enriched on ER tubules. These analyses have identified most of the known proteins with roles in ER shaping, implying that it may also be a good way to identify additional proteins with similar roles.

50  **The Abelson tyrosine kinase cooperates with the Nedd4-family ubiquitin ligase Suppressor of Deltex to regulate the late endosomal passage of Notch and modulate signaling activation**  
*Julio Miranda-Alban*, Nicelio Sanchez-Luege, Xiao Sun, Fernando Valbuena, Benjamin Glick, Ilaria Rebay University of Chicago

The conserved Notch signaling pathway controls a plethora of cellular processes during development and tissue homeostasis across animal species, and abnormal signaling activity of Notch has been linked to several developmental disorders and pathologies. The Notch protein is a transmembrane receptor and cleavage of Notch to release its intracellular domain (NICD) allows the latter to enter the nucleus and enforce the transcriptional output of signaling. NICD cleavage is best studied in response to receptor-ligand interactions but can also occur in the absence of ligand. As cells continuously traffic Notch receptor from the cell surface into endocytic compartments regardless of ligand presence, release of the NICD at different steps of the endocytic pathway is possible. Thus, Notch endocytic trafficking offers a unique opportunity to finely-tune signaling output. In this context, a key regulator of the endosomal passage of *Drosophila* Notch is the Nedd4-family ubiquitin ligase Suppressor of Deltex (Su(dx)), which recognizes a PPxY motif of the NICD, the same motif used by Su(dx) to promote internalization of Notch into the late endosomal lumen. As endocytic trafficking is a substantial contributor to the extraordinary variety of routes for Notch activation, careful and robust regulation of Notch at each step of this transport is critical for preventing non-physiological signaling activity. Our study offers important new mechanistic insight into how two post-translational modifiers, a kinase (Abl) and a ubiquitin ligase (Su(dx)), converge upon the same molecular motif (NICD PPxY) to confer robustness to the modulation of Notch activity during late steps of its endosomal passage.

51  **Understanding the Role of Loner in Myoblast Fusion.**  
*Amrita Shrikant Gokhale*, Donghoon Lee, Peter Keene, Elizabeth Chen UT Southwestern Medical Center
Cell-cell fusion is a fundamental process in the development of multicellular organisms, where it is involved in diverse processes such as fertilization, myogenesis, bone resorption and placenta formation. *Drosophila* embryonic musculature is an excellent genetically tractable model to study cell-cell fusion, as it results from the fusion between two types of cells – muscle founder cells and fusion competent myoblasts (FCMs). Previous studies in our lab have identified many molecular components in myoblast fusion and revealed an asymmetric “fusogenic synapse”, wherein the FCM generates an actin-enriched podosome-like structure (PLS) to invade the founder cell to promote cell membrane fusion. We previously identified an Arf GTPase guanine nucleotide exchange factor (ArfGEF), Loner, as an essential regulator of myoblast fusion. However, its precise molecular mechanism of action remains poorly understood. We found that actin-enriched PLS form at the fusogenic synapse between founder cells and FCMs in loner mutant embryos. However, time-lapse imaging data suggest that they are more diffused and unstable than the control embryos, failing to invade the founder cells properly to induce cell fusion. Our genetic rescue experiments showed that while Loner is required in both founder cells and FCMs, it plays a major role in the FCMs. Consistent with this, Loner is recruited to the fusogenic synapse in the FCMs but not the founder cells. Our structure-function analysis of Loner show that multiple domains of Loner can interact with FCM-specific cell adhesion molecule Sns, suggesting that Loner may be recruited to the fusogenic synapse by Sns. Thus, we hypothesize that Loner promotes actin nucleation promoting factors at the fusogenic synapse of FCMs. Currently, we are investigating how Loner and its downstream Arf GTPases interact and regulate the actin nucleation promoting factors.

52 Pericentrin-Like-Protein is a Kinesin-1 Adaptor that drives Centriole Motility. Matthew Hannaford, Rong Liu, Neil Billington, Zachary Swider, Brian Galletta, Carey Fagerstrom, James Sellers, Nasser Rusan National Heart Lung and Blood Institute, NIH, Bethesda, MD

Centrosomes are a key microtubule organizing center in the cell. They comprise a pair of centrioles surrounded by a matrix of proteins termed the pericentriolar material. Through microtubule (MT) nucleation centrosomes organize the mitotic spindle, cilia and flagella. To fulfill these functions, centrosomes must be motile to achieve proper positioning within the cell. Very little is understood about the different mechanisms of centrosome motility. Typically, motility is thought to be governed by the activity of MT motors, pushing or pulling on the microtubules anchored at the centrosome. In some cell types, centrioles lack PCM and microtubules, and are referred to as inactive centrioles. Inactive centrioles must be motile as their positioning is critical for asymmetric cell division. Despite this, the mechanisms of inactive centriole movement are not well understood. We investigated how inactive centrioles move in interphase cells. High resolution live imaging in Drosophila revealed that centrioles are microtubule cargo and move along the MT network in a manner involving Kinesin-1. Importantly, super resolution imaging demonstrated that Kinesin-1 localizes to the outside of the centriole in interphase cells. An RNAi screen identified Pericentrin-Like-Protein (Plp) as essential for interphase centriole movement. Through yeast-2-hybrid and an in vivo interaction assay we found that Plp interacts with the cargo binding region of the Kinesin-1 heavy chain. In vitro analysis showed that Plp and Kinesin-1 comigrate on MTs. Furthermore, autoinhibition of Kinesin-1 inhibits interaction with Plp. Relieving Kinesin-1 autoinhibition using hinge region mutations promotes robust PLP binding. Using random mutagenesis, we generated a series of mutations in Plp which ablate interaction with Kinesin-1, and significantly perturbs centriolar transport. Finally we show that centriole transport is essential for correct centrosome activity and inheritance in asymmetrically dividing neural stem cells. Our data support a model where Plp acts as a novel motor adaptor that links the centriole to the MT transport machinery, facilitating movement. In this work we propose the first detailed mechanism of how centrioles can move independently of their role as an MTOC. We will further discuss our recent in vitro and in vivo efforts to dissect the mechanism of Kinesin-1 activation by kinesin activators, which promote PLP binding and efficient centriole motility.

53 Uncovering the mechanism of BNIP3-mediated mtDNA selection in the female germline Anastasia Minenkova1, Swathi Jeedigunta1, Jonathan Palozzi1, Yun Li1,2, Thomas Hurd1 1) University of Toronto, Toronto, Ontario, Canada; 2) The Hospital for Sick Children, Toronto, Ontario, Canada

Mitochondria are essential, intracellular energy-producing organelles that contain their own DNA (mtDNA). In metazoans, mtDNA is subject to high mutation rates, inherited uniparentally, and undergoes little recombination. This makes mtDNA prone to the accumulation of deleterious mutations, which can cause severe disease. To prevent deleterious mutations from being inherited, the female germline has evolved a conserved quality control mechanism, termed mtDNA selection. During this process, mitochondria with deleterious mutations are purged from the germline. The mechanism of mtDNA selection is poorly understood; however, we have recently identified that mitochondrial autophagy (mitophagy) is required for this process in *Drosophila*. We find that the outer mitochondrial membrane mitophagy receptor BNIP3 is required for both selection and mitophagy. BNIP3 mediates the incorporation of mitochondria into autophagosomes through the interaction with the core autophagy machinery component Atg8a through its LC3 interacting region (LIR). BNIP3 also plays a minor role in the *de novo* assembly of autophagosomes, which independent of its LIR. Together, our results suggest that BNIP3-mediated mitophagy represents a generalizable mechanism for selection against deleterious mtDNA mutations, which may enable the development of strategies for the treatment of mtDNA disorders.
54 Tissue-specific chromatin profiling reveals a key role for Clock-dependent transcription in regulation of Drosophila photoreceptor homeostasis
Juan Jauregui-Lozano, Vikki Weake Purdue University, West Lafayette IN 47907

The chromatin landscape defines cellular identity in multicellular organisms with unique patterns of DNA accessibility and histone marks decorating the genome of each cell type. Further, modulation of the chromatin landscape contributes to the regulation of most nuclear processes, including replication, transcription and DNA repair. Thus, profiling the chromatin state of different cell types in an intact organism under disease or distinct physiological conditions can provide insight into how chromatin regulates cell homeostasis in vivo. The aging eye experiences physiological changes that include decreased visual function and increased risk of retinal degeneration. Although there are transcriptomic signatures in the aging retina that correlate with these changes, the gene regulatory mechanisms that contribute to cellular homeostasis during aging remain to be determined. To identify transcriptional regulatory mechanisms that contribute to the homeostasis of the aging eye, we utilized a tissue-specific approach to profile the transcriptomic and accessible chromatin landscape of Drosophila photoreceptor neuron in an aging time course. ATAC-seq and RNA-seq data integration identified 61 transcription factors that showed differential activity in aging Drosophila photoreceptors. These 61 age-regulated transcription factors included two circadian regulators, Clock (CLK) and cycle (CYC), that showed sustained increased activity during aging. When we disrupted Clock activity in adult photoreceptors using a CLK dominant negative mutant, we observed changes in expression of 15 – 20% of genes including key components of the phototransduction machinery and several eye-specific transcription factors. Using ATAC-seq, we showed that loss of Clock activity leads to changes in activity of 31 transcription factors and causes a progressive decrease in global levels of chromatin accessibility in photoreceptors. Supporting a key role for Clock-dependent transcription in the eye, disruption of Clock activity in photoreceptors also induced light-dependent retinal degeneration and increased oxidative stress, independent of light exposure. Since disruption of circadian rhythms has been recently associated with the onset of many age-related eye diseases, our data suggests that during normal aging, the circadian TF complex CLK:CYC protects the aging retina by directing gene regulatory networks that maintain expression of the phototransduction machinery and counteract oxidative stress.

55 Circadian autophagy drives longevity response to Intermittent Time-Restricted-Feeding (iTRF)
Matthew Ulgherait Columbia University Medical Center

Time-restricted feeding (TRF) has become an anti-aging treatment of great interest in recent years, with the potential to delay aging and improve health in diverse organisms from Drosophila to humans. TRF consists of restricting food intake to specific hours of the day. Because TRF simply controls the timing of feeding rather than nutrient or caloric content, TRF has been hypothesized to depend on circadian-regulated functions including many metabolic functions. Nonetheless, the underlying molecular mechanisms of TRF-mediated metabolic regulation remain unclear. To exploit the rapid genetic tools and well-characterized aging markers of Drosophila, we developed an alternate-day, intermittent TRF (iTRF) dietary regimen for flies that robustly extended their lifespan and delayed aging-dependent processes such as poly-ubiquitinated protein aggregation in muscles and intestinal dysplasia. We showed that iTRF treatment enhanced circadian-regulated transcription and that iTRF-mediated lifespan extension required molecular components of both the circadian clock and autophagy, a well conserved longevity pathway. Night-specific induction of autophagy was both necessary and sufficient to extend lifespan on ad lib diet and also prevented further iTRF-mediated lifespan extension. In contrast, day-specific induction of autophagy did not extend lifespan. Thus, these results identify “night-specific” circadian-regulated autophagy as a critical contributor to iTRF-mediated health benefits in Drosophila. As both circadian regulation and autophagy are highly conserved processes that play roles in mammalian aging, this work highlights the possibility that behavioral or pharmaceutical interventions stimulating circadian-regulated autophagy may provide people with similar health benefits such as delays in aging pathology and lifespan extension.

56 The Neuronal and Molecular Mechanisms by Which Death Perception Impacts Fly Behavior and Lifespan
Tuwin Chakraborty, Christi Gendron, Cathryn Duran, Scott Pletcher University of Michigan, Ann Arbor

Certain perceptive experiences, such as the sudden death of a companion or repeated exposure to traumatic events, have significant behavioral and physiological effects that are well established across taxa. Recently, we demonstrated that when Drosophila melanogaster perceive dead conspecifics in their environment, they exhibit behavioral and physiological changes that influence energy metabolism, stress resistance, and lifespan. Sight and serotonin signaling through receptor 5-HT2A are required for these effects to manifest upon exposure to dead. New data using neural tracing and genetic analyses identified the ellipsoid body, a neuropil in the central complex that is known to be important for multifaceted sensory integration and motor coordination functions, as an important structure underlying the effects of death perception on lifespan. To better understand how this structure transduces perceptual experience into physiological changes, we dissected the contributions of individual ellipsoid body ring neurons and found that activity of the R4d and R4m neurons were required for the effect of death perception on lifespan. To elucidate the molecular mechanisms, we executed a targeted screen and discovered that RNAi-mediated knock down of 5HT2A in R4d neurons abrogated the lifespan effects caused by death perception. We also discovered that FOXO expression, a transcription factor associated with the insulin-signaling pathway and a known regulator of aging, was required in R4d neurons.
Further evidence suggests a role for the insulin pathway in mediating the effects of death perception on lifespan; dIlp3 and dIlp5, but not dIlp2, showed increased mRNA levels in dead-exposed wild type flies compared to the unexposed control animals. In addition, this mRNA increase was reflected at the protein level as dIlp3 immunostaining was higher in dIlp-expressing neurons of dead exposed compared to the unexposed controls. At present we are testing a model in which visual input to the ellipsoid body influences insulin production to modulate psychological state, physical health, and aging.

57 Blocking cell fusion inhibits age-induced polyploidy and maintains epithelial organization in *Drosophila* Ari Dehn\(^3\), Navdeep Gogna\(^4\), Patsy Nishina\(^4\), Vicki Losick\(^3\) 1) Boston College, Chestnut Hill, MA; 2) The Jackson Laboratory, Bar Harbor, ME

A characteristic of normal aging and age-related diseases is the remodeling of a tissue’s cellular organization through polyploid cell growth. Polyploidy arises from an increase in nuclear ploidy or the number of nuclei per cell. However, it is not known whether age-induced polyploidy is an adaption to stressors or a precursor to degeneration. Here, we find that the adult fruit fly’s abdominal epithelium becomes polyploid with age through generation of large multinucleated cells that make up more than 40% of the tissue area. The syncytia arise by cell fusion, not endomitosis. Epithelial multinucleation is also a characteristic of macular degeneration, including *Ctnna1\(^{tmns}\)*, a mouse model for pattern dystrophy. Similarly, we find that the knockdown of alpha-catenin enhances multinucleation in the fly epithelium. We further show that age-induced polyploidy can be suppressed by inhibiting cell fusion revealing a means to maintain tissue organization in older animals.

58 Mechanisms of Systemic and Cellular Growth Control by Cholesterol Mette Lassen, Michael J. Texada, Lisa H. Pedersen, Kim Rewitz University of Copenhagen, Denmark

Growth control is fundamentally important for normal biological development, with nutrient availability being a key factor regulating cellular and systemic growth and maturation timing (including mammalian puberty and insect metamorphosis). Cholesterol is an essential lipid used as substrate for steroidogenesis and as a structural component of cellular membranes. Diet is a major source of cholesterol, and adipose tissue is a main cholesterol storage depot, especially in obesity. Emerging evidence indicates that cholesterol is an important regulator of cell growth and thereby plays a critical role in health and disease. In fact, cholesterol has recently been identified as a promoter of the occurrence, metastasis, and mortality of cancers such as breast and prostate cancer and glioblastoma, and this lipid has therefore become a clinically important therapeutic target. Despite this importance, the mechanism by which cholesterol regulates both normal cell growth and cancer development remains poorly understood. Since growth control is a fundamental process during development, the signaling pathways that regulate growth have been conserved between flies and humans. We have found that cholesterol’s influence on systemic and cellular growth in *Drosophila* larva arises primarily through effects on the conserved intracellular nutrient sensor Target of Rapamycin (TOR) and the superimposed insulin-like signaling system, the main hormonal system controlling systemic growth. Increasing intracellular cholesterol levels either by dietary supplementation or through knockdown of the lysosomal cholesterol transporter Niemann-Pick Type C-1a (NPC1a) increases TOR activity, leading to increased cell growth. These effects in the cells of the fat body and glia of the blood-brain barrier lead to increased insulin production, release and signaling, whereas in the prothoracic gland, cholesterol-driven TOR activity promotes endoreduplication and thus appears to alter or bypass the “critical weight” checkpoint. Cholesterol induced TOR activity appears to be tightly regulated and exhibits temporal dynamics that differ from those seen in amino acid-induced TOR activity. Dysregulation of the TOR and insulin/IGF signaling systems are frequently linked with cancers, and thus these findings may further our understanding of the links between cholesterol, hormonal signaling, and growth control in cancer development, as well as and the connection between childhood obesity and early puberty.

59 Hypoxia-dependent Control of Larval Maturation Michael Turingan, Tan Li, Savraj Grewal University of Calgary

When exposed to low oxygen (hypoxia), *Drosophila* larvae can survive by reducing their growth rate and delaying development to the pupal stage. While the mechanisms that mediate reduced growth have been described, it is less clear how hypoxia delays maturation from the larval to pupal stage. This maturation is controlled by a neuroendocrine circuit in which neuronal projections from the brain innervate the prothoracic gland (PG), an endocrine organ that produces the steroid hormone ecdysone. Activation of this circuit at the end of the larval period triggers a pulse of ecdysone to initiate maturation. We find that larvae reared in hypoxia (5% oxygen) from hatching show delayed ecdysone production (as evidenced by delayed induction of ecdysone biosynthetic genes) resulting in a ~2-day delay in larval maturation. Although hypoxic larvae are delayed in their attainment of critical weight (CW), switching larvae to hypoxia after the CW checkpoint still led to a strong developmental delay, suggesting that hypoxia can specifically act on the larval-to-pupal transition. Interestingly, our data suggest that the developmental delay seems to be independent of the classic hypoxia regulator, HIF-1α and nutrient/TOR signaling. Instead, we find a role for alteration of Ras/MAPK signaling. Expression of ecdysone biosynthetic genes, and thus ecdysone production, is controlled by the conserved Ras/MAPK signaling pathway. PG-specific activation of this pathway is sufficient to reverse the hypoxia-induced developmental
delay. Upstream of this MAP kinase pathway in the PG are multiple receptor tyrosine kinases (RTKs). One of these - Epidermal Growth Factor Receptor (Egfr) - has been reported as a major controller of ecdysone production at the end of the larval period. PG-specific RNAi-mediated knockdown of Egfr causes a marked developmental delay. In hypoxic larvae, however, Egfr knockdown in the PG does not delay development any more than hypoxia alone, suggesting that hypoxia and Egfr-knockdown are working in the same pathway. Moreover, Egf ligands, spitz and vein, which mediate autocrine Egfr signaling in the PG, show a delayed induction in hypoxic larvae. We thus find that the post-CW maturation defect in hypoxia may involve suppressed Egfr/Ras signaling in the PG. Owing to conservation in key aspects of steroid hormone regulation, this work adds to our understanding of whole-body responses to hypoxia during development.

60 The role of Jagunal protein in the establishment of cortical polarity in Drosophila melanogaster neuroblast Lelahiwat Legesse, Blake Riggs San Francisco state university

In the central nervous system (CNS), cell diversity is accomplished by asymmetric cell division, a fundamental and highly conserved process during development of multicellular life. Drosophila melanogaster neural progenitor cells, or neuroblasts (NB), are one of the best models for studying asymmetric division. Neuroblast undergoes asymmetric division generating one self-renewing daughter neuroblast and another cell known as a ganglion mother cell (GMC). An essential factor for asymmetric division is the correct establishment of cellular polarity which is required for the correct partitioning of cell fate determinants. However, it is unclear the pathway by which these determinants are distributed and arranged during asymmetric divisions. Apical basal polarity is established through an evolutionarily conserved protein complex that includes Bazooka (Baz), Par-6, aPKC. Previous research has determined that the endoplasmic reticulum (ER) divides asymmetrically during mitosis prior to neuroblast differentiation with the requirement of a novel ER protein Jagunal (Jagn). Here we propose to investigate if ER asymmetry during mitosis plays a role in cortical polarity. Through genetic cross and immunohistochemistry, I determined that in the larval neuroblast, ER does not co-localize with apical or basal cell fate determinants, Bazooka (Baz) or Prospero (Pros), respectively. Future studies will include embryo fixations to investigate if there could be a possible colocalization earlier in development as well examining if Baz is necessary for ER asymmetric partitioning during cell division. These approaches will help define the role of Jagn in asymmetric ER division and neuroblast development.

61 Exploring the role of dynein in transporting cen mRNA to the centrosome Hala Zein-Sabatto1, Li Jin2, Simon L. Bullock2, Dorothy Lerit1 1) Emory University Atlanta, GA, USA; 2) MRC Laboratory of Molecular Biology Cambridge, UK

The centrosome is a multi-functional organelle that plays a key role in nucleating and organizing microtubules, facilitating ciliogenesis, and organizing the bipolar mitotic spindle during cell division. A protein matrix known as the pericentriolar material (PCM) surrounds the centrosome and regulates centrosomal function. Various mRNAs also concentrate around the centrosome, but their functional significance is not yet fully understood. Our lab and others have identified centrocortin (cen) mRNA as forming micron-scale RNA granules near the centrosome in a cell cycle-dependent manner. We also showed localization of cen mRNA at the centrosomes is needed for error-free mitosis. However, how cen mRNA localizes to the centrosome is still unknown. Preliminary data from immunofluorescent imaging combined with smFISH reveals cen mRNA decorates astral microtubules. Further, biochemical studies show that Cen protein, a component of the cen mRNA granule, interacts with the dynein motor complex. Taken together, these data suggest that cen mRNA is transported to the centrosome via dynein-directed trafficking along microtubules. To test this hypothesis, cen mRNA localization was quantified in CRISPR-edited Drosophila embryos designed to disrupt a predicted dynein light intermediate chain binding site within the cen sequence. Similarly, cen mRNA localization was quantified in embryos of classical dynein heavy chain transheterozygous mutants. Moreover, the effect of destabilizing the microtubule network on cen granule localization to the centrosome was analyzed. Inhibiting the interaction of cen mRNA with dynein and microtubules resulted in a decrease in cen mRNA granules localized to the centrosome, consistent with a role for dynein in RNA trafficking to this site. Furthermore, fly lines harboring N- and C-terminal truncations of the cen transcript were made to identify the minimal sequence required for cen mRNA localization to the centrosome. This work provides insight into the dynamic localization of cen mRNA through active intracellular transport.

62 Elucidating the mechanism of coactivator Taiman/AIB1-driven cell competition and its relation to the Adenomatous Polyposis Coli (APC) tumor suppressor in Drosophila Colby Schweibenz, Ken Moberg Emory University

The phenomenon of cell competition ensures that the fittest cells populate developing primordia but is also postulated to underlie the phenomenon of “field cancerization,” in which cancer cells expressing ‘super-competitor’ genes eliminate slow growing neighbors and take over an epithelial tissue. Our prior work demonstrated that cells overexpressing the Drosophila protein Taiman (Tai; human NCOA3/AIB1), a transcriptional co-activator of the Ecdysone receptor (EcR), are able to kill wild type neighbors within the larval wing epithelium in a manner dependent on production of the Toll
ligand Spätzle (Byun et al, 2019). Here we use the wing disc to test and confirm the reciprocal hypothesis, that cells with reduced Tai expression (Tai<sup>low</sup>) are competitive ‘losers,’ and we use a genetic screen in the adult eye to identify candidate mechanisms required for elimination of these cells by wild type neighbors. This screen recovered ‘hits’ in the pro-apoptotic genes, <i>head involution defective</i>, <i>reaper</i>, and <i>grim</i>, as dominant suppressors of the loss of Tai<sup>low</sup> cells, confirming a competitive mechanism that operates through the classic DIAP-RHG-Caspase apoptotic pathway. In addition to these validating ‘hits’, we also recovered alleles of factors involved in cell:cell signaling pathways that are thus candidates to act downstream of Tai to control competitive fate. Our studies have focused on two of these: the two <i>Drosophila</i> Adenomatous polyposis coli (APC) tumor suppressor homologs, Apc1 and Apc2, which are conserved elements of the Wg pathway and inhibit field cancerization in the fly midgut (Suijkerbuijk, 2016). We find that Apc1/Apc2 loss rescues elimination of Tai<sup>low</sup> cells in both eye and wing epithelia, and couple this with evidence that Tai controls some Wg-target genes in larval wing cells. We will describe ongoing experiments to establish mechanistic links between Tai and Wg/Apc in the wing epithelium, with the goal of defining how the Tai/EcR and Wg/Apc pathways intersect to determine winner/loser status in <i>Drosophila</i> epithelia.

63  **Cell-surface proteomic profiling identifies key regulators in epithelial cell competition** Ke Li<sup>1,2</sup>, Juan Oses-Prieto<sup>2</sup>, Alma Burlingame<sup>2</sup>, Lily Yeh Jan<sup>1,2</sup>, Yuh Nung Jan<sup>1,2</sup> 1) Howard Hughes Medical Institute; 2) University of California, San Francisco

Cell competition defines an evolutionarily conserved, cell-cell interaction mechanism by which more fit cells (“winners”) eliminate and replace suboptimal cells (“losers”). One classic example is wild-type cells eliminating the adjacent cells bearing mutated or reduced ribosomal protein genes (Rp), emphasizing the significance of the cell-surface interaction between competing populations. Despite recent progress in identifying intracellular molecular events underlying cell competition, little is known regarding how distinct cell populations recognize and compare their relative fitness. Moreover, although mechanical forces have been proposed to regulate cell competition, the components mediating mechanical responses remain elusive. Given the particular interests in the competing interface, we labeled proteins on winner and loser cell surfaces in a spatiotemporal-specific manner and created a quantitative profile using a multiplexed proteomics approach. Using clonal knockdown of <i>Rp</i> in imaginal epithelia, we find ~250 and ~500 proteins are significantly enriched in winner and loser surface proteomes, respectively. Consistent with previous findings, we find autophagy-related proteins are enriched in loser cells, whereas engulfment-related proteins are overrepresented in winner cells. Remarkably, our proteomics data indicates wild-type cells have lower levels of non-muscle myosin expression compared to cells with reduced <i>Rp</i>. Wild-type cells indeed have reduced junctional tension, and restoring junctional tension suppresses cell competition. Interestingly, mechanosensitive channels are highlighted in the winner cells, coincident with their higher calcium signaling. Our results further demonstrate that mechanical stress is required for cell competition, identifying candidate genes mediating mechanical sensing. Efforts are currently underway to perform a proteome-instructed screen to identify novel regulators of cell competition and these results will also be presented.

64  **Hypoxia-dependent regulation of epithelial tissue growth and development** Abhishek Sharma, Savraj Grewal Department of Biochemistry and Molecular Biology, University of Calgary, Canada

Our cells and organs need oxygen from the air we breathe in order to survive and function. However, in certain disorders - such as stroke, heart disease and cancer - tissues are often deprived of oxygen. <i>Drosophila</i> larvae provide an excellent in vivo model to study adaptive responses to hypoxia as they have evolved to live in naturally low oxygen conditions. We previously identified two mechanisms that promote organismal hypoxia tolerance – activation of the FOXO transcription factor and Tsc1/2-mediated suppression of TOR kinase signaling. Here we examine how both mechanisms operate to control organ-level growth and proliferative adaptation to hypoxia. To do this we examined <i>tsc</i> and <i>foxo</i> function in the developing larval imaginal disc epithelial tissues. Cell clones mutant for both <i>tsc1</i> and <i>foxo</i> show an overgrowth phenotype under normal growth conditions. We find that this overgrowth is exacerbated under hypoxia and leads to altered tissue patterning. Moreover, we also find that <i>tsc1/foxo</i> double mutant clones are susceptible to high amount of apoptosis as compared to their wildtype counterparts and that this cell death is also further exacerbated under hypoxic conditions. In contrast to loss of <i>tsc1/foxo</i> we see that clones overexpressing oncopgene <i>yorkie</i> do not show enhanced growth under hypoxic conditions. We are currently exploring mechanisms by which hypoxia exposure enhances the overgrowth of <i>tsc1/foxo</i> mutant tissue. The behaviour of <i>tsc1</i>, <i>foxo</i> double mutant clones under hypoxia recapitulate various tumor phenotypes such as high proliferation and apoptosis. Hence our work will help uncover how hypoxia impacts tumor growth.

65  **Late Endosomes act as carriers for delivery of Ceramide phosphoethanolamine (CPE) with unique acyl chain anchors to cleavage furrows during male meiosis cytokinesis.** Govind Kunduri<sup>1</sup>, Si-Hung Le<sup>2</sup>, Baena Valentina<sup>3</sup>, Nagampalli Vijaykrishna<sup>4</sup>, Daniel Blankenberg<sup>4</sup>, Yoshihiro, Izumi<sup>5</sup>, Takeshi Bamba<sup>2</sup>, Kedar Narayan<sup>3</sup>, Usha Acharya<sup>1</sup>, Jaira Acharya<sup>1</sup> 1) Cancer and Developmental Biology Laboratory, Center for Cancer Research, National Cancer Institute; 2) Department of Metabolomics, Kyushu University; 3) Center for Molecular Microscopy, Center for Cancer Research, National Cancer Institute; 4) Genomic Medicine Institute and Lerner Research Institute, Cleveland Clinic, Cleveland
Cell division, wherein one cell divides into two daughter cells, is fundamental to all living organisms. Cytokinesis, the final step of cell division, begins with the formation of an actomyosin contractile ring, positioned midway between the segregated chromosomes. Constriction of the ring with concomitant membrane addition in a spatiotemporal manner generates a cleavage furrow that physically separates the cytoplasm. Unique lipids with specific biophysical properties have been shown to localize to midbodies, however, their delivery mechanisms and biological roles remain largely unknown. In this study, we show that Ceramide phosphoethanolamine (CPE), the structural analog of sphingomyelin, has unique acyl chain anchors in spermatocytes and is essential for meiosis cytokinesis. The head group of CPE is also important for spermatogenesis. We found aberrant central spindle and contractile ring behavior but not de-localization of phosphatidyl inositolts (PIPs) at the plasma membrane was responsible for the male meiosis cytokinesis defect in CPE deficient animals. Further, we demonstrate the enrichment of CPE in Rab7 and Rab11 positive endosomes which in turn translocate to the cleavage furrows to promote cytokinesis. Volume electron microscopy analysis using Correlative light and focused ion beam scanning electron microscopy showed that CPE enriched Rab7 positive endosomes appear as giant, interconnected multivesicular bodies that are juxtaposed on contractile ring material. Genetic ablation of Rab7 alone results in significant meiotic cytokinesis defects. Our results imply endosomal delivery of CPE to ingressing membranes is crucial for male meiotic cytokinesis.

### 66 Immunosstimulatory Lipids in Drosophila Bacterial Infection

Sophia Parks, Susan Nguyen, Daiki Fujinaga, Naoki Yamanaka, Adler Dillman
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Eicosanoids are C20 polyunsaturated fatty acid (PUFA) derivatives that carry out many essential roles in mammalian and insect systems including development, reproduction, and immunity. Eicosanoids such as prostaglandins and leukotrienes are synthesized from arachidonic acid that is cleaved directly from the cell membrane. Insects, however, have been shown to have low levels of C20s in circulation and it has been hypothesized that eicosanoids are synthesized from C18 precursors such as linoleic acid. Here we show that *D. melanogaster* exhibits higher levels of C18s in the hemolymph and that these are depleted after a bacterial infection. These depleted oxylipins as well as certain downstream prostaglandins are able to rescue the outcome of infection. This work identifies oxylipins that are involved in immunity and supports the notion that *D. melanogaster* utilizes immunostimulatory and immunosuppressive lipid signaling mechanisms to mitigate bacterial infections. Our understanding of immune signaling in the fly and its analogies to the mammalian system will allow for an even more detailed use as a model organism in immune studies.

### 67 Interorgan transfer of intact micron-sized lipid droplets to macrophages during the Drosophila immune response

Ishneet Kaur, Amber Myers, Eniola Ogundipe, Catherine Brennan
Cal State University Fullerton

Macrophages are white blood cells of the immune system in both mammals and Drosophila. In addition to their roles in engulfing and killing microbial invaders by phagocytosis, these cells play important roles in embryonic development, tissue repair, metabolism, and maintenance of homeostasis. Under certain inflammatory conditions, mammalian macrophages accumulate lipids in the form of lipid droplets, including in infection, atherosclerosis, and obesity; however, neither the mechanisms nor purpose of this accumulation are well-understood. We have found the fly macrophages [plasmatocytes] also accumulate lipid droplets, with astonishing speed following bacterial infection: filling up to 30% of cell volume within 3 hrs of infection. We have discovered a completely novel mechanism for this inflammatory lipid accumulation: instead of synthesizing lipids from fatty acids or glucose taken up from the blood, fly macrophages obtain intact, micron-sized lipid droplets from the fat body, through a phagocytosis-like mechanism we term “phagoliposis”. By blocking the metabolism of these lipid droplets, we have discovered some unexpected inflammatory mechanisms fueled by these lipids.

### 68 Paying the amino acid cost of the humoral immune response to bacterial infection

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Immunity and metabolism are closely intertwined. Immune responses are resource-demanding and metabolic changes are therefore required to the supply the immune system with the required energy and raw materials. In pathological infections where the immune response becomes dysregulated, these metabolic changes can contribute to pathology. Work using the model organism *Drosophila melanogaster* has been productive in uncovering how the immune system regulates metabolism, but details of actual metabolic changes that occur during infection, and their functions, are poorly known. This project seeks to uncover the metabolic changes that occur during the *Drosophila* innate immune response to infection and understand the consequences of these changes. By combining mass spectrometry-based metabolomic analysis with genetic approaches, we found that successful immune responses to systemic bacterial infection caused an increase in nitrogenous waste metabolism. Further investigations suggested that this increase in nitrogenous waste is due to upregulation of endogenous protein degradation via an ESCRT-III mediated process. Expression of CHMP2B,
encoding a core ESCRT-III component, is indicated to be required in fat body for the optimal translation of antimicrobial peptide transcripts. Together these data suggest a model in which the fly upregulates degradation of its own protein in order to rapidly provide in-demand amino acids for immune effector production, with catabolism of the resulting excess amino acids causing an increase in nitrogen waste metabolism. This project has thus revealed a profound metabolic change induced by a successful immune response in *Drosophila*. As the cost of this change is the degradation of the host’s own protein, this a probable mechanism of metabolic pathology in the context of a dysregulated immune response. Studying the etiology of such metabolic pathology will contribute to our understanding of conditions underpinned by metabolic dysregulation, such as sepsis and infection-induced cachexia.

### 69 The cytoplasmic incompatibility factor proteins CifA and CifB are both nucleases in *Drosophila melanogaster* Rupinder Kaur, J. Dylan Shropshire, Brittany Leigh, Seth Bordenstein, 1) Department of Biological Sciences, Vanderbilt University, Nashville, TN, USA; 2) Vanderbilt Microbiome Initiative, Vanderbilt University, Nashville, TN, USA; 3) Division of Biological Sciences, University of Montana, Missoula, MT, USA; 4) Department of Pathology, Microbiology & Immunology, Vanderbilt University Medical Center, Nashville, TN, USA; 5) Vanderbilt Institute for Infection, Immunology and Inflammation, Vanderbilt University Medical Center, Nashville, TN, USA

The widespread symbiosis between invertebrates and *Wolbachia* bacteria is due in part to cytoplasmic incompatibility (CI) that is significant to arthropod vector control and evolution. While CI selectively favors the fitness of *Wolbachia*-transmitting females and thus the bacterial spread, the CI mechanism is not broadly understood. In *Drosophila melanogaster*, dual expression of the genes *cifA* and *cifB* from wMel causes CI, resulting in embryonic lethality between symbiotic male and aposymbiotic female cross. These two co-diverging genes span five phylogenetic Types, and previous work suggested different CifB Types are distinguishable by the presence/absence of nuclease activity. Here, we re-evaluate this claim using transgenic, mutant, enzymatic, and cytochemical assays *in vitro* and *in situ*. We demonstrate that contrary to prior conclusions, T1 CifB is an *in vitro* DNase that cleaves single- and double-stranded DNA. Moreover, we provide the first characterization of T1 CifA as a nuclease that cleaves both DNA and RNA. We further show that *in vitro* DNase activity translates to *in situ* spermatid DNA fragmentation in Cif-expressing and *Wolbachia*-bearing testes, and CifB-mediated spermatid DNA damage alone is insufficient to cause CI. Cif enzymatic activity is ablated by deletions and amino acid substitutions. In sum, these molecular insights (i) highlight previously unrecognized nuclease functions of T1 CifA and CifB proteins (ii) reinforce a conservative gene nomenclature and mechanistic model that is agnostic to presumptions about distinct Cif enzymatic activities and (iii) support a postulate of the Host Modification model of CI that spermatid DNA is altered by the Cif proteins.

### 70 A symbiotic niche in the Drosophila gut regulates the stable association of a multispecies community Ren Dodge, Eric Jones, Haolong Zhu, Benjamin Obadia, Chenhui Wang, Andrés Aranda-Díaz, Kevin Aumiller, Zhexian Liu, Marco Voltolini, Eoin Brodie, Kerwyn Casey Huang, Jean Carlson, David Shivak, Allan Spradling, Will Ludington, 1) Carnegie Institute of Washington; 2) University of California, Santa Barbara, CA; 3) Department of Biology, Johns Hopkins University; 4) Department of Microbiology and Immunology, Stanford University School of Medicine; 5) Molecular and Cell Biology Department, University of California; 6) Howard Hughes Medical Institute; 7) Simon Fraser University, Burnaby, BC; 8) Lawrence Berkeley National Lab; 9) Chan Zuckerberg Biohub

Animal guts are colonized by a complex community of host-specific commensal bacteria that is relatively stable over time within an individual and can have life-long effects on health. The microbiome is established and maintained in the face of daily fluctuations in diet, invasion by pathogens, and disruptions by antibiotics. Many gut resident bacteria localize to specific regions of the gut that correspond to chemical environments matching the specific species’ metabolism. Certain probiotic species, namely Lactobacilli, additionally make physical attachments with host mucus, stabilizing their colonization. A key gap in our knowledge is how the host constructs microenvironments that promote colonization by specific multispecies communities of bacteria. Here, we show that a physical niche exists within the proventriculus region of the Drosophila foregut that selectively binds bacteria with exquisite strain-level specificity. Using gnotobiotic flies, microscopy, and microbial population kinetics we found that primary colonizers saturate the niche and exclude secondary colonizers of the same strain. Conversely, initial colonization by Lactobacillus physically remodels the niche to favor secondary colonization by another commensal, Acetobacter. Our results provide a mechanistic framework for understanding the establishment and stability of a multispecies intestinal microbiome. We anticipate this model will form the basis for dissecting the host genetics of an intestinal niche as well as for the discovery of similar niches in Drosophila and other animals, including humans.

### 71 The Turandot proteins promote tolerance to stress by regulating energy consumption and tracheogenesis Samuel Rommelaaere, Alexa L. Carboni, Jean-Philippe Boquete, Shu Kondo, Bruno Lemaître, 1) Global Health Institute, School of Life Science, École Polytechnique Fédérale de Lausanne (EPFL), 1015 Lausanne, Switzerland; 2) Invertebrate Genetics Laboratory, Genetic Strains Research Center, National Institute of Genetics, Mishima, Japan

*Turandot* (*Tot*) genes code for a family of height small secreted proteins. They are highly expressed in response to a variety of stresses. Tots are transcriptional targets of several stress pathways, including the JAK/STAT and p38...
MAPK pathways. Although *Tots* have been extensively used as readouts of the stress response, they have never been functionally characterized. We have generated a *Drosophila* line that lacks 6 *Tots* (*TotXMAZ*). These mutant flies are extremely susceptible to several challenges, including bacterial infection, starvation and osmotic stress. Phenotypically, *TotXMAZ* flies have reduced triglyceride content. During stress exposure, their lipid and carbohydrate stores are rapidly depleted, suggesting that energy availability is a limiting factor in the stress response of these flies. Indeed, feeding *TotXMAZ* flies a high-sugar diet improves their survival to osmotic stress. *Tots* may thus participate in the stress response by tempering energy utilization. Turandot proteins secreted by the fat body specifically bind to tracheas. Interestingly, tissue tracheation is reduced in *TotXMAZ* mutant and restoring tracheal growth in these flies is sufficient to rescue their susceptibility to stress. Altogether, our data suggest that Turandot proteins participate in the adaptation to stress by controlling tracheal growth and nutrient utilization. We propose that Turandot may adjust the fly metabolic rate by controlling tracheal homeostasis under stress conditions.

**72 Regulation of Misshapen during Border Cell Migration** Gabriela Molinari Roberto, Gregory Emery Institute for Research in Immunology and Cancer. Université de Montréal

**Introduction:** Collective cell migration is an important process during development, wound repair and metastasis. Border cells (BC) migration in Drosophila egg chamber is a powerful model to study collective cell migration in vivo. Misshapen (Msn) is a key regulator of this process that coordinates protrusion formation with contractile forces through the border cell cluster. However, the molecular mechanism regulating Msn in border cells is still unknown. Msn is composed of a kinase domain and a Citron homology (CNH) domain separated by a long coiled-coil. The kinase domain was shown to be phosphorylated by the kinase Tao in other contexts, while CNH domains are described to be bind to small GTPases. Therefore, our hypothesis is that both Tao and small GTPases regulate Msn activity during border cell migration.

**Methods and Results:** Border cell clusters depleted for Tao or Msn are phenotypically identical: they do not entirely detach from the follicle epithelium and they present ectopic protrusions. Interestingly, the expression of a Msn phosphomimetic form restores the migration phenotype induced by Tao depletion. This indicates that the main function of Tao in BCs is to phosphorylate and activate Msn. Furthermore, we found that the CNH domain of Msn is required for its recruitment at the periphery of the cluster, suggesting that its interaction with a GTPase is required for its recruitment. Using the expression of mutant form of Rho and Rap GTPases and co-immunoprecipitation, we found that Rap1 and Rap2L might regulate Msn. In particular, we found that Msn binds preferentially to Rap2L. The role of Rap2L in BC migration is unknown, but preliminary data shows that its depletion induces a migration phenotype. Further analysis of the lost-of-function phenotype and its effects on Msn localization will be performed.

**Conclusion and Relevance:** Here we presented two mechanisms of regulation of Msn in BC migration. We show that Tao acts as Msn upstream kinase to activate it in BC. Furthermore, we found that Msn binds to Rap GTPases with a preference to Rap2L, which might regulate its recruitment at the periphery of the BC cluster. The Msn ortholog, MAP4K4, is found dysregulated in many cancers and related to metastatic capacity and has been linked to Rap2 and Tao kinases. Therefore, our findings suggest that a conserved molecular cascade regulates the collective cell migration in BCs and during cancer progression.

**73 Investigating mechanisms regulating actin assembly in the early Drosophila embryo** Anna Yeh¹, Julian Eskin², Bruce Goode², Adam Martin² 1) MIT; 2) Brandeis University

Actin networks undergo dynamic rearrangements that promote morphogenesis in the developing embryo. Therefore, actin network dynamics must be tightly regulated in space and time. The range of mechanisms that control actin network assembly and organization during development is still not fully understood. The *Drosophila* Synaptotagmin-like protein, Bitesize (Btsz), has been shown to organize actin at epithelial cell apical junctions in a manner that is thought to depend on its interaction with the actin-binding protein, Moesin. Through the maternal depletion of Btsz by RNA interference (RNAi), we discovered that Btsz functions at earlier, syncytial stages of development. In the syncytial embryo, nuclei separated by metaphase furrows undergo rounds of synchronous divisions. Btsz-RNAi disrupts these metaphase furrows, leading to spindles collisions and nuclear division defects, similar to mutants in the formin Diaphanosus. Interestingly, we identified an amino acid motif in Btsz, that is similar to a formin elongation effector domain (FEED) found in yeast formin regulators. Additionally, purified Btsz protein containing FEED stimulates formin activity and directly interacts with F-actin through the C-terminal half of the protein *in vitro*. To test the functional requirement of Btsz isoforms that contain this motif, we used CRISPR/Cas9 to engineer a *btsz* mutant, that specifically targets FEED-containing isoforms. Our preliminary data suggest FEED-containing isoforms and to a lesser extent, MBD-containing isoforms are important for the syncytial stages of development. This work suggests a novel mechanism regulating actin assembly during animal development.

**74 Discovery of a novel competitive interaction between the Chlamydia trachomatis early effector Tarp and the endogenous actin bundler Singed/Fascin during mechanosensory bristle development** George Aranjuez, Travis Jewett Immunity and Pathogenesis Division, College of Medicine, University of Central Florida, Orlando, FL

*Chlamydia trachomatis* infection is the most frequently reported sexually transmitted infection in the United States.
As an obligate intracellular bacterial pathogen, *Chlamydia* injects multiple protein effectors via the type III secretion system (T3SS) into the host cell to induce its entry. The early effector Tarp is required for efficient host cell entry by *Chlamydia* since a ΔTarp mutant has significantly reduced invasion frequency. Tarp is a ~150kDa protein with a tyrosine-rich N-terminal region and a C-terminal region that holds actin nucleating as well as F-actin bundling activities. Interestingly, removing Tarp’s F-actin bundling domain also impacts *Chlamydia* invasion. Studying the F-actin bundling activity of Tarp within the host cell has not been explored. Here, we use *Drosophila* as an in vivo cell biology platform to study Tarp’s F-actin bundling activity and how it impacts host cell biology. Ubiquitous expression of Tarp results in reduced adult viability and tissue-specific expression leads to various defects consistent with its ability to manipulate host actin. We used mechanosensory bristles as a model to study Tarp’s F-actin bundling activity since the establishment of bristle shape is dependent on proper F-actin bundling. Expressing Tarp in the thorax of wildtype flies leads to increased curvature of mechanosensory bristles. The growing pupal bristle is supported by parallel longitudinal F-actin bundles and the actin bundler Singed, required for normal bristle shape, localizes with the F-actin bundles. Surprisingly, Tarp expression in the bristles displaced Singed away from F-actin bundles. Tarp’s competitive behavior against Fascin during F-actin bundling was confirmed in vitro using co-sedimentation assay. Loss of either *singed* or *forked* in flies leads to highly deformed bristles. Strikingly, Tarp partially rescued the bristle morphology defect caused by *singed* knockdown. This work demonstrates the utility of *Drosophila* in studying bacterial effector function. Moreover, we provide in vivo confirmation of Tarp’s F-actin bundling activity and further uncovers a competitive behavior against Singed/Fascin during F-actin bundle assembly.

75 **Sufficiency of active Rac to drive whole tissue phagocytosis in vivo** *Abhinava Mishra*, Lauren Penfield, Denise Montell University of California Santa Barbara

The Rho family of small GTPases, Rac, Rho, and Cdc42 represent central nodes in the cytoskeletal and signaling networks that drive cell migration and engulfment. How the cells utilize the same RhoGTPase networks to promote different processes in different contexts is not well understood. We address this question in the *Drosophila* ovary, which contains ~850 somatic follicle cells and 16 much larger germline cells. Previously we reported that local and transient activation of Rac using a photoactivatable Rac (PA-Rac) induced protrusions and motility in border cells. However, persistent expression of a constitutively active form of Rac (Rac-CA) in just a subset of follicle cells, using a cell-type-specific enhancer (*slbo-Gal4*), destroyed the entire egg chamber, a phenotype that is significantly suppressed by mutation of a single engulfment receptor, Draper. Additionally, border follicle cells expressing Rac-CA by *slbo-Gal4* engulf adjacent polar cells in early-stage 8 egg chambers. To further assess the effects of Rac-CA mediated engulfment, we generated FpoutGal4 driven clones to express Rac-CA in a subset of follicle cells. These clones displayed a non-autonomous expression of Draper in follicle cells. Therefore, the expression of Rac-CA in a few cells destroys the whole tissue likely by stimulating engulfment.

To understand the mechanisms underlying Rac-CA mediated tissue destruction, we performed fixed tissue imaging with cell death markers. We observed that expression of Rac-CA caused caspase activation in follicle cells non-autonomously and increased acidification of germline nurse cells evident by lysotracker staining. Live imaging of follicle cells expressing Rac-CA by *slbo-Gal4* unveiled rapid and synchronous death of germline nurse cells. It was initiated by biting the nurse cell membrane by border follicle cells. Expression of Rac-CA by stretch follicle cell-specific *PG150-Gal4* also results in egg chamber death suggesting that the Rac-induced egg chamber death shares some features with late-stage developmental programmed cell death.

These results indicate that focal and transient activation of Rac promotes protrusion and motility, whereas high and sustained Rac activation promotes phagocytosis, initially of living cells, ultimately destroying the whole tissue. These results offer a novel explanation for a previously mysterious human immunodeficiency.

76 **WAVE regulatory complex facilitates cell rearrangements through the generation of an F-Actin subpopulation at tri-cellular junction in the follicular epithelium** *Lisa Calvary*, Piere Pouchin, Graziella Richard, Vincent Mirouse iGReD, Université Clermont Auvergne - CNRS UMR 6293- INSERM U1103 - Clermont-Ferrand, France

How cells change their respective position to allow a tissue acquiring a specific shape is a major question in development. *Drosophila* follicle elongates through its antero-posterior axis and this elongation depends on the follicular epithelium. We recently described that this elongation relies on two steps, the first one occurring from stage 3 to 7 and involving, among other mechanisms, cell intercalations (Alégot et al., *eLife*, 2018). However, in these cells there is no junctional planar polarization of Myosin II as it is the case in some other epithelia undergoing cell intercalations. In this context, we got interested in the role of F-Actin in cell rearrangements.

By performing a reverse genetic screen of F-Actin regulators, we identified the WAVE regulatory complex (WRC), a complex involved in the generation of branched F-Actin, as required for follicle early elongation. In follicular cells, WRC is localized at tri-cellular junctions and colocalized with an intense spot of F-Actin. Tricellular junctions are important hotspots for tissue dynamics thought their function is not yet fully understood. In mutant cells for WRC subunits, this
spot of F-Actin is absent suggesting that it reflects the protrusive activity of this complex. Accordingly, high-resolution analysis reveals that these spots correspond to protrusions from one cell between the bicellular junction of the two others. In addition, mutants cells for WRC exhibit a strong packing defect, leading to the conclusion that WRC is involved in cell rearrangements. We therefore propose that the protrusive activity of WRC at tri-cellular junctions promotes cell intercalations, allowing follicle elongation. Data of our current analyze on living samples of the relationship between F-Actin protrusions induced by WRC and cell junction dynamics will be presented.

77  Fat2 polarizes the WAVE complex to align protrusions for collective cell migration Audrey Williams, Seth Donoughe, Edwin Munro, Sally Horne-Badovinac University of Chicago

Many epithelial tissues form tubes and spheres, rather than the sheets often studied using epithelial cell culture models. Despite their closed topology, examples of spherical or tubular tissues undergoing collective cell migrations have emerged. The follicle cells of the ellipsoidal Drosophila egg chamber undergo one such collective migration. These epithelial cells, which make up the outermost cell layer of the egg chamber, migrate along an encapsulating basement membrane with high directional persistence for days. In the absence of a free edge or another guidance cue to dictate their migration direction, they must coordinate their migration through local interactions within the migrating cohort. One important regulator of follicle cell migration is the atypical cadherin Fat2, which localizes to the trailing edge of each cell and promotes leading edge protrusion of the cell behind. Another is the WAVE regulatory complex, which templates the formation of lamellipodial protrusions at each cell's leading edge. Using genetic manipulations, live imaging, and quantitative image analysis, we show that Fat2 polarizes protrusive activity in trans by concentrating the WAVE complex at the leading edge, across the cell-cell interface. Without this cue from Fat2, the WAVE complex remains active and the cells are protrusive, but protrusive sites fluctuate around the cell periphery and collective cell migration fails. To stabilize the orientation of WAVE complex enrichment and protrusive activity, clusters of Fat2 recruit the WAVE complex to corresponding clusters just across the cell-cell interface, at the tips of filopodia embedded within the lamellipodium. Because the Fat2-WAVE complex signaling system is deployed at each leading-trailing interface in a planar polarized manner, it both polarizes protrusions within individual cells and couples these individual cell polarities across the epithelium. This allows the cells to exert force in a common direction and achieve a highly coordinated collective cell migration.

78  Defining the role of prostaglandins in collective cell migration Samuel Mellentine, Anna Ramsey, Omar Rabab'h, Tina Tootle University of Iowa Carver College of Medicine

Collective cell migration – the coordinated movement of associated cells – is important for both normal development and tumor invasion. Prostaglandins (PGs) are short-range lipid signals that regulate cell migration and are up regulated in many cancers. However, their mechanisms of action during collective migration are poorly understood. Here we use the native, collective migration of the Drosophila border cells to uncover the roles of PGs. During Stage 9 of oogenesis a cluster of epithelial cells becomes specified as border cells, delaminates from the epithelium, and migrates collectively and invasively between the nurse cells. Prior work found that loss of Pxt, the Drosophila cyclooxygenase-like enzyme responsible for all PG synthesis, results in delayed migration and decreased cluster cohesion. However, the particular PG or PGs controlling border cell migration remain unknown. To begin to address this, we assessed the roles of three PGE synthases, (mPGES-1, mPGES-2, and cPGES) and the sole PGF2α synthase (PGFS) in border cell migration. Our data support the model that cPGES and PGFS are required for on-time border cell migration. Specifically, loss of cPGES or PGFS delays border cell migration, but has no effect on cluster cohesion. We are currently using cell-specific RNAi knockdown to determine which cells produce PGE and PGF2α. Initial studies suggest that, cPGES acts in the nurse cells to promote border cell migration. We are also assessing genetic interaction between the synthases and Pxt. We find that co-reduction of both Pxt and PGFS results in delayed border cell migration; supporting that the phenotype due to loss of PGFS is related to loss of PG signaling. Together our data lead to the model that PGE and PGF2α both promote on-time border cell migration. As PG signaling is highly conserved, these studies provide critical insight into the specific functions of individual PG signaling cascades controlling collective cell migration and can be applied to understanding both developmental collective cell migrations and pathological migrations including cancer metastasis.

79  Microtubule acetylation promotes epithelial cell stretching and squamous cell carcinogenesis in Drosophila Rachita Bhattacharya, Nitin Mohan, Pradip Sinha Indian Institute of Technology Kanpur, India

The cytoskeleton is an essential regulator of cell shape and size changes, of which microtubules (MT) are a crucial component. Indeed, MT re-organize during epithelial cell shape transition, for instance, from columnar to the squamous cell shape. The mechanism that regulates these MT-mediated cell shape remodeling, however, is not known. Here, we show that MT acetylation is an essential step during epithelial cell stretching. For instance, MT acetylation increases remarkably during columnar-to-squamous transition. Conversely, loss of MT acetylation via genetic downregulation of Tubulin Acetyl Transferase (TAT) compromises the cell stretching by dysregulating the turnover of polarity protein, Crumbs, and cell-cell adhesion protein, cadherin. We further show that neoplastic transformation of squamous epithelium is accompanied by hyperacetylation of MT. Our results, therefore, reveal an essential mechanism of MT-
Maternal pioneer factor CLAMP regulates sex-specific transcript diversity in early Drosophila embryos. Mukulika Ray¹, Ashley Conard¹, Jennifer Urban³, Annie Huang¹, Erica Larschan¹ 1) MCB department, Brown University, Providence, RI; 2) CCMB department, Brown University, Providence, RI; 3) Johns Hopkins University, Baltimore, Maryland

Post-transcriptional processes of alternative splicing of RNA are essential in gene expression, critical for transcript diversity, eventually leading to cellular differentiation and specification. Alternative splicing is initiated during the initial embryonic stages when the key gene expression regulators are maternally deposited transcription factors (TFs). Yet, it is unknown how transcription factors (TFs) work with splicing factors to regulate alternative splicing. Also, how maternal factors shape the early embryonic transcriptome in a sex-specific manner is not well understood. Here, we show that the maternally deposited transcription factor CLAMP (Chromatin-linked adaptor for MSL proteins) is essential for sex-specific alternative splicing in early embryos of both sexes. We found that CLAMP binds along with the gene bodies of sex-specifically spliced genes in both sexes and differentially binds to RNA-binding protein components of the spliceosome complex in males and females. In females, CLAMP also modulates the chromatin environment at the sxl gene, the master regulator of sex-determination, affecting its splicing and thus downstream female-specific splicing events. In males, CLAMP influences the distribution of RNA-helicase Maleless (a component of spliceosome complex and Male Sex-lethal (MSL) complex in males) on chromatin, affecting male sex-specific splicing. We show that CLAMP is an RNA-binding protein through iCLIP (individual-nucleotide resolution Cross-Linking and Immunoprecipitation) and RNA gel shift experiments and is important for the dynamics of formation of nuclear speckles of ribonucleoprotein origin that constitute alternative splicing complexes. Using CLAMP mutants, we demonstrate that the prion-like domain (PrLD) of CLAMP inhibits aggregation of RNA-binding splicing factors that renders them non-functional and is essential for survival. We hypothesize that the PrLD domain alters the phase transition properties of CLAMP. Thus, we define a new function for a maternally deposited TF in regulating sex-specific alternative splicing via interaction with chromatin and RNA-binding proteins, which influences the distribution of RNA-binding proteins on chromatin and between protein complexes. As a part of this study, we have also developed a computational pipeline called time2splice for the identification of alternative splicing events which change over time and between sexes.

Hippo pathway transcriptional regulators alter chromatin binding dynamics of the transcription factor Scalloped Samuel Manning¹,², Benjamin Kroeger¹,², Elizabeth Hinde³, Kieran Harvey¹,² 1) Monash University, Melbourne, Australia; 2) Peter MacCallum Cancer Centre, Melbourne, Australia; 3) The University of Melbourne, Australia

The Hippo signalling pathway is a highly conserved regulator of cell fate and organ size, which regulates the nuclear-cytoplasmic shuttling of the transcriptional co-activator Yorkie (Yki), and thereby regulates gene expression. We currently have very little understanding of how the Hippo pathway modifies the nuclear biophysical behaviour and genomic interactions of Yki or its DNA binding partner Scalloped (Sd) to regulate transcription. To address this, we used cutting-edge ex vivo fluorescence microscopy approaches including Raster Image Correlation Spectroscopy (RICS), Fluorescence Recovery After Photobleaching (FRAP) and Single Molecule Tracking (SMT) to investigate the biophysical behaviour of Sd and Yki in the nuclei of living Drosophila tissues. We found that Yki moves more rapidly through the nuclear environment than Sd, and that Yki's movement is highly sensitive to Sd abundance, demonstrating the importance of Sd for Yki's navigation of the genome. Sd and Yki were highly enriched at regions of active chromatin, consistent with their role as a heterodimeric activator of transcription. Further, enrichment of Sd at these loci is driven in part by increased chromatin residence times. Yki interacts more transiently with chromatin than Sd at these sites, and the duration of this interaction is dependent on Sd levels, further highlighting the importance of Sd levels for recruitment of Yki. FRAP and SMT demonstrate that Yki and the transcriptional repressor Nerfin-1 alter Sd chromatin residence times, providing mechanistic insight into how these proteins alter Hippo pathway output. Taken together, we show that a transcriptional co-activator and repressor are able to modify the binding duration of a transcription factor, with subsequent impacts on transcription. These results also indicate that altering binding duration is one mechanism by which Hippo pathway transcription factors can be enriched at a specific locus to regulate gene expression. These results have implications for understanding how the Hippo pathway regulates normal development and disease, and also provide mechanistic frameworks for understanding how transcription may be regulated by other signalling pathways.

Hox linker domain phosphorylation alters Exd-Hox DNA-binding preferences and regulates gene expression William Glassford, Richard Mann Zuckerman Mind Brain Behavior Institute, Columbia University, New York, NY

Hox proteins are transcription factors best known for patterning segment identity along the anterior-posterior axis in early embryogenesis. As monomers, Hox proteins exhibit little difference in DNA-binding specificity, however; upon binding their common cofactor Extradenticle (Exd) each Hox paralog gains the DNA-binding specificity necessary to enact acetylation-dependent cell shape regulation as well as squamous cell carcinogenesis and hints at potential therapeutic targets.
Assembly of the Brain tumor RNA decay pre-complex expedites downregulation of Notch signaling following asymmetric stem cell division


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RNA-binding proteins assemble translational control complexes by recruiting their interactors that exert regulatory effects on target mRNAs. How these multi-protein complexes are assembled in a timely manner to promote dynamic changes in gene regulatory programs that promote proper specification of cell identity during key developmental transitions remains poorly understood. During asymmetric division of type II neuroblasts in fly larval brains, RNA-binding protein Brain tumor (BRAT) segregates into one of the neuroblast progeny in which it expedites the decay of Notch target gene transcripts allowing for the onset of differentiation less than 60 minutes after division. The tight binding protein Brain tumor (BRAT) segregates into one of the neuroblast progeny in which it expedites the decay of Notch target gene transcripts leading to supernumerary neuroblast formation.

We confirmed that NOT1 and PAN3, the scaffolding components of two main RNA deadenylation complexes, co-segregate with BRAT in mitotic neuroblasts and that their segregation is BRAT-dependent, consistent with our previous findings that Brat functions together with these proteins to promote the decay of Notch target gene transcripts. We also identified Ubiquitin specific protein 5 (USP5) as a novel BRAT interactor by IP-MS experiment using embryonic extract. We confirmed that USP5 co-segregates with BRAT into neuroblast progeny destined to differentiate, and that asymmetric USP5 separation depends on BRAT. Furthermore, reducing usp5 function enhanced differentiation defects in *brat*-hypomorphic brains, and complete loss of usp5 leads to supernumerary neuroblast formation phenocopying brat mutant. By using single-molecule fluorescent RNA *in situ* hybridization, we demonstrated that neuroblast progeny ectopically accumulate Notch target gene transcripts in *brat* or *usp5*-mutant brains, supporting the model that USP5 is a key component of the BRAT RNA decay complex. The scaffolding protein Miranda (MIRA) binds and segregates BRAT in mitotic neuroblasts. Consistent with Miranda inhibiting the RNA-binding ability of BRAT in wild-type neuroblasts, we detected co-localization of Notch target mRNAs with mutant BRAT that is defective in MIRA-binding in mitotic neuroblasts. We propose that BRAT assembles an enzymatically inactive and non-RNA-binding pre-complex that asymmetrically segregates BRAT in mitotic neuroblasts. Consistent with Miranda inhibiting the RNA-binding ability of BRAT in wild-type neuroblasts, we detected co-localization of Notch target mRNAs with mutant BRAT that is defective in MIRA-binding in mitotic neuroblasts. We propose that BRAT assembles an enzymatically inactive and non-RNA-binding pre-complex that asymmetrically segregates BRAT in mitotic neuroblasts.

Using CRISPRi to uncover mechanisms of transcriptional repression by Rb and CtBP co-repressors

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CRISPR activation and interference (CRISPRa/i) have revolutionized gene expression studies, allowing us to precisely regulate genes through targeting activator and repressor domains across the genome. With CRISPRa/i, we can turn genes on and off with high precision and perform large-scale LOF and GOF screens in many systems. Yet, this modified CRISPR system is not often used for studying the mechanisms of gene regulation by transcription factors. Here, we have adapted the CRISPRi system for targeting transcriptional co-repressors to gene promoters to uncover their mechanisms of repression. We engineered dCas9 fusions to the Retinoblastoma family proteins (RB) and the C-terminal binding protein (CtBP), which are highly conserved co-repressors across Metazoa. In *D. melanogaster*, the Rb family consists of the Rbf1 and Rbf2 paralogs, while a single CtBP gene encodes the CtBP-long and CtBP-short isoforms, which differ by about 100
residues in the CTD. We expressed the dCas9-Rb and dCas9-CtBP chimeras in the fly wing for in vivo targeting to specific promoters, and tested each protein’s effect on target gene expression and their mechanism of repression. We found that on some genes, both Rbf1 and Rbf2 are able to mediate potent repression, but that this happens in a distance-dependent manner. In contrast, we found examples where Rb proteins modestly activate the target gene, suggesting that they do not always function to turn off transcription. Notably, an Rbf1 CTD mutant lacking the Instability Element (IE), which we previously identified as being necessary for activity, was just as good a repressor as the wild type protein. This suggests that the IE is chiefly involved in recruiting and is not necessary for repression. Between the CtBP isoforms, CtBP-short was a much more potent repressor than CtBP-long, which indicates that the long CTD extension may have an inhibitory role in transcription. Our novel adaptation of a well-established CRISPR tool has allowed us to probe mechanisms of repression in vivo in the fly and compare factors in the same contexts with great precision. This molecular analysis of promoter-specific regulation by Rb and CtBP proteins will enhance our understanding of these conserved regulatory proteins in development and disease.

85 Sculpture of a sex-specific piRNA program Peiwei Chen, Alexei Aravin California Institute of Technology

Maternally deposited piRNAs act as epigenetic vectors to pass on transgenerational memory of selfish genetic elements to the offspring. This model posits that the maternal input instructs the zygotic genome to mount a piRNA program in the next generation that reflects the maternal response to genomic parasites. However, males implement a piRNA program distinct from the maternal instruction and their female siblings. Here, we disentangled the contribution of three factors—sex chromosome content, gonadal sex, and maternal input—towards a sex-specific piRNA program. While maternally deposited piRNAs can identify homologous sequences as piRNA-producing loci in the zygote, we found de novo piRNA production that does not rely on maternal input but instead depends on the chromosome content and gonadal sex. In fact, Y chromosome is both necessary and sufficient to recapitulate some key aspects of male piRNA program. Meanwhile, sexual identity exerts a major influence on divergent piRNA production from identical genomic sequences between sexes. Our work showed that it is the collective action of chromosome content, gonadal sex and maternal input that sculpts the observed sexual dimorphism in piRNA program, highlighting a previously unknown input from sexual identity into piRNA biogenesis.

86 The steroid hormone Ecdysone coordinates larval growth and development through its interaction with the transcriptional repressor Smrter. Joanna Wardwell-Ozgo, Colby Schweibenz, Douglas Terry, Ken Moberg Emory University School of Medicine

Hormones act as systemic signaling cues to coordinate developmental timing, growth, and tissue patterning. In Drosophila, pulses of the steroid hormone ecdysone (Ec) synchronize various physiological and developmental processes required for larval ecdysis, tissue growth, and morphogenesis. Binding of the bioactive form of ecdysone (20-hydroxyecdysone; 20E) to its cognate nuclear hormone receptor, Ecdysone Receptor (EcR), is proposed to switch target genes from a transcriptionally repressed to activated state. Current data indicate that EcR has tissue-specific targets, and that even within a single cell type, developmental fluctuations in 20E levels also coincide with dynamic redistribution of EcR between different loci. Despite a long history of genetic analysis of EcR, mechanisms that direct these temporal/cell-type specific transcriptional responses to 20E are not well defined. To probe this repression-to-activation switch in a single tissue, we created a Gal4/UAS-regulated “EcR sponge” transgene composed of the ligand-binding domain (LBD) region of EcR (EcRLBD) fused to RFP. Expression of EcRLBD represses heterologous EcR transcriptional reporters in wing disc cells when 20E titers are high (late L3), and reciprocally activates EcR activity in these same cells during early L3 when 20E levels are low, indicating that EcRLBD interacts with key elements of the repression-to-activation switch. Indeed, loss of the transcriptional repressor Smrter (NCoR) mimics the derepressing effect of EcRLBD in early L3 wing cells, suggesting that Smrter is an EcR-associated repressor in larval disc cells. Consistent with this hypothesis, a version of EcRLBD carrying the A483T mutation, which blocks Smrter binding, (termed ‘EcRdmber’) represses EcR transcriptional reporter activity regardless of developmental age. With regard to endogenous target genes, EcRLBD, but not EcRdmber, relieves repression of multiple EcR targets in wing disc cells and disrupts some, but not all, 20E-EcR-triggered developmental processes in the larval salivary gland and fat body. Significantly, expression of EcRLBD confers a growth advantage to larval wing cells and derepresses a subset of growth genes controlled by the Hippo pathway. In sum, these data reveal distinct molecular roles of 20E-EcR signaling at individual promoters, and suggest that 20E-EcR signaling coordinates proliferation and tissue growth in part through repression of Hippo pathway target genes. Current work centers on proteomic analysis of EcRLBDinteractors in larval cells to identify key activators and repressors that are active in these mechanisms.

87 Embryo development requires histone acetylation by Nejire during the maternal-to-zygotic transition Audrey Marsh1, George Hunt2, Tyler Gibson3, Elizabeth Larson3, Katherine Hullin1, Mattias Mannervik2, Melissa Harrison1 1) Department of Biomedical Chemistry, University of Wisconsin School of Medicine and Public Health, Madison, WI; 2) Department of Molecular Biosciences, The Wenner-Gren Institute, Stockholm University, 10691, Stockholm, Sweden

The sperm and egg are differentiated cell types, individually specialized for the purpose of fertilization. After merging...
to form the diploid genome, the new embryo must remove signatures in the parental genomes to reprogram to a totipotent state. Reprogramming is essential for the embryo to give rise to all the cell types of the adult organism. This reprogramming process occurs during a period called the maternal-to-zygotic transition (MZT). Initially transcriptionally quiescent, the zygotic genome is gradually activated by maternally encoded factors loaded into the oocyte prior to fertilization. Activation of the genome is controlled by a specialized set of transcription factors called pioneer factors, defined by their unique ability to bind and increase chromatin accessibility. In *Drosophila melanogaster*, the pioneer factor Zelda (ZLD) is an essential activator of zygotic transcription. Prior to ZLD-mediated genome activation, chromatin is largely comprised of nucleosomes devoid of post-translational modifications. Intriguingly, ZLD-bound sites are enriched with active histone acetylation marks, suggesting that a histone acetyltransferase (HAT) is functioning with ZLD to induce transcription. We showed that during the MZT Nejire (NEJ), the *Drosophila* homologue of the HAT family p300/CBP, bound to thousands of genomic loci that are also occupied by ZLD and that expression of ZLD in tissue culture resulted in increased acetylation at ZLD-bound loci. In the early embryo, ZLD knock down resulted in decreased acetylation and NEJ recruitment at shared sites. Together these data suggest ZLD mediates activation of the zygotic genome, at least in part, by recruiting NEJ to acetylate histones. To test whether NEJ-mediated acetylation during the MZT is required for development, we used the CRY2 optogenetic system to specifically inactivate NEJ during the period of widespread zygotic genome activation. Embryos treated with blue light through the MZT failed to hatch as compared to sibling controls left in the dark, indicating that NEJ is necessary during this early time in development. Our findings suggest a model in which pioneer factors are first to prime the genome for activation by recruiting histone acetyltransferases to cis-regulatory regions and that this is required for genome activation. Ongoing work is investigating the role of NEJ in activating the zygotic genome and establishing the mechanistic relationship between NEJ and the pioneer factor ZLD.

88 Color augments motion vision for detecting approaching objects in *Drosophila* Kit D Longden, Edward M Rogers, Aljoscha Nern, Heather Dionne, Michael B Reiser HHMI Janelia Research Campus

Color and motion are used by many species to identify salient moving objects. They are processed largely independently, but color contributes to motion processing in humans, for example, enabling moving colored objects to be detected when their luminance matches the background. In *Drosophila*, motion vision was thought to be independent of color vision, based on experiments with blue and green gratings; color inputs to the luminance system had been identified through neurogenetics and connectomics, but these were thought to expand the wavelength sensitivity of the luminance channel. Here, we demonstrate an unexpected, additional contribution of color to motion vision in *Drosophila*, using a novel display system to present ultraviolet (UV) and green visual patterns to tethered flying flies. We show that *Drosophila* respond to looming UV discs at all intensities, indicating that color contributes to their motion vision. We further show that behavioral ON-motion responses are more sensitive to UV than for OFF-motion, and this difference is present in all six *Drosophila* species we tested. Using genetic rescue and silencing, we show that ON-motion UV-sensitivity depends on the UV-sensitive R7 photoreceptors, and using calcium imaging of neural activity we show that ON-motion encoding T4 cells are more sensitive to UV than OFF-motion encoding T5 cells. Calcium imaging of the cells connecting the photoreceptors to T4 reveals a diversity of UV-sensitivity, in which the UV-sensitivity of medulla T4 input cells correlates with that of their most numerous lamina monopolar cell inputs, notably M19 and L3. In a parallel connectomics study, we discovered that the majority of R7 inputs to L3 are found in the optic chiasm between the medulla and lamina, and so had been missed in previous work. We hypothesized that if the ability of flies to respond to all intensities of looming UV discs was explained by ON-motion sensitivity to UV, then flies should be motion blind to looming green discs with a UV background, even when they have the same chromatic contrast as UV discs that drive behavioral responses. Remarkably, behavioral experiments confirmed this hypothesis. Together, these results demonstrate that color contributes to motion vision in *Drosophila* in a way that favors the detection of approaching UV objects. We propose that UV objects are of particular interest to flies, and illustrate how this mechanism generalizes for the detection of approaching colored objects in other visual systems.

89 A hymenopteran odorant alerts flies to bury eggs Shaun Davis1, Gregory Chism3, Megan Maurer2, Julio Trejo1, Ricardo Garcia3, Todd Schlenke1 1) University of Arizona, Tucson, AZ; 2) Arizona State University, Tempe, AZ

Ants are ubiquitous and consume insects at all life stages, presumably creating a strong selective pressure for ant avoidance behaviors across insects. The insect egg stage can be especially defenseless against predation given that eggs are usually immobile and unguarded, suggesting insect mothers may have evolved oviposition strategies to minimize the ant predation risk to their offspring. Given the lack of parental care in most insects, these oviposition strategies would likely be innate rather than learned, since insect mothers are not usually present to assist predation of their eggs. Here, we use the vinegar fly *Drosophila melanogaster* as a model system for examining parental defensive responses to ant presence. Flies usually lay eggs partially inserted into the food substrate, although some are laid on top of the food and a few are inserted deeply into the food. We found that exposure to ants significantly alters fly oviposition depth: the proportion of eggs on the food surface decreased while the proportion of buried eggs increased. Buried eggs survive ant foraging bouts better than surface eggs, showing that this oviposition depth behavior is adaptive. This induced behavior is conserved across the genus Drosophila and is dependent on the fly olfactory system: anosmic mutant flies fail to bury
The investigations into the functional role of these genetic changes will allow us to identify how aversion to some toxins is mediated by gustatory receptors whose homologs in *D. melanogaster* are expressed in all, or most, bitter sensitive sensilla. Further work will be needed to identify the families involved in gustatory reception in the peripheral nervous system. Notably, many of these changes were among those associated with the herbivorous lineage experienced significant gene losses, gains, and changes in selective constraint among several gene families involved in detoxification. These changes were likely driven by co-diversification with their host plants over the last 400 million years. During this dynamic dietary shift, herbivores need to evolve detoxification mechanisms to overcome toxic plant secondary compounds, but also behavioral changes to preferentially feed and lay eggs on their new host. While highly specialized herbivores use these toxins or their precursor compounds as feeding and oviposition cues, the extent of behavioral plasticity is not well understood. The extent and nutritional quality of food ingested as larvae determines the adult size and ultimately fecundity. Studies on *Drosophila* larval feeding behavior have shown that under different nutritional conditions, multiple neuronal circuits regulate and maintain feeding. But most of these studies focused on the later stage of larval feeding and less is known on how constant feeding is maintained in second instar larvae. Here we have identified a subset of dopaminergic (THD') neurons that maintain constant feeding in early larvae. The function of STIM protein (an ER calcium sensor) in this subset of dopaminergic neurons is important to maintain their excitability and dopamine release. Increasing the activity in THD'-dopaminergic neurons through genetic means rescued the feeding deficit of *STIM*KO larvae. Through optogenetic methods, we have established that THD'-dopaminergic neurons are synaptically connected to a subset of neuroptidergic cells. Although dopaminergic neurons are known to regulate feeding behavior under different conditions, this is the first study showing how a subset of dopaminergic neurons regulate larval growth and pupariation by maintaining constant feeding.

Feeding is a vital and complex behavior essential to acquire nutrition. In *Drosophila melanogaster*, a holometabolous insect, the extent and nutritional quality of food ingested as larvae determines the adult size and ultimately fecundity. Studies on *Drosophila* larval feeding behavior have shown that under different nutritional conditions, multiple neuronal circuits regulate and maintain feeding. But most of these studies focused on the later stage of larval feeding and less is known on how constant feeding is maintained in second instar larvae. Here we have identified a subset of dopaminergic (THD') neurons that maintain constant feeding in early larvae. The function of STIM protein (an ER calcium sensor) in this subset of dopaminergic neurons is important to maintain their excitability and dopamine release. Increasing the activity in THD'-dopaminergic neurons through genetic means rescued the feeding deficit of *STIM*KO larvae. Through optogenetic methods, we have established that THD'-dopaminergic neurons are synaptically connected to a subset of neuroptidergic cells. Although dopaminergic neurons are known to regulate feeding behavior under different conditions, this is the first study showing how a subset of dopaminergic neurons regulate larval growth and pupariation by maintaining constant feeding.

Herbivorous insects account for half of all insect species—their evolutionary success is hypothesized to have been driven by co-diversification with their host plants over the last 400 million years. During this dynamic dietary shift, herbivores need to evolve detoxification mechanisms to overcome toxic plant secondary compounds, but also behavioral changes to preferentially feed and lay eggs on their new host. While highly specialized herbivores use these toxins or their precursor molecules to identify their hosts, in the early stages of herbivore evolution, the taste system must first lose or reduce their ancestral aversion to these toxins, while maintaining discriminatory capabilities to distinguish varying levels or different chemical classes of the toxin. The latter may be advantageous for evolutionarily young herbivores lineages that lack sophisticated detoxification systems. How these gustatory adaptations are achieved at the molecular and neural level remains unclear. The herbivorous drosophilid *Scaptomyza flava* belongs to a lineage that has recently evolved herbivory (~15 million years). This species feeds on mustard plants (Brassicaceae), whose main defense compounds are isothiocyanates, highly reactive electrophiles, derived from glucosinolates, which are used by many mustard specialists as feeding and oviposition cues. Using a dye-based feeding assay, we demonstrate that *S. flava* has lost aversion to several glucosinolates, compared to closely related non-herbivores (*S. pallida*, *S. shui*, and *Drosophila melanogaster*), while maintaining aversion to the aliphatic glucosinolate sinigrin and other non-mustard plant toxins (caffeine and several glucosinolates). Further investigations into the functional role of these genetic changes will allow us to identify how aversion to some toxins is maintained, while lost for others.

The mRNA-binding protein *Pumilio* pleiotropically regulates food-related phenotypes through foraging. The regulation of two or more phenotypes by a single gene, behavioural pleiotropy, is a common occurrence. At the molecular level, these genes need to have structural and/or regulatory features that allow for this pleiotropy to occur. Here we dissect the regulation of independent *Drosophila* larval phenotypes through gene product specific regulation of the foraging gene by the 3'UTR binding protein *Pumilio*. foraging regulates larval movement on food (pathlength), food-intake, fat stores and nociception. We found that *Pumilio* binds the 3'UTR of foraging mRNA transcripts, regulating expression of the gene in a transcript and protein isoform-specific manner. While overall protein and mRNA levels are upregulated in *Pumilio* hypomorph mutants, specific transcripts are downregulated. We find that this transcriptional
regulation of foraging by Pumilio plays a role in regulation larval food-intake, fat stores and nociception, but not larval pathlength. Interestingly, we find that in the case of larval fat stores and nociception, reduced Pumilio phenocopies the effect of a foraging over-expression phenotype, while in the case of larval food-intake reduced Pumilio phenocopies the effect of a foraging null phenotype. Together these findings suggest that Pumilio is a transcriptional regulator involved in the molecular regulation of pleiotropy by independent regulation of distinct products of a single gene, foraging.

93 Genetic Variation in Cocaine Preference in the Drosophila melanogaster Genetic Reference Panel Jeffrey Hatfield, Trudy Mackay, Robert Anholt Clemson University, Center for Human Genetics, Greenwood, SC

Studies on the genetic basis of susceptibility to cocaine addiction in human populations are quite challenging due to limited sample sizes, heterogeneity of genetic backgrounds, and environmental variability. Drosophila present a powerful model for investigation into the genetic underpinnings of cocaine addiction, using preference for cocaine as a proxy for addiction behavior. Utilizing the Microplate Feeder Assay (MFA), I quantified cocaine preference for 16,442 flies across 103 distinct genetic backgrounds of the Drosophila melanogaster Genetic Reference Panel (DGRP). I provided individual flies with an equal choice between 10µL of a control liquid food (4% sucrose, 1% yeast extract, and 0.004% FD&C Blue #1) and 10µL of the same liquid food supplemented with 0.02% cocaine. Solutions were delivered with the MFA, and consumption of each solution was quantified using a plate reader following a 22-hour exposure. Normalized cocaine preference was calculated for each fly as the difference in consumption between the two solutions divided by their total consumption. I found significant, naturally-occurring genetic variation for cocaine preference across this subset of the DGRP. I also observed significant sexual dimorphism across tested DGRP lines, where male flies on average exhibit higher cocaine preference than female flies of the same line. Overall, males of 18.4% of tested DGRP lines preferred cocaine-supplemented food over control food, while females of only 3.9% of lines preferred the cocaine solution. Estimates of broad sense heritability of consumption were calculated using individual level data, as well as using DGRP line means, and were found to be $H^2 = 0.19$ and $H^2 = 0.97$, respectively. Here, we demonstrate the impact of genetics on susceptibility to cocaine addiction behavior in D. melanogaster. These data will facilitate future genome-wide association analyses, as well as identify D. melanogaster genetic backgrounds from the DGRP that can better model cocaine addiction.

94 Descending neurons coordinate anterior grooming behavior in Drosophila LI GUO, Neil Zhang, Julie Simspoon University of California, Santa Barbara

The brain coordinates the movements that constitute behavior, but how descending neurons (DNs) convey the myriad of commands required to activate the motor neurons of the limbs in the right order and combinations to produce those movements is not well understood. Here, we used fly anterior grooming to address this question. Anterior grooming consists of two basic movements, head sweeps and front leg rubbing. The analysis of the anterior grooming structure by Automatic behavior Recognition System (ABRS) showed that head sweeps and front leg rubbing are normally coupled, but alternate for the efficient dust removal. Using the optogenetic activation, we identified 3 different groups of command-like DNs that can induce front leg rubbing (DNg11), or head sweeps (aDN), or the alternation of both (DNg12). This demonstrates that there are multiple control modes for anterior grooming at the level of descending neurons, which DNs can evoke either the whole alternation program or the specific subroutines. Head sweeps and front leg rubbing both use the front legs and so are mutual exclusive. Parallel optogenetic activation of DNg11 and aDN resulted in normal alternation between head sweeps and front leg rubbing, and DNg12 itself can induce both movements. These results demonstrate that the VNC can resolve conflicting descending drives. Using FlyWire to reconstruct neurons in a whole-brain electron microscope dataset (EM), we identified a novel inhibitory circuit connecting head sensory neurons to DNg11 that command front leg rubbing. This suggests that decisions between actions can also be made in the brain. Lastly, we explored descending control of limb coordination in these movements by unilateral activation of DNs. Interestingly, unilateral activation of DNg12 or aDN1 induced single-side head sweeps, while unilateral activation of DNg11 induced bilateral front leg rubbing. Our results suggests that left and right leg movements can be decoupled in head sweeps but not in front leg rubbing. Using EM, we found a neural circuit in the brain indirectly connecting the left and right DNg11, which can potentially explain how unilateral activation can result in bilateral execution. Taken together, these results demonstrate that distinct descending neurons can orchestrate the complex alternation between the movements that make up anterior grooming.

95 Drosophila females receive male substrate-borne signals through specific leg neurons during courtship Caroline Fabre, Eleanor Mckelvey University of Cambridge

Substrate-borne vibratory signals are thought to be one of the most ancient and taxonomically widespread communication signals among animal species, including Drosophila flies. During courtship, the male Drosophila abdomen tremulates to generate vibrations in the courting substrate. These vibrations coincide with nearby females becoming immobile, a behavior that facilitates mounting and copulation. It was unknown how the Drosophila female detects these substrate-borne vibratory signals. We present data showing that the immobility response of the female to the tremulations is not dependent on any air-borne cue. We show that substrate-borne communication is used by wild Drosophila and that the vibrations propagate through those natural substrates (e.g., fruits) where flies feed and court.
We examine transmission of the signals through a variety of substrates and describe how each of these substrates modifies the vibratory signal during propagation and affects the female response. Moreover, we identify the main sensory structures and neurons that receive the vibrations in the female legs, as well as the mechanically gated ion channels Nanchung and Piezo (but not Trp) that mediate sensitivity to the vibrations. Together, our results show that Drosophila flies, like many other arthropods, use substrate-borne communication as a natural means of communication, strengthening the idea that this mode of signal transfer is heavily used and reliable in the wild. Our findings also reveal the cellular and molecular mechanisms underlying the vibration-sensing modality necessary for this communication.

96 **chimmo-mutant spermatogonial stem cells cause mitotic drive by evicting non-mutant neighbors from the niche** CHEN YUAN TSENG1, Michael Burel3, Michael Cammer1, Sneh Harsh1, Maria Sol Flaherty1, Stefan Baumgartner2, Erika Bach1 1) NYU Grossman School of Medicine, New York, NY; 2) Department of Experimental Medical Sciences, Lund University, 22184 Lund, Sweden

Niches maintain a finite pool of stem cells via restricted space and short-range signals. Stem cells compete for limited niche resources, but the mechanisms regulating competition are poorly understood. Using the *Drosophila* testis model, we show that germline stem cells (GSCs) lacking the transcription factor Chinmo gain a competitive advantage for niche access. Surprisingly, chimmo1− GSCs rely on a new mechanism of competition in which they secrete the extracellular matrix protein Perlecan to selectively evict non-mutant GSCs and then upregulate Perlecan-binding proteins to remain in the altered niche. Over time, the GSC pool can be entirely replaced with chimmo1− cells. As a consequence, the mutant chimmo allele acts as a gene drive element: the majority of offspring inherit the allele despite the heterozygous genotype of the parent. Our results suggest the influence of GSC competition may extend beyond individual stem cell niche dynamics to population-level allelic drift and evolution.

97 **Role for local ecdysone signaling in *Drosophila* imaginal wing disc regeneration** Douglas Terry, Can Zhang, Joanna Wardwell-Ozgo, Colby Schweibenz, Ken Moberg Emory University

Ecdysone steroid hormone (Ec) signaling is required to drive *Drosophila* developmental transitions and to promote growth of larval imaginal discs. Previous studies investigating imaginal wing disc regeneration have demonstrated the importance of inhibiting systemic Ec production to delay development and allow time for regeneration to occur. However, local roles for Ec signaling within regenerating tissues are not well understood. Based on our observation that Ec-responsive transcriptional reporters can be induced locally in injured larval wing discs, we have tested elements of the Ec pathway for roles in regeneration. We find that local knockdown of the Ec biosynthetic enzyme phantom (phm) during injury is sufficient to impair wing disc regeneration, and that local knockdown of the Ec-degradation enzyme cyp18a1 reciprocally enhances regeneration. In parallel, we find that injury induces a transient ring of Ec receptor (EcR) activity encircling the regenerating blastema which can be blocked by phm knockdown. In the absence of this EcR-activity pulse, we find a failure of intra-organ growth inhibition as cells within the uninjured compartments of the wing disc fail to slow their proliferation. Further, injured animals with phm knockdown lack a robust injury-induced developmental delay, while cyp18a1 knockdown has the opposite effect of extending injury-induced developmental delay. We will discuss mechanistic links between local Ec/EcR and signals produced by injured tissues that act systemically to control tissue homeostasis.

98 **Blocking the native differentiation program recapitulated in yki3S/A-induced midgut tumor alters the tumor cells’ capacity to disseminate and induce cachexia-like wasting** Inez Pranoto, Young Kwon University of Washington, Seattle WA

The developmental program generates heterogeneous cell types that form and maintain tissue structure and physiology. Many tumors have been shown to recapitulate the developmental program of the tissue of origin, thus generating heterogenous cells within the tumors. Although tumor cells with stem cell-like properties have been extensively studied, how developmental programs recapitulated in tumors or tumor cells undergoing differentiation contribute to the adverse phenotypes associated with malignant tumors, such as metastasis and cachexia, are only poorly understood. We employed *Drosophila* midgut tumors driven by expression of the active form of yorkie (yki3S/A) and revealed the striking heterogeneity of yki3S/A midgut tumor cells, which aberrantly reconstituted the differentiation program to generate enterocytes (ECs)– polyploid absorptive intestinal epithelial cells. Blocking the EC differentiation program by expressing dominant negative Notch (N1DM) significantly reduced the heterogeneity of yki3S/A tumor cells by eliminating the differentiating yki3S/A tumor cells, leading to formation of fully grown tumors comprising only intestinal stem cell-like yki3S/A tumor cells. Of note, blocking the EC differentiation program was sufficient to eliminate a population of yki3S/A tumor cells that can invade and migrate to disseminate from the midguts. Interestingly, eliminating yki3S/A tumor cells undergoing EC differentiation significantly altered the expression of multiple tumor-driven secreted factors and the tumor’s propensity for inducing the tumor non-autonomous and systemic phenotypes. In particular, blocking the EC differentiation program significantly decreased the expression of wasting factors in tumors and suppressed multiple wasting phenotypes, including ovary atrophy, fat body degeneration, and ‘bloating syndrome’. Our study demonstrates that the EC differentiation program recapitulated in yki3S/A midgut tumors is crucial for the tumor cells’
 capacity to disseminate from the midguts and induce certain wasting phenotypes. This work provides insights into how the contextual information in the tissue of origin can give rise to tumor’s ability to induce the phenotypes associated with malignant tumors. Thus, we propose that manipulating the developmental/differentiation programs recapitulated in tumors is a way to modulate tumor behavior for the treatment of the complications associated with the advanced cancers.

Enterocytes are a well conserved cell type and make up the majority of the digestive system across metazoans. In Drosophila, enterocytes comprise nearly 90% of the adult gut epithelium, and play important roles in digestion such as secretion of digestive enzymes and absorption of nutrients, maintaining gut epithelial barriers and regulating tissue turnover. Enterocytes are also the first line of defense against oral pathogens. However, despite their critical functions, enterocytes are not as well studied as the other cells in the midgut epithelium. Enterocytes are also subject to frequent cell turnover in the midgut epithelium, a process that is acutely accelerated upon infection. What determines the decision of an enterocyte to remain in the epithelium or to delaminate? To understand this, we have developed a number of genetic tools as well as microscopy and flow cytometry based assays to study epithelial dynamics and cell turnover in the midgut over a comprehensive time course following oral infection with multiple pathogens such as Erwinia carotovora 15 (Ecc15) and Pseudomonas entomophthila (Pe). We have identified key differences between the anterior and posterior midgut in the kinetics and amplitude of cell loss and enterocyte turnover upon infection in both male and female adult flies. We observe differences in enterocyte loss in response to infection with Ecc15 and Pe. At the tissue level, we have identified mechanical changes that the midgut epithelium undergoes in response to oral pathogens such as muscle constrictions, gut contraction and region-specific changes in length. Additionally, at the cellular level, we have identified acute and dynamic changes in enterocyte cell shape, size and the nature of cell-cell junctions. We are currently investigating the genetic factors responsible for these changes, and how they impact gut physiology and infection response in the adult fly. We are also employing live imaging strategies to monitor enterocyte dynamics in vivo under various conditions. Our work will provide several key insights into the biology of enterocytes under physiological as well as pathological conditions.

PAAC to the new normal: Intravital imaging of dynamic brush border repair in the adult Drosophila intestine Anthony Galenza1, Yu-Han Su1, Paola Moreno-Roman2, Irina Kolotuev2, Lucy O’Brien1 1) Stanford University, Stanford, CA; 2) Universite de Lausanne, Lausanne, Switzerland

Frequent exposure of the intestine to hostile pathogens and harsh toxins demands rapid and seamless repair. Because the intestine is located inside the body cavity, examination of the intestine’s physiological repair processes has traditionally been limited to fixed tissues. Here we use an intravital imaging methodology, Bellymount, to monitor real-time intestinal injury and subsequent repair in individual animals over multiple days. Within 24 hours of toxin feeding, components of the intestinal brush border, such as Moesin::GFP and Meduse::GFP, relocalize from lumenal cell surfaces to lateral cell surfaces; after 48 hours, they disappear entirely. Concomitant with loss of the intestinal brush border, Moesin::GFP and Meduse::GFP appear in nascent brush border membranes associated with new stem cell progeny. Unlike new brush border formation during normal intestinal renewal, which involves a strikingly large, Pre-Assembled Apical Compartment (PAAC) at tri-cellular junctions (Moreno-Roman et al., bioRxiv 457819), new brush border formation during repair involves linear clusters of 4-8 miniature PAACs at bi-cellular junctions. We call these clusters multi-PAACs. Either genetic ablation of the brush border in mature cells or activation of the injury response pathway JAK-STAT in stem cell progeny is sufficient to induce multi-PAACs in otherwise healthy guts. These findings delineate how damage signals trigger an injury-specific morphogenetic process. Together, these results reveal spatiotemporal dynamics of brush border damage and regeneration and provide insights into fundamental mechanisms of epithelial barrier repair.

Asymmetric nucleosome density and differential condensation of sister chromatids coordinates with Cdc6 to ensure distinct cell fates Rajesh Ranjan1, Xin Chen1,2 1) Johns Hopkins University; 2) Howard Hughes Medical Institute (HHMI)

Stem cells undergo asymmetric division to produce both a self-renewing stem cell and a differentiating daughter cell. During Drosophila male germline stem cell (GSC) asymmetric division, preexisting old histones H3 and H4 are enriched in the self-renewed stem daughter cell, whereas the newly synthesized H3 and H4 are enriched in the differentiating daughter cell. However, the biological consequences in the two daughter cells resulting from asymmetric histone inheritance remained to be elucidated. In this work, we track both old and new histones throughout GSC cell cycle using high spatial and temporal resolution microscopy. We find several unique features differentiating old versus new histone-enriched sister chromatids, including nucleosome density, chromosomal condensation, and H3 Ser10 phosphorylation. These distinct chromosomal features lead to their differential association with Cdc6, an essential component of the pre-replication complex, which subsequently contributes to asynchronous initiation of DNA replication in the two
resulting daughter cells. Disruption of asymmetric histone inheritance abolishes both differential Cdc6 association and asynchronous S-phase entry, demonstrating that asymmetric histone acts upstream of these critical events during cell cycle progression. Furthermore, GSC defects are detected under these conditions, indicating a connection between histone inheritance, cell cycle progression and cell fate decision. Together, these studies reveal that cell cycle remodeling as a crucial biological ‘readout’ of asymmetric histone inheritance, which precedes and could lead to other well-known readouts such as differential gene expression. This work also enhances our understanding of asymmetric histone inheritance and epigenetic regulation in other stem cells or asymmetrically dividing cells in multicellular organisms.

102 Rab35 mediates two distinct pathways that regulate actin modification through Mical/SelR and actin remodeling through Septins during cell wound repair Mitsutoshi Nakamura, Justin Hui, Tessa Allen, Susan M. Parkhurst Basic Sciences Division, Fred Hutchinson Cancer Research Center, Seattle, WA

Cell wound repair is essential to cope with daily wear and tear, especially when cells are fragile as a result of trauma or from disease conditions. The three main steps of wound repair are highly conserved among different organisms: rapid membrane resealing; dynamic cytoskeletal reorganization for wound closure; and cell cortex (membrane and underlying cytoskeleton) remodeling to restore the cell to its unwounded state. We recently found that Rab35 is required for the cell cortex remodeling phase following wound closure in the Drosophila cell wound repair model. Rab35, a member of Rab family GTPases, regulates membrane trafficking and cytoskeleton dynamics in many cellular processes such as cytokinesis. We find that Rab35 is recruited to wounds and its RNAi knockdown exhibits premature actomyosin ring disassembly. Interestingly, we find that Rab35 functions in this context through two different downstream pathways. One pathway involves Mical, an oxidation-reduction enzyme that binds and disassembles F-actin, and SelR, a methionine sulfoxide reduction enzyme that works as a counterpart to Mical. Both Mical and SelR are recruited to wounds and their RNAi knockdowns exhibit opposite phenotypes in actomyosin ring remodeling. The other pathway involves Septins, a family of GTP-binding proteins that form hetero-oligomeric structures at the cell cortex to regulate cytoskeleton. We find that all five Drosophila Septins are recruited to wounds with similar, but not identical, spatial and temporal patterns. Consistent with this, they exhibit different mutant phenotypes with respect to actin dynamics, suggesting that Septins have non-redundant and complex-independent functions during cell wound repair. We are currently investigating the coordination between these pathways needed for proper remodeling of the cell cortex upon wound closure.

103 Re-entry into mitosis and regeneration of intestinal stem cells through enteroblast dedifferentiation in Drosophila midguts Aiguo Tian1, Virginia Morejon1, Sarah Kohoutek1, Yi-Chun Huang1, Wu-Min Deng1, Jin Jiang2 1) Tulane University; 2) UT Southwestern Medical Center

Colorectal cancer (CRC) is the 3rd most common cancer diagnosed in the United States. Intestinal dysplasia is considered as a precursor lesion of CRC in mammals and can be mimicked in Drosophila. Dysplasia is characterized by hyperproliferation and accumulation of intestinal stem cells (ISCs) and enteroblasts (EBs). As a result of dysplasia, the lifespan is shortened. Current studies in Drosophila indicate that only resident ISC can divide and their over-proliferation results in intestinal dysplasia. In addition, enteroblasts (EBs) as one type of ISC progenies are accumulated under intestinal dysplasia. Thus, our studies use the Drosophila intestine as a model to understand how EBs contribute to dysplasia in response to infection of pathogenic bacteria. Here we find that infection of pathogenic bacteria induces enteroblasts as one type of ISC progenies to re-enter the mitotic cycle in the Drosophila intestine. The re-entry into mitosis is dependent on epithelial growth factor receptor (EGFR)-Ras signaling and ectopic activation of EGFR-Ras signaling in EBs is sufficient to drive EBs cell-autonomously to re-enter into mitosis. In addition, we examined whether EB gain ISC identity as a prerequisite to divide, but the immunostaining with stem cell marker Delta shows that these dividing EBs do not gain ISC identity. After employing lineage tracing experiments, we further demonstrate that EBs dedifferentiate to generate functional ISC after symmetric divisions of EBs. Together, our study in Drosophila intestines uncovers a new role of EGFR-Ras signaling in regulating re-entry into mitosis and dedifferentiation during regeneration and reveals a novel mechanism by which ISC progenies undergo dedifferentiation through mitotic division, which has important implication to increase of ISC pool and intestinal dysplasia.

104 Cellular and molecular basis of detection of acidic pH in fly gustatory system Anindya Ganguly1, Avinash Chandel1, Shan Wang1, Heather Turner2, Emily Liman2, Craig Montell1 1) University of California, Santa Barbara; 2) University of Southern California

Drosophila melanogaster use their sense of taste to forage for food, avoid toxic or hazardous substances, to select partners for courtship and mating and to choose suitable oviposition sites. Although the cellular and molecular basis of sweet, bitter and salt tastes have been well characterized in flies a detailed understanding of how they sense acid is lacking. Based on past studies, members of Gustatory Receptor (GR) or Ionotrophic Receptor (IR) families are unlikely to function broadly as acid receptors. Hence, we elected to test whether fly genes that are distantly related to mammalian Otop1, a proton channel required for sour taste in mammals, are required for acid taste in flies. RNAi-mediated silencing as well as CRISPR/Cas9-mediated mutagenesis of OtopLA, but not of OtopLB or of OtopLC caused a severe reduction in behavioral aversion to sweet solutions in the presence of low pH or high concentrations of carboxylic acids as well as
action potentials induced by acids on neurons in labellar taste bristles. We identified a novel OtopLA isoform from the proboscis (OtopLAp) that rescued the behavioral deficit as well as the reduced neuronal sensitivity to acids exhibited by the OtopLA mutant. Although high concentrations of HCl caused behavioral aversion, we observed mild attraction to low concentrations of HCl. This attraction was also impaired in OtopLA mutant. By conducting cell-type specific genetic rescue experiments in various subsets of taste neurons, we demonstrated that OtopLA functions in different neuronal subsets to cause different behavioral outcomes. This study demonstrates functional conservation of a taste receptor between flies and mammals. It also elucidates that the same taste receptor is required both for appetitive and repulsive gustatory behaviors.

105 Developmental mechanisms regulating the formation and function of drosophila sleep-wake circuit Adil R Wani1, Aisha Hamid1, Hannah Deutsch 2, Gonzalo M Chaya1, Matthew S Kayser1, Mubarak H Syed1 1) University of New Mexico, Albuquerque, NM; 2) University of Pennsylvania, PA

Molecular mechanisms regulating the formation and function of neural cell types are not fully understood. Studying the programs by which neuronal types are specified, connected to form unique functional circuits regulating discrete behaviors is an important area of neuroscience. We are investigating this long-standing question using the recently identified and conserved Drosophila sleep-wake circuit. We utilize novel lineage filtering tools to show that sleep-promoting dorsal fan-shaped body (dFB) neurons are born from late-type II neural stem cells (NSCs) that express ecdysone induced protein (E93). Using mosaic analysis, we show that majority of the dFB neurons are generated from the dorsolateral 1 (DL1) type II NSCs, and E93 is essential for the specification of these neurons. Interestingly, the knockdown of E93 in type II NSCs affects adult sleep behavior. Our data show that we have identified steroid hormone-mediated NSC-specific developmental programs that regulate the formation and function of the sleep-wake circuit.

106 Associative learning drives longitudinally-graded presynaptic plasticity of neurotransmitter release along axonal compartments Aaron Stahl1, Nathaniel Noyes1, Tamara Boto1, Miao Jing1, Jianzhi Zeng4,5,6, Lanikea King1, Yulong Li1,4,5,6, Ronald Davis1, Seth Tomchik1 1) The Scripps Research Institute, Jupiter, FL; 2) Trinity College, Dublin, Ireland; 3) Chinese Institute for Brain Research, Beijing, China; 4) State Key Laboratory of Membrane Biology, Peking University School of Life Sciences, Beijing, China; 5) Peking-Tsinghua Center for Life Sciences, Beijing, China; 6) PKU- IDG/McGovern Institute for Brain Research, Beijing, China

Anatomical and physiological compartmentalization of neurons is a mechanism to increase the computational capacity of a circuit, and a major question is what role axonal compartmentalization plays. Axonal compartmentalization may enable localized, presynaptic plasticity to alter neuronal output in a flexible, experience-dependent manner. Here we show that olfactory learning generates compartmentalized, bidirectional plasticity of acetylcholine release that varies across the longitudinal compartments of Drosophila mushroom body (MB) axons. The directionality of the learning-induced plasticity depends on the valence of the learning event (aversive vs. appetitive), varies linearly across proximal to distal compartments following appetitive conditioning, and correlates with learning-induced changes in downstream mushroom body output neurons (MBONs) that modulate behavioral action selection. Potentiation of acetylcholine release was dependent on the Ca2.1 calcium channel subunit cacophony. In addition, contrast between the positive conditioned stimulus and other odors required the inositol trisphosphate receptor (IP3R), which was required to maintain responsivity to odors in untrained conditions. These data demonstrate that learning drives valence-correlated, compartmentalized, bidirectional potentiation and depression of synaptic neurotransmitter release, which rely on distinct mechanisms and are distributed across axonal compartments in a learning circuit.

107 A conserved RNA binding protein regulates RNAs critical for neurodevelopment Carly Lancaster, Ken Moberg, Anita Corbett Emory University

Inherited forms of intellectual disability (ID) are common in the general population and have been linked to lesions in >700 genes. Emerging evidence suggests that this diverse group of genes converge on a limited set of neurodevelopmental pathways, including those that rely on RNA binding proteins (RBPs) to guide spatiotemporal patterns of neuronal mRNA trafficking and translation. Our labs co-discovered a monogenic form of ID caused by loss-of-function mutations in the ubiquitously expressed RBP ZC3H14. Studies exploiting the conserved ZC3H14 ortholog in Drosophila, Nab2, reveal that Nab2 localizes to neuronal nuclei and cytoplasmic ribonucleoprotein granules and is required specifically within brain neurons for olfactory memory and proper patterns of axon projection. At a molecular level, Nab2 can act as a translational repressor in conjunction with the Fragile-X mental retardation protein homolog Fmr1 and shares target RNAs with the Fmr1-interacting RBP Ataxin-2. However, neuronal signaling pathways regulated by Nab2, as well as mechanisms that elevate ZC3H14/Nab2 function in neurons relative to other cell types, remain elusive. We will present evidence that Nab2 controls neuronal expression of a well-conserved growth cone guidance factor, the guanine-nucleotide exchange factor (GEF) Trio, whose vertebrate homolog TRIO acts through the F-actin regulatory GT-Pases RHO and RAC to guide axon projection. Nab2 controls Trio levels in the fly brain by modulating an intron-retention event within the 5’ UTR of trio pre-mRNA. Data will be presented on the role of m6A and Nab2 in
controlling Trio splicing and expression, along with Nab2-Trio coregulation of axonal development in the CNS. Given that human TRIO is mutated in a dominant form of ID, this potential link between Nab2 and Trio in Drosophila could suggest that Nab2/ZC3H14 and Trio/TRIO act in a conserved ID pathway required to pattern neuronal processes in the developing nervous system.

108 Recovery from cold-induced reproductive dormancy is regulated by temperature-dependent AstC signaling Matthew R. Meiselman1, Michael H. Alpert1, Xinyue Cui2, Jamien Shea1, Ian Gregg1, Marco Gallio2, Nilay Yapici1 1) Department of Neurobiology and Behavior, Cornell University, Ithaca, NY, NY, USA; 2) Department of Neurobiology, Northwestern University, Evanston, IL, 60208, USA

Many species have evolved a variety of behaviors to cope with adverse environmental conditions. Similar to many insects, Drosophila melanogaster responds to sustained cold temperatures by reducing locomotion, metabolic rate, and arresting reproduction. Here, we show that a subset of dorsal neurons (DN3s) that express the neuropeptide Allatostatin-C (AstC) in the fly brain facilitates recovery from cold-induced reproductive dormancy. Activity of AstC-expressing DN3s is modulated by temperature and AstC peptide levels are suppressed at cold temperatures in Drosophila and in vector mosquitos. The stimulatory effect of AstC on egg production during and after cold-induced dormancy is mediated by AstC-R2-expressing CCHA2 neurons, whose activation is sufficient to induce reproductive dormancy in warm temperatures. Our results demonstrate that DN3s coordinate female reproductive capacity with environmental temperature via AstC signaling. AstC/AstC-R2 is conserved across many insect species and their role in regulating female reproductive capacity makes them an ideal target for controlling the population of agricultural pests and human disease vectors.

109 Orion bridges phosphatidylserine and Draper in the phagocytosis of somatosensory neurons in Drosophila Hui Ji1, Bei Wang1, David Labib1,2, Joyce Lei1,4, Xinchen Chen1, Maria Sapat1,3, Ana Boulanger2, Jean-Maurice Dura2, Chun Han1 1) Well Institute for Cell and Molecular Biology and Department of Molecular Biology and Genetics, Cornell University, Ithaca, NY; 2) IGH, Centre National de la Recherche Scientifique, Université de Montpellier, Montpellier, France; 3) New York Stem Cell Foundation, New York, NY; 4) Tisch MS Research Center of New York, New York, NY

Phagocytic clearance of degenerating neurites is critical for the prevention of neuroinflammation and neuronal regeneration. Phosphatidylserine (PS), an ‘eat-me’ signal exposed on degenerating neurites, and Draper (Drpr), an engulfment receptor expressed by phagocytes, are known to mediate the phagocytosis of degenerating neurites in Drosophila. However, how PS is sensed by Drpr-expressing phagocytes remains poorly understood. A novel secreted protein, Orion, was recently found to be required for axon pruning of mushroom body neurons during metamorphosis. In this study, we investigated whether Orion functions as a bridging molecule between PS and Drpr using larval dendritic arborization (da) neurons and phagocytic epidermal cells as a model of neuronal phagocytosis. First, using multiple models of dendrite degeneration, we demonstrate that Orion is required for Drpr-mediated phagocytosis of dendrite debris. Orion loss-of-function (LOF) did not interfere with PS exposure on dendrites but abolished recruitment of epidermal Drpr to degenerating dendrites, suggesting that Orion acts downstream of neuronal PS exposure and upstream of Drpr. Second, we found that Orion is secreted by many non-neural tissues and shows PS-dependent binding to degenerating dendrites. Orion outcompetes Annexin V in binding to injured dendrites, suggesting a direct interaction with PS. Third, we found that Orion functionally interacts with Drpr: Orion accumulates on epidermal cells that overexpressed Drpr. Importantly, a membrane-tethered Orion induced PS-independent and Drpr-dependent dendrite degeneration when expressed in neurons but blocked engulfment of dendrites when expressed in epidermal cells. Lastly, we found that the Orion dosage modulates the sensitivity of the epidermal cell to PS: While Orion heterozygosity reduced epidermal engulfment of PS-exposing dendrites, Orion overexpression caused engulfment-dependent, ectopic degeneration of wildtype dendrites. Together, these findings argue that Orion functions as a bridging molecule between PS and Drpr and fine-tunes the competence of phagocytes in neuronal engulfment. The shared features between Orion and human chemokines that were recently demonstrated to function as “find-me” signals by binding to PS-exposing vesicles suggest conserved mechanisms between insects and humans in phagocytosis and homeostatic regulations of diverse body systems.

110 The Circular RNA Edis-Relish-Castor Axis Regulates Neurodevelopment Wei Liu1, Weihong Liang1, Xiao-Peng Xiong2, Jian-Liang Li1, Rui Zhou1 1) Depts. of Medicine, Biological Chemistry & Oncology, Johns Hopkins University School of Medicine, Baltimore, MD; Johns Hopkins All Children’s Hospital, St. Petersburg, FL; 2) Sanford Burnham Prebys Medical Discovery Institute, La Jolla, CA; 3) NIEHS, Durham, NC

We have identified and validated a collection of circular RNAs in Drosophila melanogaster in response to immune challenge. We show that depletion of the brain-enriched circular RNA Edis (Ect4-derived immune suppressor), but not its linear sibling Ect4, causes hyperactivation of antibacterial innate immunity signaling both in cultured cells and in vivo. Conversely, ectopic expression of Edis blocks innate immunity signaling. Our analyses reveal that Edis is enriched in neurons, and that neuron-specific depletion of Edis leads to defective axon guidance, impaired locomotive activity and shortened lifespan. Importantly, restoration of Edis expression suppresses the innate immunity hyperactivation
and neurodevelopment phenotypes elicited by Edis depletion. Mechanistically, Edis encodes a functional protein that associates with and blocks the processing/activation of the innate immunity transcription factor Relish. We further show that the Castor gene, which encodes a transcription factor involved in neurodevelopment, is upregulated in Edis knockdown neurons. Notably, Castor overexpression phenocopies Edis overexpression in neurons, and reducing Castor levels suppresses the neurodevelopmental phenotypes in Edis knockdown neurons. Chromatin immunoprecipitation assay reveals that Relish associates with the promoter of Castor. In addition, we show that Relish-binding sites are required for optimal Castor promoter activity induced by either peptidoglycan treatment or Edis depletion. Lastly, Relish mutation rescues both Castor hyperactivation and neurodevelopmental phenotypes in Edis knockdown neurons. We conclude that the circular RNA Edis serves as a key modulator of innate immunity, and that the Edis-Relish-Castor axis regulates neurodevelopment.

111 Netrins and receptors control Drosophila optic lobe organization and transmedullary neuron axon targeting Yu Zhang, Scott Lowe, Xin Li The School of Molecular and Cellular Biology

During development, integration of temporal patterning and spatial patterning of neural progenitors as well as Notch-dependent binary fate choice between sister neurons contribute to generation of neural diversity. How these upstream neural fate specification programs regulate downstream effector genes to control axon targeting and neuropil assembly remains less well-understood. Here we show that Notch-dependent binary fate choice in Drosophila medulla neurons regulates the expression of Netrin, and that Netrin pathway controls axon guidance of transmedullary (Tm) neurons and contributes to the organization of optic lobe neuropils. Netrins are enriched in the lobula where Tm axons target, and the attractive receptor Frazzled is expressed broadly in medulla neurons, while the repulsive receptor Unc-5 is excluded from Tm neurons and this is necessary for their correct targeting to the lobula. Frazzled is required collectively in a group of early-born Tm neurons for their axons to establish the inner optic chiasm (IOC). In addition, Frazzled acts in the layer-specific targeting step of Tm3 and Tm4 cell-autonomously, and is also required for the formation of the lobula branch of TmY3. Moreover, we show that the diffusibility of Netrins is necessary for Netrin enrichment in the lobula, the IOC formation and layer-specific targeting of Tm3 and Tm4. Netrin enrichment in the lobula is promoted by Frazzled expressed in Tm neurons, while Unc-5 appears to have an opposite role in Netrin distribution. Furthermore, we show that Netrin B is expressed in the Notch-on hemilineage of medulla neurons including most Tm and TmY neurons that target lobula, and loss of Su(H) abolished NetB expression in the medulla. Without medulla-originated NetB, Tm axons from late-born medulla columns cannot join the IOC. Therefore, the Notch-dependent binary fate choice regulates the assembly of the optic lobe neuropils by controlling the expression of Netrin.

112 Functional divergence of the bag of marbles gene in the Drosophila melanogaster species group Jaclyn Bubnell, Cynthia Ulbing, Paula Fernandez Begne, Charles Aquadro Cornell University

In Drosophila melanogaster, a key germline stem cell (GSC) differentiation factor, bag of marbles (bam) shows rapid bursts of amino acid fixations between sibling species D. melanogaster and D. simulans, but not in the outgroup species D. ananassae. We previously hypothesized that a change in function and/or a genetic conflict with the intracellular bacteria W. pipientis could be driving the adaptive evolution of bam, however bam function has only been defined in D. melanogaster. Here, we test the null hypothesis that bam’s differentiation function is conserved in four additional Drosophila species in the melanogaster species group spanning approximately 15 million years of divergence. Surprisingly, we demonstrate that bam is not necessary for oogenesis or spermatogenesis in D. teissieri nor is bam necessary for spermatogenesis in D. ananassae. Remarkably bam function may change on a relatively short time scale. We further report tests of neutral sequence evolution at bam in additional species of Drosophila and find a positive correlation between evidence for positive selection at bam and its essential role in GSC regulation and fertility for both males and females. Further characterization of bam function in more divergent lineages will be necessary to distinguish between bam’s critical gametogenesis role being newly derived in D. melanogaster, D. simulans and D. yakuba or it being basal to the genus and subsequently lost in numerous lineages.

113 Cross-species incompatibility between a DNA satellite and the Drosophila homolog of Spartan poisons germline genome integrity Cara Brand, Mia Levine University of Pennsylvania

Satellite DNA spans megabases of eukaryotic genome sequence. These vast stretches of tandem DNA repeats undergo high rates of sequence turnover, resulting in radically different satellite DNA landscapes between closely related species. Such extreme evolutionary plasticity suggests that satellite DNA accumulates mutations with no functional consequence. Paradoxically, satellite-rich genomic regions support essential, conserved nuclear processes, including chromosome segregation, dosage compensation, and nuclear structure. A leading resolution to this paradox is that deleterious proliferation of satellite DNA variants trigger adaptive evolution of satellite-associated chromosomal proteins to preserve these essential functions. Here we experimentally test this model of intra-genomic coevolution between chromosomal proteins and DNA satellites by conducting an evolution-guided manipulation of both. The 359bp satellite spans an 11Mb array in D. melanogaster that is absent from its sister species, D. simulans. This species-specific satellite array colocalizes with the Drosophila protein, Maternal Haploid (MH), an essential, adaptively evolving member of the
Spartan family of proteases that cleave DNA-protein crosslinks. To determine if MH and 359 coevolve, we swapped the *D. simulans* version of MH (“MH[sim]”) into *D. melanogaster*. We discovered that MH[sim] triggers elevated ovarian cell death, reduced ovary size, and loss of mature eggs. In contrast, mh null females have no such ovary phenotypes, suggesting that MH[sim] is toxic. Using both cell biological and genetic approaches, we demonstrate that MH[sim] poisons oogenesis through a DNA damage pathway. Remarkably, deleting the *D. melanogaster*-specific 359 satellite array from mh[sim] females completely restores female germline genome integrity and fertility, consistent with a history of coevolution between these two fast-evolving components of the *Drosophila* genome. Genome integrity and female fertility are also restored by overexpressing Topoisomerase II (Top2). Top2 resolves DNA entanglements by generating transient double stranded breaks while crosslinked to DNA. Intriguingly, these DNA-Top2 crosslinks are resolved by the MH homolog in worms and humans. We propose that MH[sim] prematurely cleaves DNA-Top2 crosslinks that form during Top2-mediated processing of 359 entanglements. Under this model, MH evolved adaptively along the *D. melanogaster* lineage to avoid 359 in the ovary, leading to the observed cross-species incompatibility between the 359 satellite and MH. Our study offers rare experimental evidence that intra-genomic coevolution between ostensibly inert repetitive DNA and essential chromatin proteins preserves germline genome integrity.

### 114 A putative de novo evolved gene is essential for male fertility via a paternal effect

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*De novo* evolved genes arise from previously non-coding DNA regions and are found in only one or a few species. In *D. melanogaster*, many *de novo* genes are expressed in the testis, suggesting they may affect male fertility and be subjected to sexual selection. An RNAi and CRISPR knockout screen identified *katherine johnson* (*kj*) as one of several putative *de novo* genes essential for male fertility. Males knocked down or knocked out for *kj* show a 90% reduction in fertility. Curiously, dissection of mutant testes revealed high numbers of mature sperm in the seminal vesicle. Furthermore, these sperm are transferred to females and enter the seminal receptacle at rates equivalent to sperm of controls. Instead, measurements of egg-to-adult viability showed that the loss of *kj* causes a fertility defect because of a failure of embryos fathered by *kj* null males to develop into larvae. Preliminary evidence indicates that embryos fathered by *kj* nulls initiate mitotic divisions, confirming that *kj* null male sperm are fertilization competent. However, by 6 hours after fertilization, embryos of *kj* null males show a significantly reduced rate of developmental progression. To understand the normal function of the *kj* protein, and how its loss causes this developmental defect, we constructed a transgenic fly line expressing a 6xHis-tagged *kj* transgene under the control of its own regulatory sequences. We are currently examining *KJ-HA* localization and further evaluating the exact stage of embryonic development at which *kj* null-fathered embryos begin to show defects. While several other putative *de novo* genes are required for successful sperm production, *kj* represents the first such gene to our knowledge to be required for the proper use of the paternal genome after fertilization. These findings further establish the ability of newly evolved genes to evolve essential roles in reproduction.

### 115 The Y-linked gene, WDY, is necessary for sperm storage in *Drosophila melanogaster*.  

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Only 13 single-copy protein-coding genes are known on the *Drosophila melanogaster* Y-chromosome -- a chromosome that is 40 Mb, haploid, paternally inherited, entirely heterochromatic, repeat-dense, and lacks recombination. These genes display remarkable “functional coherence,” with five Y-linked genes (three of which encode dynein motor components) being necessary to produce mature sperm past the early individualization stage of spermatogenesis. The significance of placing these fertility-essential genes in a heterochromatic and degenerating region of the genome is not understood. We are now discovering additional, more subtle functions associated with the Y chromosome. We and others recently demonstrated by knockdown that the gene, *WDY*, is responsible for the sterility of the *kl-1* region of the Y chromosome. Here we further investigate the phenotype of *WDY* using CRISPR mutants we generated. Using compound sex chromosomes, we established several stable mutant lines for *WDY*. Mutant males were sterile, as expected, but surprisingly produced mature, motile sperm. *WDY* mutant sperm were transferred to females, albeit at lower rates. Consistent with prior reports associated with the *kl-1* region, we found that *WDY* mutant sperm are unable to enter storage in the female. This failure may be caused by a motility defect of the sperm or an inability to interact with female molecules that enable storage. Further analysis of *WDY* may reveal mechanisms by which sperm are stored, as well as clues to the underlying evolutionary logic of locating a gene on the Y chromosome.

### 116 Investigating the evolution of new body parts in the rapidly evolution genitalia of *Drosophila*  

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Recently evolved traits i.e., novelties, often represent key features that allow animals to exploit new ecological niches (e.g. feathers in birds) and can even help them find a mate (e.g. bioluminescence in fireflies). The rapidly evolving genitalia of Drosophila provides a powerful system to study the developmental basis of qualitative changes in morphology. However, this high morphological diversity also poses a distinct challenge, since it can be difficult to disentangle which structures are homologous and which represent novelties.

To determine whether genital structures of different species were homologous or novel, we compared the development of phallic outgrowths in the Oriental lineage, which contains Drosophila melanogaster. We found that most phallic outgrowths are formed by multicellular projections. However, several multicellular outgrowths that were thought to be homologous, are in fact formed by different tissues of the phallus in different species, indicating that they are heterologous structures. Furthermore, we uncovered evidence that phallic outgrowths found in the Drosophila eugracilis phallus are formed by large unicellular projections, suggesting that they have evolved convergently to the previously described multicellular outgrowths. These unicellular outgrowths are likely novel as they are absent the homologous tissue of five other species of the Oriental lineage. We found that the trichome genetic network is expressed in these unicellular phallic outgrowths, suggesting that the co-option of this network underlies this dramatic phenotype. In fact, activation of the trichome genetic network in the phallus of Drosophila melanogaster induces a phenocopy of the unicellular outgrowths found in Drosophila eugracilis.

Our work highlights that understanding development can be key to discern between homology vs heterology, allowing us to formulate testable evolutionary models (i.e. co-option vs convergence).

**117 Do supergenes mediate seasonal adaptation in overwintering Drosophila?** Joaquin Nunez1, Alyssa Bangerter1, Connor Murray1, Yang Yu1, Benedict Lenhart1, Priscilla Erickson1,2, Alan Bergland1 1) University of Virginia, Charlottesville, VA; 2) University of Richmond, Richmond, VA

Vinegar flies (Drosophila melanogaster) living in temperate regions experience strong fluctuations in the strength and direction of selection due to seasonality and evolve rapidly to track the changing fitness landscape. Recent work has shown the existence of genomic loci whose seasonal signal is replicated across continents. Despite these advances in identifying the presence of fluctuating selection and adaptation between seasons, we still lack a detailed understanding of the functional genetic architecture of seasonal adaptation. This gap in knowledge is further compounded by the fact that wild Drosophila populations experience demographic “boom-and-bust” cycles coconmitant with seasonality, a process which may alter levels of standing genetic variation. We characterized both the genomic consequences of boom-and-bust cycles as well as the targets of seasonally varying selection. We accomplished this by leveraging temperature data combined with genomic sequences from densely sampled orchard populations collected within (June-November) and across growing seasons over multiple populations on two continents. Principal component analysis reveals temporal structure, and using simulation, we show that this temporal structure is consistent with cyclical winter population collapse. This temporal signal is observed genome-wide except for chromosome 2L, where allele frequencies of the cosmopolitan inversion Inv(2L)t drive genetic structure. We further investigated these patterns at Inv(2L)t by regressing the mean temperature, 30 days prior to sample collection, onto allele frequencies of each SNP using a generalized linear model. This analysis reveals an enrichment of loci responding to temperature clustered within Inv(2L)t. Linkage analysis suggest that the loci associated with temperature changes lie within the inversion but is not the inversion per se. Additional analyses show multiple mutations in high linkage at distances ~0.1 – 0.5 Mb, suggesting long range associations among seasonal loci inside the inversion. Enrichment analysis show an excess of non-synonymous mutations among the Inv(2L)t seasonal outliers. Overall, our data provides two insights about Drosophila seasonality. First, we show that severe demographic contractions and expansions are an important, and measurable, component of the seasonal dynamics of temperate Drosophilids. Second, our data supports the general hypothesis that supergenes are important drivers of adaptation in highly fluctuating environments.

**118 The evolution and genetic mechanism of sex-ratio meiotic drive in Drosophila affinis** Wen-Juan Ma, Robert L. Unckless University of Kansas

Meiotic drivers are selfish genetic elements that promote their own transmission to the gametes. Sex-ratio (SR) meiotic drive occurs when a selfish genetic element on the X chromosome manipulates gametogenesis to prevent the maturation of Y-bearing sperm in males, resulting in the production of predominant female progeny. The spread of SR meiotic drive can affect host genetic diversity, sex chromosome evolution, and even cause host extinction if it reaches high enough frequency. SR meiotic drivers have evolved independently several times, however, the underlying genetic mechanism is only known in few cases. In this study we use a combination of genomics, transcriptomics, single nuclei RNA-seq (snRNA-seq), and CRISPR-Cas9 to identify the genetic mechanism responsible for the SR meiotic drive in Drosophila affinis. We found X chromosome is enriched for differentially expressed (DE) transcripts and that many DE genes had elevated Ka/Ks values between SR and ST. We identified top DE candidate genes, including two X-linked
dissimilar and testis-specific: encoding a chromosomal protein involved in spermatogenesis (thmg2), and encoding the regulator of chromatin condensation (rcc1). we used crispr-cas9 knockout experiments to test whether these genes were involved in the sex ratio phenotype. lastly, snrna-seq analysis recapitulates transcriptional patterns associated with gene expression changes during spermatogenesis between st and sr testes. our results suggest the two candidate genes’ significant role involving in the drive in this species, providing evidence that rapid evolution of genes disrupting spermatogenesis is important source of intragenomic conflict.

119 the genetic basis of cardiac glycoside resistance in wild-caught Drosophila melanogaster Arya Rao, Peter Andolfatto Columbia University

A significant number of unrelated herbivorous insects have repeatedly and independently evolved the ability to feed on plants that produce toxic secondary compounds called cardiac glycosides (CGs). CGs inhibit the function of Na+,K+-ATPase (NKA), a medically important enzyme in animals that is necessary for many processes including neural function and muscle contraction. Previous work identified a small number of adaptive amino acid substitutions in the alpha-subunit (ATPα) of NKA that confer the enzyme with resistance to CG inhibition. Further, engineering the native ATPα of D. melanogaster to carry one or more of these adaptive substitutions results in flies that are substantially resistant to CG toxicity. We show that wild-collected D. melanogaster strains harbor substantial genetic variation in sensitivity to CG-toxicity. By conducting GWAS on 180 Drosophila melanogaster Genome Reference Panel lines, we show that this CG resistance does not map to ATPα, but maps to a number of genes implicated in maintaining physiological barriers to solute diffusion and cellular and behavioral stress response. This information can be used to predict additional targets of recurrent adaptation in diverse CG-tolerant taxa. In addition, these results yield insight into the development of drugs to treat a number of Na+,K+-ATPase-associated neurological and physiological disorders in humans.

120 old hormones, new tricks: juvenile hormones ensure primordial germ cells reach the embryonic somatic gonad Lacy Barton1, Justina Sanny1, Emily P Dawson1, Rebecca Spokony2, Ruth Lehmann3 1) New York University, Skirball Institute; 2) Baruch College, CUNY; 3) The Whitehead Institute, MIT

Proper germ cell development is crucial for the health and fertility of subsequent generations. Emerging evidence suggests that secreted isoprenoids, such as retinoic acids (RAs) in vertebrates and juvenile hormones (JHs) in insects, impact several aspects of fertility. However, the mechanisms by which these potent molecules support reproductive development remain unclear. Here, we sought to elucidate the functions of JHs in primordial germ cell development. We made a GFP-based JH reporter, which revealed that JH signaling is first active in the Drosophila embryonic mesoderm as primordial germ cells emerge from the endoderm and begin to migrate within the mesoderm toward the somatic gonad. Consistent with this germ cell-proximal signaling pattern and previous work (Niwa, et al., 2008), we find that expression of a gene encoding the key JH biosynthesis enzyme, jhamt, is enriched near germ cells as they migrate toward the gonad and that JH degradation enzymes are expressed in areas that germ cells migrate away from. By generating several new double and triple mutant animals, we find that both JH biosynthesis and degradation enzymes are required for germ cells to efficiently colonize the developing somatic gonad. Together, these data suggest that JHs, which act systemically in larval and adult animals, act locally in the Drosophila embryo and that local dynamics in JH bioavailability may be needed to facilitate germ cell migration. To isolate autonomous vs non-autonomous requirements, we turned to an in vitro migration assay using primordial germ cells isolated by FACS, finding that pure JH III or the JH mimic, methoprene, is sufficient for germ cell migration in vitro. Interestingly, results from several experiments suggest that this newly uncovered function for JHs does not involve classical, nuclear receptor-mediated transcription, but rather calcium-linked cytosolic factor, phospholipase C. Collectively, these findings add to the growing appreciation of the diverse roles of secreted steroid and isoprenoids throughout the germline lifecycle in species that span both invertebrates and vertebrates. In mice, enzymes that synthesize RAs are also expressed in the developing somatic gonad as germ cell colonize this tissue. We find that, like JH in Drosophila, RA is sufficient for mouse germ cell migration in vitro, leaving open the possibility that this newly uncovered role in reproductive development may be conserved.

121 the role of Drosophila germ granules in regulating mRNA stability during germ cell development Anna Hakes, Elizabeth Gavis Princeton University

Germ granules, membraneless organelles containing mRNAs and proteins required for germline development, are a characteristic feature of germ cells. In Drosophila, germ granules form at the posterior of the oocyte and are subsequently segregated to the germ cell progenitors, called the pole cells, which bud from the posterior of the embryo. After pole cell formation, the germ granules increase in size and persist at least until the pole cells coalesce in the gonad. The persistence and evolutionary conservation of germ granules suggest that they play an important role in RNA regulation during germ cell development, but this role is not fully understood. We found that components of the mRNA decay pathway are sequentially recruited to the germ granules as these granules begin to increase in size. This association coincides with a dramatic decrease in the levels of two granule mRNAs, nos and pgo. By contrast, another germ granule mRNA, cycB, is protected from this degradation throughout embryogenesis. Our findings thus suggest that after pole cell formation, the germ granules play a dual role, selectively protecting some mRNAs while promoting...
the degradation of others, which is distinct from their protective role prior to pole cell formation. Additionally, we have found that the recruitment of mRNA decay proteins to the germ cells is dependent on translation. Together, these data indicate that there is a temporally regulated change in germ granule function in the pole cells, suggesting that the germ granules evolve throughout development to meet the needs of the germ cells. Experiments are in progress to determine what factor is translationally activated to trigger this shift in germ granule function. Since germ granules are one of many types of RNA-rich membraneless organelles, understanding their function in regulating RNA stability can provide general insights into the roles of membraneless organelles in post-transcriptional regulation.

122 A feedback loop between heterochromatin and the nucleopore complex controls germ-cell to oocyte transition during Drosophila oogenesis Kahini Sarkar1, Noor M Kotb2, Alex Lemus1, Elliott T Martin3, Alicia McCarthy3, Justin Camacho3, Ayman Iqbal3, Alex M Valm3, Morgan A Sammons1, Prashanth Rangan1 1) Department of Biological Sciences/RNA Institute, University at Albany SUNY, Albany, NY; 2) Department of Biomedical Sciences, The School of Public Health, University at Albany SUNY, Albany, NY; 3) Current address: 10x Genomics Headquarters, 6230 Stoneridge Mall Rd, Pleasanton, CA

Germ cells differentiate into oocytes that become totipotent upon fertilization. How the highly specialized oocyte acquires this distinct cell fate is poorly understood. During Drosophila oogenesis, H3K9me3 histone methyltransferase SETDB1 translocates from the cytoplasm to the nucleus of germ cells concurrent with oocyte specification. Here, we discovered that nuclear SETDB1 is required to silence a cohort of differentiation-promoting genes by mediating their heterochromatinization. Intriguingly, SETDB1 is also required for the upregulation of 18 of the ~30 nucleoporins (Nups) that comprise the nucleopore complex (NPC). NPCs in turn anchor SETDB1-dependent heterochromatin at the nuclear periphery to maintain H3K9me3 and gene silencing in the egg chambers. Aberrant gene expression due to loss of SETDB1 or Nups results in loss of oocyte identity, cell death and sterility. Thus, a feedback loop between heterochromatin and NPCs promotes transcriptional reprogramming at the onset of oocyte specification that is critical to establish oocyte identity.

123 Single-cell testes expression of ampliconic meiotic drivers on the sex chromosomes of Drosophila miranda Kevin Wei, Kamalakar Chatla, Doris Bachtrog University of California Berkeley

Sex-ratio meiotic drivers act during spermatogenesis to cause unequal production of X- and Y-bearing sperm. The presence of a driver on one chromosome reduces the fitness of the other and puts selection pressure for a suppressor to emerge counteracting the sex-ratio imbalance. This conflicting dynamic of X- and Y-linked drivers and suppressors is thought to underlie copy number amplifications of meiotic genes on sex chromosomes which has been observed all across animals. The recently formed sex chromosomes of Drosophila miranda is an extreme case; on both the neo-X and neo-Y chromosomes, hundreds of previously autosomal genes with meiotic function have massively amplified within ~1.5 million years. This system provides the opportunity to elucidate how these gene amplons manipulate spermatogenesis to confer transmission advantage. To this end, we applied single-cell RNA-seq on D. miranda testes to understand their spatiotemporal regulation. 10,095 cells were recovered and placed into 14 clusters corresponding to different testis cell types including sperm cells at varying stages of spermatogenesis. Of the 128 multi-copy gene families, the majority are most highly expressed in either the spermatocytes (n = 74), where meiosis occurs, or the premeiotic spermatogonia (n = 28). Interestingly, the expression of ampliconic X-linked genes, on average, peaks in early spermatocytes, while ampliconic Y-linked genes lags behind, peaking in late spermatocytes and lasting through sperm individualization. Because X-linked drivers can be suppressed via complementary small RNAs, we further generated D. miranda testes small RNA-seq and found abundant production of 21bp small RNAs from the ampliconic genes strongly implicating down regulation via the small interfering RNA silencing pathway. Moreover, genes that are co-amplified on both sex chromosomes and have pre-meiotic expression show disproportionally high siRNA production from the Y-linked ampliconic gametologs. These results suggest that Y-linked ampliconic genes may act to downregulate their X-linked counterparts, and maintenance of equal sex ratio likely depends on the premeiotic dosage balance of X-linked expression and Y-linked siRNA production.

124 Ecdysone signaling times border cell migration by regulating protrusive activity and cell-cell adhesion Mallika Bhattacharya, Michelle Starz-Gaiano University of Maryland, Baltimore County

Migratory cells play a significant role in spatiotemporally-regulated physiological processes such as normal embryonic development and wound healing. Dysregulation of these cells has severe implications in birth defects and diseases such as cancer. Border cell migration, during Drosophila oogenesis, serves as a useful model to study collective cell migration in vivo. Derived from the anterior epithelium of the egg chamber, the border cell cluster migrates toward the posteriorly-located oocyte in a temporally regulated manner. Developmental timing in flies is largely controlled by the sole steroid hormone ecdysone, which binds to a heterodimeric receptor complex that regulates transcription. Our work focuses on understanding how signaling by the ecdysone receptor (EcR), a conserved nuclear hormone receptor, affects border cell migration. Downregulation of ecdysone signaling results in delayed migration as a result of late detachment from the epithelium and slower migration. The directional translocation of the cluster, in response to chemotactants,
requires the extension of actin-rich protrusions that adhere to other cells in the migratory path. Live imaging video analysis revealed that clusters expressing transcriptionally-inactive EcR extend fewer functional protrusions. Protrusion stabilization, cluster integrity, and migration are contingent on optimal distribution of the adhesion molecule, E-cadherin. Immunofluorescence characterization of E-cadherin showed variations in the levels and localization in border cells with reduced ecdysone signaling. We hypothesize transcription by EcR is important for migration and could explain some of the above phenotypes. Thus, we are studying potential migratory roles of EcR targets. Additionally, we are assaying EcR binding sites within migration-specific target gene loci using bioinformatic and chromatin immunoprecipitation analyses. Interestingly, an EcR transcriptional reporter is specifically activated in the cluster despite the apparent availability of the signalling components throughout the egg chamber. We are investigating possible reasons for transcriptional specificity such as genetic interactions between chromatin regulators and EcR. Elucidating the role of EcR in cell migration kinetics will be a useful guide to improve our understanding of nuclear hormone receptors in development and disease.

125  Cell intruder targeting system mediates paternal mitochondrial destruction after fertilization in *Drosophila* Sharon Ben-Hur, Sara Afar, Eli Arama Weizmann institute of science, Israel

Uniparental inheritance of mitochondria is one of the most conserved phenomena across evolution, occurring in diverse species, ranging from fungi and plants to mammals. In almost all animals, the sperm enters the egg with its mitochondria, resulting in a short period where mitochondria from both parents reside within the forming zygote. However, rapidly after fertilization, sperm mitochondria are eliminated, leaving the zygote solely with maternal mitochondria, which will further propagate in the developing embryo. Why and how paternal mitochondria are eliminated remain a matter of speculation and controversy, respectively. It is believed that massive heteroplasmy (i.e. the presence of more than one mitochondrial (mt)DNA type in a cell) is deleterious to the organism, but the overwhelming number of egg mitochondria (up to 600,000 in human) as compared with sperm mitochondria (50-75 in human) could not lead to massive heteroplasmy. Furthermore, in some organisms, such as *Drosophila*, sperm mtDNA is eliminated already during spermatogenesis, yet sperm vacuolar mitochondria are still targeted for destruction after fertilization. Whether the mechanism of paternal mitochondrial elimination is conserved across species is unclear. Some studies suggest a passive mechanism whereby paternal mitochondria are diluted beyond detection by the overwhelming number of maternal mitochondria. On the other hand, other studies, including in *Drosophila*, suggest an active egg-derived paternal mitochondrial destruction mechanism.

Here, I will present our recent unpublished discovery of a non-canonical cell intruder targeting system, consisting of combined elements from the endocytic, phagocytic, and autophagic systems, which specifically targets sperm mitochondria for destruction after fertilization in *Drosophila*. These elements are conserved between *Drosophila* and human, implying that similar mechanisms might also operate to eliminate paternal mitochondria, at least in all flagellated sperm organisms.

126  Transcriptome analysis implicates circadian clock genes in Sex Peptide-dependent post-mating responses in *Drosophila melanogaster* females Sofie Delbare, Sara Venkatramen, Martin Wells, Sumanta Basu, Mariana Wolfner, Andrew Clark Cornell University

Sex Peptide (SP), a seminal fluid protein of *D. melanogaster* males, elicits an array of post-mating responses in females, including increased egg laying, activity and food intake, with a preference for proteins and a switch from carbohydrate to amino acid metabolism. To determine how one protein can have such widespread effects, we set out to dissect the genetic architecture of the female’s response to SP, and determine whether SP alters the expression of several regulators targeted to specific post-mating responses or acts on a pleiotropic regulator that controls multiple responses. We performed bulk RNA-seq of female heads at 10 time points within the first 24 hours after mating, sampling virgin females, females mated to control males and females mated to SP-null males. Using gene- and exon-level differential expression (DE) analyses, we find 666 genes and 86 exons that are DE across the 3 treatments. The earliest DE features include 3 regulators of circadian rhythms (*cwo, Clk, Pdp1*), leading us to hypothesize that SP acts on the circadian clock to induce at least a subset of post-mating responses. Consistent with this, we find that 16% of the 752 DE features have circadian expression patterns in virgin females, and their rhythms are altered after mating. Cluster analysis shows that features with altered circadian rhythms follow either of two patterns: upregulated during both day and night, or downregulated specifically during the night, relative to virgin females and females that did not receive SP. The downregulated genes include ones involved in glycogen biosynthesis, transcription factors involved in energy metabolism, and *Acer*, whose downregulation is known to cause increased activity at night. Upregulated genes include ones involved in amino acid metabolism, transcription factors including *Clk*, and signaling molecules. Among these are exons of *foraging*, a well-studied gene with roles in feeding behavior, of which some exons follow a circadian rhythm in virgin females, but no longer after receipt of SP. Overall, our transcriptome analysis suggests that SP might act on genes in the circadian clock to alter the expression of target genes with various functions. We are further investigating networks of genes and exons using an empirical Bayes clustering approach and transcription factor motif enrichment analyses, to better understand how SP orchestrates multiple post-mating responses.

127  Genetic coordination of sperm morphology and seminal fluid proteins promotes male reproductive success
Spermatozoal morphology is highly variable both among and within species and in ways that can significantly impact fertilization success. In *Drosophila melanogaster*, paternity success depends on sperm length of both competing males and length of the female's primary sperm storage organ. We found that genes upregulated in long sperm testes are enriched for IncRNAs and seminal fluid proteins (Sfps). Transferred in seminal fluid to the female during mating, Sfps are secreted by the male accessory glands (AG) and affect female remating rate, physiology, and behavior with concomitant advantages for male reproductive success. Despite being upregulated in long sperm testes, they have no known function in tests tissue. We found that Sex Peptide and ovulin (Acp26Aa) knockouts resulted in shorter sperm, suggesting that Sfps may regulate sperm length during spermatogenesis. However, knockout of AG function did not affect sperm length, suggesting that AG expression has no influence on spermatogenic processes. We also found that long sperm males are better able to delay female remating, suggesting higher Sfp expression in AG. These results might suggest that long sperm males have a double advantage in sperm competition by both delaying female remating, likely through transfer of more Sfps, and by resisting sperm displacement. However, we also found that this extra advantage does not necessarily translate to more progeny or higher paternity success. Thus, we found that multiple components of the ejaculate coordinate to promote male reproductive success at different stages of reproduction, but the realized fitness advantages in sperm competition are uncertain.

136  Visceral organ morphogenesis via calcium-patterned muscle constrictions Noah Mitchell1, Dillon Cislo1, Suraj Shankar1,2, Yuzheng Lin1, Boris Shraiman1, Sebastian Streichan1 1) University of California Santa Barbara, Santa Barbara, CA; 2) Harvard University, Cambridge, MA

During development, interacting tissue layers orchestrate complex shape changes to form visceral organs. Tracing the dynamics and mechanical interactions across tissue layers and across scales -- from cell to tissue, to entire organs -- remains an outstanding challenge. Here, we study the embryonic *Drosophila* midgut as a model visceral organ. Using light-sheet microscopy, genetics, computer vision, and tissue cartography, we reconstruct in toto the dynamics of individual tissue layers to map the time course of shape across scales. We identify the kinematic mechanism driving shape change by linking out-of-plane motion to active contraction patterns, revealing a convergent extension process in which cells deform as they flow into deepening folds. Optogenetic perturbations of muscle contractility reveal that muscle activity drives these contraction patterns and induces cell shape changes in the adjacent endoderm layer. This induction cascade relies on calcium pulses in the muscle, under the control of hox genes. Our study of multi-layer organogenesis reveals how genetic patterning in one layer triggers a dynamic molecular mechanism to control a physical process in the adjacent layer, orchestrating whole-organ shape change.

137  Maintaining symmetry in morphogenetic movements Celia Smits1,2, Sayantan Dutta1,3, Sebastian Streichan4,5, Stanislav Shvartsman1,2,3,6 1) Lewis Sigler Institute for Integrative Genomics, Princeton University, NJ; 2) Department of Molecular Biology, Princeton University, NJ; 3) Department of Chemical and Biological Engineering, Princeton University, NJ; 4) Department of Physics, University of California Santa Barbara, CA; 5) Biomolecular Science and Engineering, University of California Santa Barbara, CA; 6) Center for Computational Biology, Flatiron Institute, Simons Foundation, NY

Morphogenetic processes must resist biological noise and environmental perturbations to faithfully construct an adult organism. We are investigating the mechanisms underlying such robustness in *Drosophila* embryogenesis. Here, we focus on bilateral symmetry maintenance during body axis extension. In *Drosophila*, the body axis extends parallel to the long axis of the embryo during gastrulation. However, we present evidence that there is a certain amount of left-right variation in midline position that is corrected as the tissue elongates. To determine the origin of this correction, we investigated a class of mutants where embryos exhibit a characteristic “corkscrew” phenotype. We find that in these mutants, the midline begins twisting along the left-right axis shortly after the onset of axis extension. Using a combination of classical genetics, light sheet microscopy, particle image velocimetry, dimensionality reduction and computation modelling, we characterize the mechanics and genetics that underlie this twisting. Finally, we propose a model where invagination of the posterior midgut acts to disperse the forces of convergent extension in a bilaterally symmetric manner, which is required to maintain bilateral symmetry in the face of stochastic differences in lateral extension rates. Overall, this work indicates that coordination of morphogenetic movements and mechanics act to rein in random biological noise and ensure correct tissue flow and proper morphogenesis.

138  Malvolio, a Fork head target metal ion transporter, is required for salivary gland morphogenesis Srihitha Akula, Rajprasad Loganathan, Tony Zou , Rika Maruyama, Deborah Andrew Johns Hopkins University

Malvolio (Mvl) is a member of the SLC11 family of metal ion transporters, and is the Drosophila ortholog of the mammalian natural resistance-associated macrophage proteins (NRAMPs). The family members (NRAMP1 and NRAMP2) function as general metal ion transporters that use proton motive force to transport Fe2+, Cu2+, Mn2+, Zn2+, Cd2+, Ni2+, and...
Adherens junctions play a major role in assuring cell-cell adhesion, and through their linkage to the cytoskeleton, mediate dramatic shape changes and rearrangements during embryonic morphogenesis. The junctions need to be strong yet also dynamic, to allow cell rearrangement without tissue rupture. We want to understand how junction-associated proteins, like the multi-domain scaffold Polychaetoid (Pyd; fly homolog of ZO-1), work together to allow junctions to react dynamically to cell shape change and force generation. We hypothesize that the junctional complex and its linkage to the cytoskeleton involves multivalent connections among multiple junction-associated proteins, assembled into an underlying structured molecular complex that mediates binding to the cytoskeleton. I am using structured illumination microscopy (SIM) to explore the junctional localization of these proteins in high resolution. My preliminary data suggest that the core junction-associated proteins Canoe (Cno) and Armadillo (β-Catenin) localize to alternating clusters along the zonula adherens of cell-cell borders, often with little overlap. Furthermore, I found that Pyd surrounds these core junctional proteins in a cloud-like pattern of puncta. I am now broadening my analysis to include other junction-associated proteins like Bazooka (Par-3), and Non-muscle myosin II. Next, I will analyze how these patterns change in pyd mutants. In parallel, I am exploring the functional role of Pyd as one player in this robust network. While Pyd is not absolutely essential for embryonic viability, mutants display defects in cell rearrangements during germband extension. Mutants have stacks of elongated cells, an elevated number of rosettes and junctional disruptions at tricellular junctions and anterior-posterior cell borders, suggesting defects in responding to elevated force on cell-cell junctions. However, these occur without major tissue rupture, suggesting other proteins act in parallel. To explore a hypothesized change in junction dynamics in pyd embryos, I am applying live imaging during germband elongation. Furthermore, we are now expanding our analysis to include other players in this protein network, including the tricellular junction protein Sidekick, and Canoe’s regulators Rap1 and the GEF Dizzy.

A feedback mechanism mediated by actomyosin activity-dependent apical targeting of Rab11 vesicles reinforces apical constriction

Wei Chen, Bing He Dartmouth College

During tissue morphogenesis, cell shape changes resulting from cell-generated forces often require active regulation of intracellular trafficking. How mechanical stimuli influence intracellular trafficking and how such regulation impacts tissue mechanics are not fully understood. Non-muscle myosin II (Myosin II) is an important force generator in many cell and tissue types that acts through its contraction on actin network. In this study, we identify an actomyosin activity-dependent mechanism involving Rab11-mediated trafficking in regulating apical constriction in the Drosophila embryo. During Drosophila mesoderm invagination, apical actin and Myosin II (actomyosin) contractility induces accumulation of Rab11-marked vesicle-like structures (“Rab11 vesicles”) near the apical membrane. These Rab11 vesicles are not derived from apical endocytosis. Instead, apical accumulation of Rab11 vesicles is associated with constant vesicle transport along cell apical-basal axis, which is mediated by microtubule motor dynein. Interestingly, actomyosin contractility promotes apical targeting of Rab11 vesicle by inhibiting the reverse transport, the resulting directional bias towards apical side therefore contributes to the apical accumulation of Rab11 vesicles. At the apical domain, Rab11 vesicles are targeted to the adherens junctions (AJs). The apical accumulation of Rab11 vesicles is essential to prevent fragmented apical AJs, breaks in the supracellular actomyosin network and therefore promotes the stepwise, ratchet-like cell area.
Mechanical cues planar polarize Pins and orient divisions during *Drosophila* gastrulation Jaclyn Camuglia, Soline Chanet, Adam Martin 1) Massachusetts Institute of Technology; 2) Collège de France

Epithelial morphogenesis and homeostasis are properly achieved by oriented cell divisions that occur along specific directions relative to both the epithelial plane and the polarity of an organ/organism. In epithelial sheets of cells, divisions can occur either perpendicular or parallel (planar) to epithelial sheets. Planar cell divisions can further exhibit polarity in specific orientations along epithelial tissues (planar polarization). Spindle orientation is often achieved by a complex of Pins/LGN, Mud/NuMa, Gai, and Dynemin, which interacts with astral microtubules to rotate the spindle. Here, we identify the mitotic domains of the early embryo as a system to study Pins-mediated planar polarized divisions during morphogenesis. Mitotic domains (MDs), 1, 3, 5, and 14 exhibit oriented division and these oriented divisions are dependent on proper Pins localization and activity. We find that the divisions within these domains are oriented within 30 degrees of the anterior-posterior (AP) axis of the embryo and that Pins is localized in planar polarized crescents within the domains. Disruption of Pins localization and activity via expression of a myristoylated version of Pins leads to misoriented divisions. It is currently unknown what causes Pins localization in this context. Both planar polarity proteins and mechanical force have been shown to localize Pins and cue spindle orientation during division. We find that in MDs 1, 3, 5, and 14 canonical planar cell polarity pathways are not solely responsible for the orientation of the divisions. Further, we find that adherens junctions, which mechanically couple cells within epithelia, are necessary for the orientation of the divisions. Disruption of a-catenin disrupts division orientation and pins polarization. Our findings suggest mechanical forces, specifically those generated during morphogenesis are responsible for Pins polarization and spindle orientation. Disruption of forces through a chemical inhibitor, laser ablation, and genetic perturbation causes divisions to fail to orient along the AP axis and abolishes Pins polarization. To our knowledge, demonstrating that mechanical force polarizes Pins to mediate division orientation has not yet been shown in vivo.

Tissue geometry reorients in-plane homeotic tension to promote folding. Aurélien Villedieu, Lale Alpar, Isabelle Gaugue, Amina Joudat, François Graner, Floris Bosveld, Yohanns Bellaiche 1) Institut Curie, PSL Research University, CNRS UMR 3215, INSERM U934, Paris, France.; 2) Sorbonne Universités, UPMC Univ Paris 06, CNRS UMR 3215, INSERM U934, Paris, France.; 3) Matière et Systèmes Complexes, Université de Paris - Diderot, CNRS UMR 7057, Paris, France

Shape is a conspicuous and fundamental property of biological systems entailing the function of organs and tissues. While much emphasis has been put on how tissue tension and mechanical properties drive shape changes, how a given initial shape influences subsequent morphogenesis remains poorly characterized. Here, we analyzed invagination of the *Drosophila* neck epithelium, which exhibits a curved radial geometry prior to its formation. We found that neck folding depends on an in-plane tension promoted by the *Deformed* (*Dfd*) homeotic gene. By varying tissue curvature, we established that tissue geometry promotes an inward force driving folding. Accordingly, the interplay between *Dfd*-dependent in-plane tension and curvature explains the spatiotemporal dynamics of tissue invagination. Collectively, our work highlights how tissue geometry and homeotic regulation of tissue tension provide a simple design principle contributing to folding morphogenesis during development.

The flipside of tissue growth: how the two layers of the wing imaginal disc keep pace with each other. Sophia Friesen, Iswar Hariharan University of California, Berkeley

Epithelial sheets of cells are a fundamental building block of animal tissue. To build functional, proportional organs, epithelia must coordinate their growth with the growth of other epithelial and non-epithelial tissues. How this synchronized growth occurs is a key question in developmental biology. To understand coordinated epithelial growth, we turned to the wing imaginal disc, a larval organ that develops into the adult wing and thorax. The wing disc consists of two layers of simple epithelia that are connected at their edges (similar to a pita pocket). These two layers have very different morphologies – the disc proper (DP) is composed of many tightly-packed columnar cells, while the peripodial epithelium (PE) is made up of relatively few cells that are very flat and broad – but despite their differences, the two layers grow at the same speed throughout larval development. We wanted to know what regulates growth of the PE, and how it keeps pace with the DP, in order to better understand how epithelial proportionality is maintained during development.

We found that even though the two layers grow at the same speed, they use very different pathways to do so. Growth of the DP absolutely requires the secreted morphogens Hedgehog and Dpp, but we found that most PE growth can occur without these signals. In contrast, several components of the mechanosensitive Hippo pathway, including Yorkie (Yki) and multiple transcriptional cofactors, are critical for growth of the PE. Yki is required for PE survival and proliferation, as it is in many tissues, but it also mediates the dramatic cell shape changes that are unique to this layer of the disc. Yki activity can be upregulated by mechanical stretching forces, and so could directly link growth of the PE to growth.
of the DP. We speculate that DP growth stretches the PE, which increases PE Yki activity. Increased Yki activity, in turn, promotes growth of the PE, so that the two layers grow at the same speed. In this case, use of two very different growth paradigms—one based on morphogen gradients, and one based on mechanical force—seems to be the key to synchronized epithelial growth.

144 Uncovering how the pioneer transcription factor Grainy head binds and opens chromatin Meghan Freund, Andrew Rashoff, Tyler Gibson, Peter Lewis, Melissa Harrison University of Tennessee, Knoxville

Multicellular organisms are made up of numerous differentiated cell types that perform unique functions. Beginning from a single cell, organisms undergo a journey of differentiation, resulting in the formation of this variety of unique cell types. Because these cells all possess the same genome, distinct cell types are not due to differences in genotype, but rather differences in gene expression. Transcription factors bind DNA and drive this differential gene expression. However, compacted chromatin structure can act as a barrier to transcription-factor binding. Specialized transcription factors, known as pioneer transcription factors, can overcome this barrier by binding to condensed chromatin and increasing chromatin accessibility at previously inaccessible loci. While pioneer factors share the ability to bind condensed chromatin, the exact mechanisms are not understood as these factors all act through divergent DNA-binding domains. Grainy head (GRH) is an essential pioneer factor that drives epithelial cell fate and when misexpressed, can lead to cancer. It is conserved across species ranging from worms to humans and binds the same canonical DNA sequence in all species studied. These features make GRH a great candidate to elucidate the basic mechanisms by which pioneer factors engage the genome and their role in shaping cellular identity. Furthermore, while mammals have three GRH-like proteins, Drosophila encode GRH from a single gene, simplifying functional studies. To understand the properties of GRH that allow it to access compacted chromatin, we used in vitro studies to show that GRH can bind DNA in the context of nucleosomes. Using structural analysis, we identified mutations in the DNA-binding domain that mediate either sequence-specific or non-specific interactions and confirmed these properties in vitro. We are using these mutants to determine how this conserved protein interacts with DNA through its DNA-binding domain. We have further developed a tissue-culture system to investigate these mechanisms in vivo. Together our data will determine how GRH scans the genome and recognizes its motifs within inaccessible chromatin.

145 Image-based investigation of enhancer-promoter bridging in the Drosophila genome Aleena Patel1, Leslie Mateo 1,2, Sedona Murphy1, Angela Pogson1, Alistair Boettiger1 1) Stanford University, Stanford, CA; 2) Genentech, South San Francisco, CA

In metazoans, cis-regulatory DNA sequences (enhancers) often lie kilobases away from the target genes’ proximal transcription start sites (promoters), where transcriptional machinery is assembled. Various assays of the physical arrangement of sequences in 3D space have not yet solved controversial debates about which mechanism, or combination of mechanisms mediate such long-range communication in Drosophila. In mammals, ATP-dependent motors on the cohesin complex walking in opposite directions along DNA while tethered to one another may bridge distally located enhancers and promoters. This mechanism requires a DNA-bound factor such as the CCCTC binding factor (CTCF) to stall cohesin at the right loci. Despite a non-conserved function for CTCF in Drosophila, perturbations of cohesin components and regulators quantitatively change gene expression and suggest a possible role for active loop extrusion in enhancer-promoter contact. Alternatively, point-to-point loops could be formed by protein tethers that interact when thermal fluctuations in the DNA polymer bring their binding sites together. Several Drosophila DNA-binding factors are found near the boundaries of topologically associating domains (TADs) and may mediate physical tethering of select DNA loci by interacting or phase separating with one another. We investigate these proposed mechanisms for Drosophila genome folding with optical reconstruction of chromatin architecture (ORCA), a super-resolution imaging technique. We have fluorescently labeled 80 5-kilobase regions spanning TADs that harbor enhancer-promoter loops for the genes scyl and chrb in Drosophila tissues. From the imaging data, we have reconstructed individual polymers tracing the DNA path in single cells. ORCA has been performed on embryos depleted of loop-associated factors, including a candidate tether, GAGA binding factor (GAF), the cohesin component, Rad21, the cohesin loader, Nip8, and the cohesin unloader, WAPL. We find that GAF depletion specifically disrupts enhancer-promoter loop formation. Positive and negative perturbations of cohesin function have opposing effects on overall TAD compaction without disrupting the GAF-dependent loops. I will discuss how these results, and the insights of the Drosophila model system, are promising to expand current models of genome organization.

146 Drosophila genome architectural proteins form in vivo liquid -liquid phase separating Bright Amankwaa, Ryan Simmons, Ran An, Mariano Labrador University of Tennessee, Knoxville

Drosophila insulator proteins, which share genome architectural properties with their vertebrate counterparts, coalesce to form nuclear foci known as insulator bodies in response to osmotic stress. However, the mechanism behind the formation of these bodies remains unknown. Here, we identify signatures of liquid-liquid phase separation (LLPS) in insulator bodies. Mounting evidence implicates LLPS in the formation of membraneless organelles utilized for various
biological functions, including genome structure and function. We show that insulator proteins have a high disorder tendency, assemble into insulator bodies in a scaffold-client dependent manner, have extensive fusion behavior, and show sensitivity to 1,6-hexanediol. Further, we characterize the Cohesin subunits Rad21 and Wapl as well as the phosphorylated version of the histone variant H2Av (γH2Av) as novel components of insulator body condensates. Our data suggest a concerted role of Cohesin, γH2Av and insulator proteins in the formation of insulator bodies and under physiological conditions. We propose a model whereby these architectural proteins modulate the 3D genome organization of *Drosophila* through LLPS.

### Chromatin state transitions in the *Drosophila* intestinal lineage gives new insights into cell type specification

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Adult stem cells self-renew and differentiate into one or several cell types, thus ensuring tissue homeostasis. Understanding their regulation is crucial to have a better comprehension of uncontrolled proliferation and altered differentiation mechanisms occurring during tumorigenesis and age-dependent functional decline of tissues. Here, we aim to better understand what chromatin states are associated with adult stem cell activity in vivo in a homeostatic tissue using the *Drosophila* adult intestine as a model. Our lab has previously provided evidence of roles of conserved chromatin factors in controlling intestinal stem cell (ISC) proliferation (Gervais et al, 2019), highlighting their importance in the regulation of the intestinal lineage. Here, we expanded on these studies to investigate chromatin state changes associated with stem cell differentiation at the genome-wide scale. By generating cell-type specific whole-genome binding maps of 5 chromatin proteins (RNA Pol II, Brahma, Polycomb, HP1 and H1) using Targeted DamID and performing subsequent Hidden Markov modelling to define chromatin states, we found that 7 major chromatin states exist in the ISC lineage. Examining these states at specific genes revealed dynamic changes within the lineage. First, ISC-specific transcription factors such as *esg, klu, Sox100B* become marked by the Blue (Polycomb-enriched) state in differentiated cells. In contrast, components of signaling pathways regulating ISC activity are found in distinct states depending on the differentiated cell type, suggesting that they are repressed in different ways. In addition, the key transcription regulators of lineage specification including *pros* and *pdm1* undergo a transition from the Blue state (Polycomb-enriched) to active states upon differentiation, suggesting a role for Polycomb group proteins in repressing these genes in ISCs. While these data suggest a regulatory function of Polycomb-marked chromatin for control of the transcriptional hierarchy within the ISC lineage, we find that genes involved in differentiated cell physiology are instead associated with Histone H1-enriched Black chromatin. Indeed, physiology and metabolic activity-related genes follow a transition from the Black state in ISCs to active states upon their activation, suggesting a previously uncharacterized mode of regulation of physiology-related genes. On-going work will highlight the biological relevance of these chromatin transitions by perturbing specific states and studying the subsequent transcriptional changes and effects on proliferation and differentiation in the intestinal lineage. Overall, the extensive characterization of chromatin state changes during differentiation will provide a valuable resource to better understand the regulatory programs that control cell fate and identity, as well as physiological functions in this homeostatic tissue.

### Condensin II loss ameliorates long-range chromosomal interactions in both active and inactive physical compartments within a chromosome territory

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Metazoan genomes are folded into structures at a variety of length-scales, from chromatin loops and domains to higher-order interactions that connect distal genomic regions to form chromosome territories. Polymer modeling of Hi-C data suggests that chromosome folding is driven by a loop extrusion mechanism to both fold entire chromosomes into territories and restrict their interactions with other chromosomes. Recent work from our lab and others suggest that the condensin II loop-extruding complex specifically regulates chromosome territory formation and levels of inter-chromosomal interactions during interphase in *Drosophila* and human cells. To dissect the mechanism by which condensin II modulates genome folding, we are using a combination of Oligopaint FISH and Hi-C in *Drosophila* cultured cells to show that condensin II is required for proper inter-domain interactions in a chromatin state-nonspecific manner without altering interactions within domains. Optimized ChIP-Seq of a condensin II component reveals a novel class of genomic binding sites that are associated with long chromatin loops and colocalize with the gypsy complex, a strong insulator at the interface of active and inactive chromatin. *In vivo* data suggest that these condensin II-dependent interactions are important for proper silencing of heterochromatic sequences that are distal, but not proximal, to the pericentromeric heterochromatin. Overall, we propose a model in which condensin II extrudes chromatin into large loops until blocked by an insulator prior to cohesin activity early in interphase, thereby bridging distal interactions that fold chromosomes into distinct territories. Ongoing experiments will determine the functional consequences of condensin II loss on gene regulation.

### Histone gene replacement reveals functional independence, redundancy and synergism between lysine 36 of...
Polycomb silencing is an epigenetic mechanism for determining cell-type specific gene expression programs, ensuring that genes specifying alternative lineages remain repressed. The best-studied examples of this phenomenon are the Drosophila Hox genes, which specify the body plan. To ensure proper development of body segments, Hox genes must be expressed at certain points along the body axis and repressed at others.

One critical step in this process is trimethylation of histone H3 lysine 27 (H3K27me3) by the Polycomb repressive complex 2, PRC2. Importantly, PRC2 can also “read” the H3K27me3 mark, enabling cis-linked spreading of H3K27me3, and formation of repressive chromatin domains essential for proper Hox gene expression. To antagonize the spreading of H3K27me3 into other active genes, evidence suggests that cells utilize di- and tri-methylation of H3 lysine 36 (H3K36me2/3) to abut these repressive domains. A mechanism by which H3K36me2/3 halts spreading of facultative heterochromatin was unknown until recent structural studies revealed that unmethylated H3K36 occupies a crucial role in positioning H3K27 into the active site of its cognate methyltransferase, EZH2 (the mammalian ortholog of Drosophila enhancer of zeste E(z)). This finding suggests that H3K36me2/3 sterically hinders E(z) activity and that mutation of H3K36 would elicit strong Polycomb phenotypes. One puzzle that emerged from this work is that when Hox gene expression was examined in vivo, mutants of the replication-dependent histone H3 genes (H3.2K36R and H3.2K36A) displayed relatively mild Polycomb phenotypes, despite inhibitory effects of these mutations on H3K27 trimethylation in vitro.

In addition to H3.2, metazoan genomes also contain the replication-independent histone variant, H3.3, which differs from H3.2 by only 4 amino acids. We hypothesized that H3.3 might function redundantly to enable Polycomb mediated gene repression in the H3.2 K36R mutants. Therefore, we generated an H3.3K36R mutant to directly compare Polycomb mediated gene silencing between H3.3K36R, H3.2K36R, and H3.3K36R/H3.2K36R combined mutant animals. Our studies in embryos, first instar larvae, and adults demonstrate that H3.2K36 and H3.3K36 cooperate to maintain H3K27 trimethylation and Hox gene silencing by different, but synergistic mechanisms.

Frontotemporal dementia (FTD) is a neurodegenerative disease associated with mutations in the microtubule binding protein Tau. The clinical presentation of FTD is heterogeneous with patients exhibiting parkinsonism, dementia, atrophy in the temporal lobes, and personality changes. Current treatments can mitigate aspects of the behavioral changes associated with FTD, however, no therapies are available to slow the progression. Since patient sample procurement is restricted to post-mortem tissue, our understanding of the progression and underlying pathogenic mechanisms of this disease is limited. Recent work in model systems and post-mortem tissue has shown that expression of FTD-associated mutant Tau may lead to epigenetic modifications that alter gene expression. In our lab, we model FTD using Drosophila, which allows us to conduct longitudinal studies to observe FTD progression throughout the adult lifespan. The adult Drosophila expressing FTD-associated mutant human Tau (hTau) have age-dependent neurodegenerative vacuoles, axonal changes, locomotion defects and impaired memory while flies expressing normal hTau did not. This confirms that our models show pathogenic phenotypes associated with Tauopathies and it provides the basis to now use these models to identify molecular mechanisms of pathogenicity. We also found changes in a chromatin associated protein, Heterochromatin Protein 1 and hypothesized that FTD mutant Tau could alter chromatin accessibility. We used Single-cell Combinatorial Indexed Assay-for-Transposase-Accessible Chromatin using sequencing (sci-ATAC-seq) to assess how human Tau FTD mutations alter chromatin accessibility in the young and aged adult Drosophila brain for three clinically distinct Tau mutations (K369I, P301L, V337M). We generated >90,000 single-cell chromatin accessibility profiles from whole adult heads. Comparing our wildtype hTau insertion line to the FTD mutants revealed differentially accessible regions in both neuronal and glial cell populations. This work will further our understanding of chromatin structure in hTau knock-in models at single-cell resolution and provide insight into cell-type specific variation.

Developmental progression and cellular diversity are largely driven by transcription factors (TFs), yet characterizing their loss-of-function phenotypes remains challenging and often disconnected from their underlying molecular mechanisms. To overcome this, we have combined single cell regulatory genomics with loss-of-function mutants to simultaneously assess both cellular and molecular phenotypes.
We performed single cell chromatin accessibility profiling by sci-ATAC-seq for over 20,000 cells at eight overlapping time-points during *Drosophila* mesoderm development, ensuring that key developmental transitions are captured and can be directly traced, including stages of multipotency, specification and terminal differentiation. This dataset captures the establishment of regulatory landscapes in the nascent embryonic mesoderm at high resolution and its progression along developmental time. We demonstrate that this information can reconstruct multiple developmental trajectories of all major and rare cell-types, including the heart, and uncover the enhancers and TFs involved.

To systematically assess the TFs mutant phenotypes, we developed a streamlined strategy to robustly profile chromatin accessibility and digitally genotype mutant cells coming from embryo collections of mixed genotypes. Applying this approach to four TF mutants could uncover and quantify their characterized phenotypes *de novo*, and discover new ones, while simultaneously revealing their regulatory input and mode of action. We observe strikingly different outcomes in cell fate, as one mutant (*Mef2*) results in abnormal ‘new’ cell states, while other TF mutants appear to block cell populations much earlier along their developmental trajectory.

Our approach is a general framework to dissect the functional input of TFs in a systematic unbiased manner, uncovering both cellular and molecular phenotypes at a scale and resolution that was not feasible before.

152  **A KDM5-Prospero transcriptional axis functions during early neurodevelopment to regulate mushroom body formation**  
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Intellectual disability (ID) disorders affect up to 2% of the population and are characterized by an IQ score less than 70 with deficits in adaptive functioning. Our research focuses on the KDM5 family of transcriptional regulators (KDM5A, KDM5B, KDM5C, and KDM5D), mutations in which account for 1-3% of inherited ID ranging from mild to severe. Although recent advances in comparative genomic hybridization and whole exome sequencing have revealed approximately 70 unique genetic variants in *KDM5C* segregating in families with inherited ID, the molecular mechanisms by which this family of proteins impacts neuronal function remain largely unknown, leaving patients without effective treatment strategies.

Here, we utilize the *Drosophila* Mushroom body (MB), a major learning and memory center within the fly brain, to demonstrate that *Drosophila* KDM5 is specifically required within ganglion mother cells (GMCs) and immature neurons for proper neurodevelopment. Utilizing Targeted DamID (TaDa), we identify a core network of KDM5-regulated genes within GMCs and immature neurons that are critical modulators of neurodevelopment. Significantly, we find that a majority of these genes are direct targets of Prospero (Pros), a transcription factor with well-established roles in neuronal growth and guidance. We further demonstrate that *pros* genetically interacts with *kdm5* to orchestrate a transcriptional program critical for proper MB development.

To better understand the molecular mechanisms through which KDM5 functions to regulate MB development, we generated a library of fly strains each bearing a conserved ID patient-derived *kdm5* missense mutation. We demonstrate that a subset of these strains presents with MB structural defects and deficits to long-term memory, as assessed *via* appetitive olfactory conditioning assays. Interestingly, fly strains bearing patient mutations that disrupt KDM5’s histone demethylase activity do not present with MB morphological defects, yet have impaired learning and memory. These data suggest that KDM5 may function through demethylase-independent mechanisms to regulate MB development, independent of its effect on cognitive function. This is significant, as the prevailing model linking KDM5 dysfunction to ID assumes that altered demethylase activity of KDM5 is largely responsible for such deficits. We are currently investigating how KDM5 may utilize specific protein domains in coordination with Pros to guide neuronal development.

153  **Neuronal mechanisms of neurofibromin dependent metabolic regulation**  
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Neurofibromatosis type 1 (NF1) is a genetic disorder predisposing patients to a range of complications, including benign tumor formations in the nervous system, altered cellular function, short stature, bone abnormalities, and increased rates of cognitive and developmental disorders. NF1 is caused by mutations in the *NF1* gene, which encodes neurofibromin (NF1), a large protein that functions as a negative regulator of Ras signaling and mediates pleiotropic organismal and cellular functions. Emerging data suggest that NF1 regulates metabolism: NF1 patients show a reduced body mass index, alterations in metabolites, lower incidences of diabetes, and increased resting energy expenditure. These changes
in metabolism may contribute to complications and symptoms associated with NF1. The mechanisms by which NF1 affects metabolism and energy expenditure are not well understood. Using the Drosophila melanogaster NF1 ortholog, we show that NF1 regulates metabolic homeostasis via neuronal mechanisms. Drosophila NF1 is ~60% identical to the human protein and similarly mediates Ras signaling. Our data show that the loss of NF1 increases metabolic rate via a Ras-GAP-related domain, increases metabolic rate, feeding rate, starvation susceptibility, and alters lipid stores and turnover kinetics. These metabolic alterations map to a restricted subset of neurons in the ventral nervous system and are independent of locomotion and grooming activity. Activation of this restricted neuronal circuit mimics the loss of NF1 function by increasing metabolic rate when stimulated. These data indicate that NF1 may regulate changes in neuronal metabolic control, suggest that cellular and systemic metabolic alterations may be a pathophysiological mechanism of NF1, and provide a platform for investigating the cellular role of neurofibromin in metabolic homeostasis.

154 Traip controls brain size via suppression of mitotic DNA bridges Ryan O’Neill, Nasser Rusan National Heart, Lung, and Blood Institute, NIH, Bethesda, MD

Microcephaly is a failure to achieve proper brain size and neuron number during development. Most microcephaly-linked genes function either at the mitotic spindle or in DNA damage repair (DDR). Spindle proteins are thought to control brain size via their functions in neural stem cell (NSC) mitosis, whereas DDR proteins are thought to suppress DNA damage and cell death. However, since few microcephaly genes are well-studied in neurogenesis, we set out to characterize microcephaly genes in Drosophila brains. Here, we investigated the microcephaly gene Traip, known to function in DDR, and found that Traip promotes proper brain size by suppressing DNA bridges in mitosis. We show that traip flies have microcephaly-like brain defects, including fewer neurons and marked loss of NSCs via caspase-dependent cell death. traip NSCs have normal levels of DNA damage in interphase, suggesting that Traip does not fit the canonical model of DDR functions in microcephaly. In contrast, traip NSCs are frequently aneuploid, multinucleate, or have micronuclei, suggesting mitotic failure. Live fluorescence microscopy of traip NSCs revealed frequent mitotic DNA bridges, providing a possible explanation for the observed nuclear defects via chromosome fragmentation and cytokinesis failure. We characterized fluorescence-tagged Traip transgenes: in interphase, Traip is nuclear, whereas in mitosis Traip localizes on spindles, furrow, and midbody. A Traip variant lacking the nuclear localization signal is absent from the nucleus, but localizes properly in mitosis and rescues traip brain phenotypes, showing that a mitotic Traip function is sufficient to suppress microcephaly. Together, our work challenges current thinking about the relationships between DDR, mitosis and microcephaly by showing that, instead of repairing DNA damage during interphase, Traip primarily functions to preserve NSC genome integrity by resolving mitotic DNA bridges. Now, using Traip as a model microcephaly gene, we are using whole brain imaging and 3D analysis to screen for suppressors and uncover downstream pathways that mediate microcephaly phenotypes. To date, we have found roles for MAPK and Toll signaling, abscission kinase, and caspase-dependent cell death. We are now testing whether these pathways also mediate the phenotypes of other microcephaly genes, including both DDR and mitotic spindle genes, and targeting these pathways as potential therapeutic targets to minimize neuron loss in microcephaly.

155 Dietary restriction ameliorates TBI-induced phenotypes in Drosophila melanogaster. Rebecca Delventhal1,2, Emily Wooder2, Maylis Basturk2, Mohima Sattar2, Jonathan Lai2, Danielle Bolton2, Gayathri Muthukumar2, Matthew Ulgherait2, Mimi Shirasu-Higa2 1) Lake Forest College; 2) Columbia University Irving Medical Center

Traumatic brain injury (TBI) shares molecular and cellular hallmarks with neurodegenerative diseases (NDs), and is a major risk factor for developing neurodegeneration later in life. While our understanding of genes and pathways that underlie neurotoxicity in specific NDs has advanced, we still lack a complete understanding of molecular and physiological changes that drive neurodegeneration as an individual ages following a TBI. In recent years, many different studies have introduced Drosophila as a model organism for studying closed-head TBI, though they have primarily focused on acute or short-term outcomes of the TBI. In our research, we delivered a TBI to flies early in adult life, and then measured molecular and physiological phenotypes at short-, mid-, and long-term timepoints following the injury. We aim to identify the timing of changes that may contribute to progressive neurodegeneration, following a TBI. We confirmed prior work demonstrating a TBI-induced decline in lifespan, and uncovered evidence of a progressive decline in locomotor function, robust acute and modest chronic neuroinflammation, and a late-onset increase in protein aggregation. We also found evidence of metabolic dysfunction, in the form of starvation sensitivity and decreased lipids, that persisted beyond the immediate injury response, but resolved long-term. An intervention of dietary restriction (DR), or a low-protein diet, partially ameliorated some TBI-induced phenotypes, such as lifespan and locomotor function, though it did not alter the starvation sensitivity of injured flies. In the future, identifying molecular pathways altered in the short- or mid-term following TBI could suggest causality and point towards potential therapeutic targets.

156 Cyclin-dependent kinase 8 regulates mitochondrial morphology and modulates a Parkinson’s disease model in Drosophila Zhe Liao1, Kin Lam Wong2, Esther Verheyen1 1) Simon Fraser University, Canada; 2) Stanford University Stanford, USA

Cyclin-dependent kinase 8 is a serine/threonine kinase, which functions in regulating RNA polymerase II mediated...
transcription. Cdk8 forms a complex with Cyclin C, Mediator 12 (Med 12), and Mediator 13 (Med 13) that interacts with the core Mediator complex in a reversible fashion. While the regulatory role of Cdk8 in transcription is well-studied, we identified novel functions of Cdk8 in regulating mitochondrial morphology in Drosophila. When cdk8 is knocked down ubiquitously, we observe phenotypic effects including held-up and droopy wing postures, reduced life span, and defects in both flight and climbing abilities. Surprisingly, the observed phenotypic effects are characteristics of flies with either PTEN-induced putative kinase 1 (pink1) or parkin mutations, which are two well-known players associated with Parkinson’s disease (PD). Pink1 and Parkin normally function in regulating the homeostasis of mitochondria in a process known as mitophagy. Impaired or dysfunctional mitochondria will be recognized by Pink1 and Parkin and targeted for degradation by autophagy. Since tissue specific knock down of Cdk8 in either muscles or neurons also resulted in impaired climbing ability, we hypothesized that Cdk8 functions in a common pathway with Pink1 and Parkin in regulating mitochondria. Ectopic expression of Cdk8 in muscles significantly suppressed defects caused by the loss of function pink1 allele, pink1<sup>1<sup>, including rescue of the thorax indentation phenotype which is due to muscle degeneration, and rescued climbing activity relative to pink1<sup>1<sup>mutants. In addition, mitochondrial and muscle fiber morphologies were restored when Cdk8 was overexpressed in the pink1<sup>1<sup>mutant background. Finally, we examined the effect of Cdk8 on mitochondria under physiological conditions. Expression of Cdk8 is tightly associated with mitochondrial dynamics in a kinase-dependent fashion, as fission-like alterations were found after elevating Cdk8 expression and fusion-like morphology was induced after depleting cdk8 or expressing a kinase dead form of the protein. Together, we demonstrate that Cdk8 plays a novel function in regulating the homeostasis of mitochondria, and is able to revert the defective phenotypes caused by pink1<sup>1<sup>mutant allele. Our findings suggest that modulation of Cdk8 may serve as a potential therapeutic target for patients with PD.

157  **Endurance exercise ameliorates phenotypes in Drosophila models of Spinocerebellar Ataxias**

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Endurance exercise is a potent intervention with widespread benefits proven to reduce disease incidence and impact. While endurance exercise supports neural plasticity, enhanced memory, and reduced neurodegeneration, less is known about the effect of chronic exercise on the progression of movement disorders such as ataxias. Here, we focused on three different types of ataxias, Spinocerebellar Ataxias Type (SCAs) 2, 3 and 6, belonging to the polyglutamine (polyQ) family of neurodegenerative disorders. In Drosophila models of these SCAs, flies progressively lose motor function and accumulate levels of toxic SCAProteins. Excitingly, we observe dramatic protection of speed and endurance in exercised SCA2 flies and modest protection in exercised SCA6 models, while no benefit is observed in SCA3. Importantly, causative protein levels are reduced in SCA2 flies after chronic exercise, but not in SCA3 models, linking protein levels to exercise-based benefits. Currently, we are focusing on the activation of exercise-mimicking genes in SCA-model flies to define the mechanisms by which exercise preserves function in polyQ ataxias. The exercise-inducible protein dSestrin suppresses longitudinal mobility defects and improves early mortality in SCA2 flies, even without exercise. Furthermore, overexpression of dSestrin mimics exercise-induced reductions in disease protein in SCA2 flies by increasing autophagy. These improvements critically depend on previously-established functions of dSestrin that reduce oxidative damage and modulate mTOR activity. Our study suggests differential responses of ataxias to exercise, highlighting the potential for more extensive application of exercise-based therapies in the prevention of polyQ neurodegeneration.

158  **Loss of function variants in TIAM1 are associated with developmental delay, intellectual disability and seizures**

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The actin cytoskeleton is essential to maintain basic cell functions. Its dysregulation in neurons often leads to neurological diseases. T-lymphoma invasion and metastasis 1 protein (TIAM1) regulates Rac1 signaling pathways that affect the control of neuronal morphogenesis and neurite outgrowth by modulating the actin cytoskeletal network. To date, TIAM1 has not been associated with a Mendelian disorder. Here, we describe five individuals with biallelic TIAM1 missense variants who have developmental delay, intellectual disability, speech delay and seizures. We
found that the *Drosophila* ortholog of *TIAM1*, still life (*sif*), is expressed in larval and adult central nervous system (CNS), and is mainly expressed in a subset of neurons but not in glia. Loss of *sif* causes a severe reduction in viability, and the surviving adults exhibit climbing defects, are prone to severe seizures, and have a short lifespan. Both *sif* and *TIAM1* are toxic when ubiquitously overexpressed in *Drosophila*. Hence, either loss or gain of function of the gene affects development and survival. We assessed the toxicity associated with three *TIAM1* variants carried by two of the probands and compared them to the *TIAM1* reference cDNA function *in vivo*. The data suggest that they are all loss-of-function (LoF) variants: *p.L862F* is the most severe LoF variant, while *p.R23C* and *p.G328V* are less severe LoF variants. In summary, we provide evidence that *sif* is important for appropriate neural function and that *TIAM1* variants observed in the probands are disruptive, thus implicating loss of *TIAM1* in neurological phenotypes in humans.

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<th>Extremely rare variants in EIF4A2 are associated with a neurodevelopmental disorder characterized by hypotonia, intellectual disability and epilepsy</th>
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Eukaryotic Initiation Factor-4A2 (EIF4A2) encodes EIF4A2, an ATP-dependent RNA helicase subunit of the eIF4F complex, which recognizes the 5'cap structure of mRNAs and is required for mRNA binding to the ribosome. The fruit fly homolog *elf4a* mediates the negative regulation of Decapentaplegic (Dpp) signaling. In the fly, Dpp-signaling regulates embryo patterning, eye and wing morphogenesis, and stem cell identity determination. The vertebrate homolog of Dpp, TGF-β/BMP, is a key regulator of neuronal differentiation, development, and function and is associated with various neurological disorders. Prior fly studies revealed that both gain and loss of function (GOF and LOF) *elf4a* alleles modulate the rough eye and wing serration phenotypes associated with Dpp GOF and LOF, respectively. Despite the role of EIF4A2 homologs in key developmental processes, human disease-causing variants have not previously been identified. Here, we report eleven individuals with extremely rare variants in EIF4A2 who all present with global developmental delay or intellectual disabilities, significant hypotonia, and epilepsy in most cases. To determine the pathogenicity of the *EIF4A2* variants *in vivo*, we generated flies expressing human *EIF4A2* wild-type (WT) and variants with a C-terminal HA tag for four *de novo* variants. We used the GAL4-UAS system to selectively express human *EIF4A2* in fly neurons, wing, or eye. First, we conducted climbing assays to determine the impact of neuronal expression of *EIF4A2* *p.L344F*, *p.G364E*, and *p.T243I* and found that these variants resulted in motor defects. Second, we found that the wing-specific expression of *EIF4A2* *p.T216I* caused wing serration, which is consistent with loss of Dpp signaling. Third, we found that the eye-specific overexpression of *EIF4A2* *p.L344F*, *p.G364E*, and *p.T243I* exacerbates the rough eye phenotypes associated with Dpp GO, suggesting these variants are loss of function in nature. Finally, GMR GAL4 mediated knockdown of fly *elf4a* using two different RNAi lines results in pupal lethality that can be rescued using the expression of human *EIF4A2* WT. However, the *EIF4A2* *p.T243I* and *p.T216I* variants fail to rescue the pupal lethality. Together, these findings reveal that these *de novo* *EIF4A2* variants are pathogenic and alter fruit fly development in a dominant manner through either GO or LOF mechanisms. Our work establishes a role for *EIF4A2* dysfunction in human neurodevelopmental disorders.

168 | Cactin, a component of spliceosome C complex, is required for collective border cell polarization and migration in the *Drosophila ovary* Guangxia Miao, Li Guo, Denise Montell University of California-Santa Barbara |
Border cell migration in the Drosophila ovary serves as an in vivo model for the identification of molecular mechanisms that drive collective cell migration. While much is known concerning the physical, mechanical, and adhesive factors that steer the border cells and the cytoskeletal determinants of their morphology, less is known about the mechanisms that govern their initial delamination from the follicular epithelium and the coordination of individual cell polarization, morphology, and behavior. Here we report the identification of a gene, cactin, that is essential for these features of border cell migration. Cactin is conserved in evolution from yeast to man, yet has been ascribed to different biochemical activities in various cells, tissues, and organisms. We show that it is cactin’s conserved spliceosome function that is required in border cells, rather than its reported effects on the Dorsal inhibitor Cactus, or its effect on Rac-mediated actin dynamics. Whole transcriptome analysis shows widespread alterations in splicing, and surprisingly, transcription read-through. Viability and cell fate specification were normal in Cactin knockdown cells, which exhibited individual cell mobility. However, Cactin knockdown caused specific defects in generating a dominant lead cell protrusion and in cluster cohesion. As a result, the cells frequently failed to delaminate. Apical polarity complex proteins were more tightly concentrated in each individual cell, and lost from apical cell-cell junctions and protrusions. These results elucidate a requirement for the spliceosome complex in preventing transcription readthrough and show cell-type-specific defects caused by depletion of a general splicing factor.

169 The cytoskeletal mechanics that shape a stem cell niche Bailey Warder, Kara Nelson, Justin Sui, Lauren Anllo, Stephen DiNardo University of Pennsylvania

Stem cells often rely on signals from a niche, which in many tissues adopts a precise morphology. What remains elusive is how niches are formed, and how niche morphology impacts function. I use the gonadal niche to study mechanisms of niche formation, combining genetic tractability with powerful live-imaging techniques pioneered in our lab. This niche adopts a distinct morphology, with a smoothened boundary between itself and adherent germ line stem cells (GSCs). The niche plays key roles in regulating GSC behavior and it is thus vital to identify mechanisms of niche formation. We have found that the niche-GSC boundary is enriched for F-actin and Myosin II (MyoII). I therefore hypothesize that actomyosin contractility (AMC) shapes the niche, and makes it more efficient in guiding GSC behavior. Through transgenic and pharmacological manipulations, I show that precise levels of AMC are required for niche shape. Further, live-imaging shows that proper niche shape is crucial for function. Current work addresses mechanisms that robustly polarize F-actin and MyoII in the niche. Interestingly, AMC can be regulated by mechanical forces exerted on a cell. I have evidence that GSC divisions are required for niche morphogenesis, and I suspect that AMC in the niche is induced by forces inherent to GSC divisions. Our work suggests a unique feedback mechanism where stem cells shape the niche that guides their behavior. T32 GM007229 (BW), F32 GM125123 (LA), R35 GM136270 (SD)

170 Endocytic regulation of Fat protocadherins in tissue growth and morphogenesis Artem Gridnev, Jyoti Misra University of Texas at Dallas

Proper coordination of growth and morphogenesis during development is critical to formation of appropriately proportioned organs. The evolutionarily conserved protocadherins, Dachsous (Ds) and Fat constitute a signaling pathway that coordinates growth and morphogenesis by regulating the Hippo pathway and planar cell polarity (PCP) respectively. However, there are critical gaps in our understanding of how they regulate growth and morphogenetic processes. Further, little is known about how the spatial organization of the pathway is established and maintained and how the Ds-Fat junctions get coordinately remodeled along with other adhesion complexes to allow morphogenesis. To address these critical gaps, we use the Drosophila wing disc which provides a robust model system to study this pathway. In the developing Drosophila wing disc epithelium, Ds is expressed in a steep gradient with high expression from the periphery to the very low expression at the center of the wing pouch. In contrast, Fat is mostly expressed uniformly. The interaction between Ds and Fat is modulated by phosphorylation by the Golgi-resident kinase Four-jointed, which is expressed in an opposing decreasing gradient from the center to the periphery. The graded expression of Ds and Fj along with uniform expression of Fat results in an increasing gradient of Fat activity from the center to the periphery and slope of the gradient of Fat activity regulates cell proliferation by influencing Hippo signaling pathway. Membrane localization of Fat and Ds is dynamically regulated, where unliganded Fat is rapidly endocytosed, compared to the liganded population. Given that Ds level is very low at the center of the pouch region, most of the Fat should be unliganded and unstable. Therefore, there must be a mechanism to protect the unliganded Fat from endocytic turnover. We have identified a critical motif in the Fat cytoplasmic domain that bind to a key endocytic adapter and also have identified that the Lix1 homolog Low Fat competitively inhibit Fat endocytosis. Further, we have identified that an intricate interplay between recycling and endocytosis plays a central role in maintaining Fat levels. Taken together, these results indicate that vesicular trafficking provides an important layer of regulation in organization of the Fat signaling pathway, an aspect that has been overlooked so far. Furthermore, these studies provide novel mechanistic insight into Fat signaling pathway and address several longstanding questions in the field and will help explain the developmental disorders resulting from dysregulation of this pathway.

171 Inter-organ signaling regulates the onset of myoblast fusion Zhi-Rong Ruan1, Adwait Amod Sathe2, Chao Xing2, Elizabeth Chen1 1) Department of Molecular Biology, UT Southwestern Medical Center, Dallas, TX; 2) Eugene McDermott
Myoblast fusion, in which mononucleated myoblasts fuse to form multinucleated myotubes, is a critical step in skeletal muscle development and regeneration. In *Drosophila*, myoblast fusion is initiated by the interaction between muscle cell type-specific adhesion molecules. Here, we show that ecdysone signaling provides an extrinsic cue to regulate the onset of myoblast fusion by activating the expression of a cell adhesion molecule (CAM). Ecdysone is a steroid hormone best known for transcription regulation during pupal metamorphosis via binding to the ecdysone receptor (EcR). Surprisingly, we found that mutation in an ecdysteroidogenic enzyme, Spook (Spo), caused partial myoblast fusion defects in *Drosophila* embryos. Spo was expressed in amnioserosa, a major source for ecdysone production in early embryogenesis. The fusion defect in spo mutant was rescued by expressing spo in amnioserosa, but not in muscle cells, demonstrating a non-autonomous function of ecdysone at the tissue level. Since ecdysone is also provided maternally, we further disrupted ecdysone signaling by expressing a dominant negative form of EcR in muscle cells of spo mutant, which led to a severe fusion defect. Interestingly, the expression of Dumbfounded (Duf), a CAM required for the initiation of myoblast fusion, was significantly decreased in the spo;EcR<sup>om</sup> mutant. Conversely, overexpressing Duf in the spo;EcR<sup>om</sup> mutant significantly rescued the myoblast fusion defect, demonstrating that Duf is a major downstream target of ecdysone signaling in muscle cells. Indeed, we identified EcR-binding sites in the promoter of the duf gene and showed by in vitro luciferase assays that these sites mediated transcription activation by EcR. Furthermore, the EcR-binding sites are adjacent to the binding sites of a muscle-specific transcription factor, Twist. EcR and Twist synergistically activated duf transcription in vitro and in vivo. Taken together, these findings reveal an indispensable role for the extrinsic ecdysone signaling in regulating myoblast fusion and identify duf as one of the direct targets of ecdysone signaling in muscle cells. In addition, our studies demonstrate that the extrinsic ecdysone signaling coordinates with the intrinsic transcription factor Twist to synergistically activate the expression of duf to control the onset of myoblast fusion.

**172 Distinct contributions of ECM proteins to basement membrane mechanical properties in *Drosophila***

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The basement membrane is a specialized extracellular matrix (ECM) crucial for the development of epithelial tissues and organs. In *Drosophila*, the mechanical properties of the basement membrane play an important role in the proper elongation of the developing egg chamber, however, the molecular mechanisms contributing to basement membrane mechanical properties are not fully understood. Here, we systematically analyzed the contributions of individual ECM components towards the molecular composition and mechanical properties of the basement membrane underlying the follicle epithelium of *Drosophila* egg chambers. We find that the Laminin and Collagen IV networks largely persist in the absence of the other component. Moreover, we show that Perlecan and Collagen IV, but not Laminin or Nidogen, have a major contribution towards egg chamber elongation. Similarly, Perlecan and Collagen, but not Laminin or Nidogen, contribute towards the resistance of egg chambers against osmotic stress. Finally, we show using atomic force microscopy that basement membrane stiffness mainly depends on Collagen IV. Our analysis reveals how single ECM components contribute to the mechanical properties of the basement membrane controlling tissue and organ shape.

**173 Affinity-driven germline-soma interactions mediate *Drosophila* oogenesis***

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Germline-soma interactions are central to reproduction. In most vertebrate and invertebrate species, differentiating gametes are enveloped by a somatic epithelium, where the apical surface interacts with the germline cells. While signaling pathways acting on the two lineages have been described, little is known about how morphogenesis is coordinated between them. In the developing *Drosophila* egg chamber, the somatic follicle epithelium envelops germline-derived nurse cells and the growing oocyte to produce fertile eggs. We show that egg chamber morphogenesis can be divided into three major phases, that all require coordinated germline-soma morphogenesis. The expression of the transcriptional regulator Eya in follicle cells controls coordination between the germline and somatic follicle cells by differentially modulating affinity of the apical epithelial surface for nurse cells and oocytes. In all three phases, Eya expression patterns and levels are crucial to guarantee the right segregation of follicle cells over nurse cell and oocyte surfaces. Importantly, Eya-mediated affinity furthermore controls the spatial organization of nurse cells and the oocyte within the germline. Consequently, Eya is a master regulator of the soma-germline interface and through that controls all stages of egg chamber development.

Our work uncovers a new morphogenetic principle that enables the coordination of soma and germline and emphasizes the plasticity of epithelial behaviors during interactions with their adjoining tissues.
NGLY1 in ERAD

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N-glycanase 1 (NGLY1) deficiency is a rare disease caused by autosomal recessive loss of function mutations in the NGLY1 gene and is the only known congenital disorder of deglycosylation. Patients suffer from movement disorder, developmental delay, liver dysfunction, and alacrima. NGLY1 removes N-linked glycans from glycoproteins in the cytoplasm and is thought to help clear misfolded glycoproteins from the endoplasmic reticulum (ER) through the ER associated degradation (ERAD) pathway. Despite this, NGLY1’s physiological significance in ERAD is not understood. One way to understand disease pathogenesis is to investigate the effects of genetic variation and modifier genes on disease presentation. Our lab created a Drosophila model of NGLY1 deficiency that faithfully recapitulates several disease phenotypes observed in patients, including movement disorder, seizures, and lethality. We crossed this Drosophila model of NGLY1 deficiency with the Drosophila Genetics Reference Panel (DGRP), a collection of ~200 inbred fly strains with fully sequenced genomes, and scored for proportion of NGLY1 knockdown flies surviving to adulthood. The genetic variation of the DGRP led to a spectrum of lethality ranging from strains that had 100% lethality to 100% viability. A genome wide association study (GWAS) generated a list of 61 candidate modifier genes of NGLY1 deficiency. Nine of these candidate genes encode ER resident proteins, proteins with known ERAD functions, or are involved with protein homeostasis. CG31690 and CG4341 were two independent unlinked hits in the GWAS that are orthologs of a single human gene, TMTC2, which encodes an ER adaptor protein involved with calcium homeostasis. Hrd3 and CG8405, orthologs of SEL1L and TMEM259, were both hits and are known components of ERAD complexes that retrotranslocate misfolded proteins from the ER to the cytoplasm for degradation. We are functionally characterizing several of these candidate modifier genes of NGLY1 deficiency using Drosophila and cell culture models. The study of modifier genes can provide new insights into the etiology of the disease and functions of NGLY1, laying the foundation for the development of personalized therapeutics.

Identifying the genetic links between insomnia and cardiovascular disease using Drosophila models of sleep and cardiac physiology

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Insomnia is a common disorder defined by constant difficulty in falling and/or staying asleep. Studies have shown that insomnia is associated with cardiovascular disease (CVD) where insomnia confers more than a 2-fold increased risk of incident CVD. However, the specific shared causal pathways remain poorly understood, making it difficult to identify new therapeutic targets for insomnia that ameliorate CVD risks. Recently, our collaborators found 57 genome-wide significant genetic loci for insomnia. To identify causal genes under these association peaks and understand the mechanisms linking insomnia with CVD, we use Drosophila melanogaster models, which are well-established model systems for sleep and cardiac studies. We hypothesized that suppressing Drosophila orthologs of causal human insomnia genes will lead to compromised sleep and cardiac function. To assess the role of genes associated with the identified loci on cardiovascular physiology, we obtained RNAi stocks of 72 orthologs of genes under the 57 human genetic association peaks. Knockdown (KD) of these genes was done using the cardiac-specific Hand-Gal4 driver. 1-week-old Drosophila progeny were then used for semi-intact microscopic heart preparation followed by high-speed videography to assess cardiac physiological parameters. Similarly, to assess their role in sleep, KD of these genes was done using the neuron-specific ELAV-Gal4 driver. Sleep and locomotor activity of 3 to 7-day-old flies was monitored using the Drosophila Activity Monitoring System. Interestingly, our initial results show that 3 of 4 genes with fly orthologs, ATPSynC, Bruce and Larsen, within one insomnia genetic locus, show sleep and cardiovascular phenotypes. Mainly, cardiac-specific KD of Larsen led to significant cardiac dilation and reduced cardiac performance, while KD of Bruce led to significantly reduced cardiac performance without significant dilations. Moreover, KD of ATPSynC led to a compromised beating pattern with an elongated diastolic interval and shortened life span. Additionally, neuronal-specific KD of all 3 genes led to sleep fragmentation with only Larsen resulting in a significant decrease in overall sleep which was primarily due to a decrease in nighttime sleep. Assessments of inter organ crosstalk for this locus and measurements of sleep and cardiac parameters of other genetic lines are still in progress. This work will reveal mechanisms linking insomnia and CVD which would help prevent and treat both diseases.

A multi-model system approach identifies genetic interactions underlying atrial fibrillation susceptibility

James Kezos, Anais Kervadec, Alex Colas, Chris Larson, Karen Ocorr Sanford Burnham Prebys Medical Discovery Institute, La Jolla, CA

Atrial fibrillation (AF), the most common heart rhythm disorder, is reaching epidemic proportions in the aging population. In both human and model systems, it is not understood how aging, genetic predispositions, and external environmental factors synergize to promote arrhythmia, nor which gene regulatory networks initiate and maintain AF. Over 200 genetic variants have been associated with increased AF susceptibility, suggesting that the underlying cause is multifactorial, involving networks of interacting genes. Resolving complex interactions modulating cardiac function
in AF is difficult in mammalian systems, but approachable using Drosophila. We utilize a multi-platform approach encompassing the genetically tractable Drosophila cardiac-aging model and hIPSC-atrial-like cardiomyocyte (ACM) model. The fly model allows for high-throughput quantification of aging effects in conjunction with genetic insults, whereas the ACM model allows for high-throughput combinatorial gene knockdown (KD). High-speed imaging of ACMs and fly hearts permit quantification of cardiac parameters such as action potential duration and arrhythmicity in ACMs, as well as contraction intervals and arrhythmicity in flies. Preliminary screens of candidate AF genes in both platforms have identified a network of seven corroborating hits. Network analysis has linked these 7 genes to ion channels, such as Sarcoplasmic (SclA). Single genetic insults rarely produce robust arrhythmicity in our models, but we do see arrhythmia when testing interactions between genes in our network. Aging and pharmacological stressors (isoproterenol in ACMs, octopamine in flies) also interact with genetic insults to cause arrhythmia. For example, KD of Doc in a Sh heterozygote background produces robust arrhythmicity, even at younger ages, whereas knockdown of SclA in a Sh mutant background mitigates cardiac alterations. Our approach has been validated in an atrial-specific double knockout mouse model of Doc and Sh, where atrial P-waves were deleted and QRS complex was broadened, both signatures of AF. Further identification and characterization of components from this Shaker-centric AF network across multiple platforms will not only improve our genetic and molecular understanding of AF pathogenesis but also guide novel experimental and therapeutic strategies to treat AF.

177 Exploring the effects of diet-induced obesity on the invasiveness of Drosophila tumours Cecilia Cabrera1,2, Susumu Hirabayashi1,2 1) Medical Research Council (MRC) London Institute of Medical Sciences, London; 2) Institute of Clinical Sciences (ICS), Faculty of Medicine, Imperial College London, London

Epidemiological studies have shown that obesity promotes the development and progression of various cancers, including colorectal cancers. However, while cancer is a genetically heterogeneous disease, little is known about the genetic characteristics of tumours that associate with obesity. Here, we combine a series of Drosophila models of colorectal cancer with a Drosophila model of diet-induced obesity to explore whether and how tumours of different genetic profiles exhibit different invasiveness in response to obesity.

Using the dissemination of tumour cells into the abdominal cavity as a quantitative readout to assess tumour invasiveness, we demonstrate that obesity modulates the invasiveness of different tumours to different degrees. Furthermore, we found that tumours of different genetic profiles respond to obesity with different directionality; some tumours exhibited increased invasiveness in response to obesity, while other tumours were resistant to the effects of obesity.

RNA-Sequencing and comparative analysis of different tumours revealed Reactive Oxygen Species (ROS) as an important mediator that modulates the invasiveness of tumours in response to obesity. We demonstrate that dietary and genetic manipulation of ROS levels has a functional effect on the invasiveness of tumours.

Our observations indicate that the genetic profiles of tumours have a significant impact on the tumour’s response to obesity. This highlights the importance of stratification of tumours in understanding the relationship between obesity and cancer. Our work also demonstrates the usefulness of Drosophila as a model system to study the connection between obesity, tumour heterogeneity, and tumour invasiveness.

178 The Clot Thickens: Tumor-induced coagulopathy is a conserved driver of host mortality Katy Ong, Tsai-Ching Hsi, David Bilder University of California, Berkeley, Berkeley, CA

Malignant tumors trigger a complex network of inflammatory and wound-healing responses, prompting Dvorak’s characterization of tumors as ‘wounds that never heal’. Some of these responses lead to profound clotting defects displayed by cancer patients, such as Disseminated Intravascular Coagulopathy (DIC), which correlate with poor prognoses. However, evidence suggests that lethality from coagulopathy is not fully explained by hemostatic dysregulation alone, raising the question of alternative mechanisms. Here, we demonstrate that tumor-induced coagulopathy is a paraneoplastic syndrome displayed by Drosophila as well as mammals. Fly tumors overproduce multiple components of the coagulation cascade, including the conserved Factor XIIIa homolog, and tumor-bearing flies exhibit abnormalities in clotting and wound healing that mirror DIC pathology. Critically, depleting coagulation components within the tumor significantly improved host survival, suggesting that coagulopathy is an unappreciated driver of mortality, even in insects that have an open circulatory system. We will discuss the underlying mechanism and the resulting insight into conserved responses of animals to malignancy.

179 Modeling Paraneoplastic Diabetes in Drosophila Jyoti Tripathi, Pradip Sinha Department of Biological Sciences and Bioengineering, Indian Institute of Technology Kanpur

Cancer patients display multiple metabolic disorders induced by humoral factors secreted by the growing cancerous...
mass. Collectively termed as paraneoplastic syndromes, these systemic metabolic disorders often contribute to morbidity and even mortality of the patients: anorexia and cachexia being some of the cited examples. The onset of such paraneoplastic syndromes is marked by poor prognosis and often represents end-stage cancer. In certain cancers with the highest mortality rate, new-onset diabetes has been reported as another paraneoplastic phenomenon. Thus, management of cancer-induced diabetes remains a potential route to alleviate morbidity of cancer patients, improve their prognosis and, in certain instances, facilitate their early diagnosis and management. Here we show the systemic fallouts of oncogenic Ras signaling underlies cancer-linked diabetes in *Drosophila*. We found that a localized Ras activation in a non-vital organ decreases the adult eclosion rate. Further, oncogenic Ras signaling systemically perturbs host metabolism, particularly insulin signaling, with the accompanying onset of obesity, dyslipidemia, hyperglycemia, and, finally, insulin resistance and diabetes. Our results further reveal putative candidate secreted factors from the oncogenic Ras signaling cells as an inducer of these paraneoplastic syndromes. Our findings, therefore, reveal for the first time a causal underpinning of cancer-linked diabetes and the prospect of using *Drosophila* model for the exploration of therapeutic interventions.

180  **Ionizing Radiation induces cells with past caspase activity that contribute to the adult organ in *Drosophila* and show reduced Loss of Heterozygosity** Sarah Colon Plaza, Tin Tin Su University of Colorado Boulder

There is increasing recognition that cells may activate apoptotic caspases but not die, instead displaying various physiologically relevant consequences. We know very little, however, of mechanisms that underlie the life/death decision in a cell that has activated apoptotic caspases. By optimizing a published reporter of past caspase activity, we have been able to visualize such cells that result specifically from exposure to ionizing radiation (IR) in *Drosophila* larval wing discs. We found that cells with X-ray-induced past active caspases (XPAC) do not arise at random but are born at specific locations within the developing wing imaginal discs of *Drosophila* larvae. Furthermore, this reporter has allowed us to gather pre-liminary data on genes that might influence the number of XPAC. Axin, a negative regulator of *wingless* signaling, increases the number of XPAC cells suggesting that *wingless* signaling regulates whether cells with caspase activity live or die. A dosage response curve shows that the number of XPAC peaks and then decreases at higher radiation doses. We also found that heterozygotes of H99 chromosomal deficiency that removes pro-apoptotic genes hid, rpr and grim show reduced number of XPAC cells. Yet, XPAC cells appear in stereotypical patterns that do not follow the pattern of IR-induced apoptosis, suggesting additional controls at play. By following irradiated larvae into adulthood, we found that XPAC cells contribute to the adult wing. By combining a reporter for past caspase activity with mwh, an adult marker for Loss of Heterozygosity (LOH), we addressed the relationship between XPAC and genome stability after irradiation. We find that XPAC cells show reduced LOH relative to the rest of the wing, suggesting a physiological role for non-lethal caspase activity during recovery from radiation damage.

181  **Irradiation-Induced Cell Migration: An Epithelial-Mesenchymal Transition Process Regulated By Low-Level Caspase Activity** Lena Sapochnikov, Eli Arama The Weizmann Institute of Science

Apoptosis, the most common form of programmed cell death in animal development and homeostasis, is mediated by the activation of a unique family of proteases called caspases. However, an ever-growing list of studies suggest that caspases have an essential role in ensuring non-lethal cellular functions during normal development, tissue repair and regeneration, including in cell differentiation, proliferation, migration, signaling and cellular remodeling. Moreover, upon dysregulation, these caspase-dependent non-lethal cellular processes (CDPs) can also instigate disease. We previously discovered and characterized a novel non-apoptotic role of caspases in maintaining epithelial tissue integrity in *Drosophila*. We showed that upon ionizing irradiation or even spontaneously during development, epithelial cells compromised for caspase activity gain high migratory and invasive capacities. Furthermore, low levels of effector caspase activity, far below the threshold required to induce apoptosis, potently inhibit this process, which we termed irradiation-induced cell migration (ICM). Previous work showed that ICM is mediated by the DNA damage response (DDR) pathway, but the molecular mechanisms and signaling pathways underlying ICM downstream of the DDR remained largely elusive. Here we show that ICM shares many features with the process of epithelial-mesenchymal transition (EMT). Inactivation of key EMT-driving genes, which are also potential caspase substrates, including *twist* and *zfh1* (*ZEB1*), attenuates ICM. We further show evidence that the Notch signaling pathway is involved in ICM, and that effector caspases are important regulators of Notch-mediated ICM. Furthermore, members of the ADAM family of metalloproteinases, which among other functions are also responsible for the activation of the Notch signaling pathway, are also involved in ICM and may function as potential extracellular matrix modifiers during this process. Collectively, our findings indicate that ICM is an EMT-like process activated by genotoxic stress and is potently attenuated by non-lethal levels of caspase activity. Given that ionizing irradiation has been a common treatment for many cancers, and that one of the hallmarks of cancer cells is their ability to evade cell death, these findings may have implications for cancer malignancy and the induction/aggravation of metastasis in some patients undergoing radiotherapy.

182  **A genome-wide CRISPR screen identifies DPM1 as a modifier of DPAGT1 deficiency and ER stress** Hans Dalton¹, Raghuvir Viswanatha², Roderick Brathwaite, Jr.², Jeesophia Zuno¹, Stephanie Mohr², Norbert Perrimon³,⁴, Clement Chow¹ 1) Department of Human Genetics, University of Utah School of Medicine, Salt Lake City, UT, USA; 2) Department
Partial loss-of-function mutations in glycosylation pathways underlie a set of rare diseases called Congenital Disorders of Glycosylation (CDGs). Glycosylation is a broad category of sugar modifications on proteins and lipids, with functions ranging from complex post-translational modifications (e.g. N-glycosylation) to single additions involved in cell signaling (e.g. O-GlcNAcylation). CDGs have a range of symptoms, but commonly include severe epilepsy, developmental delay, and disability. CDG Type Ij is caused by loss-of-function mutations in DPAGT1 – the first step in N-glycosylation. Our goal is to better understand the pathways connected to DPAGT1 loss to develop potential treatment options.

We performed a CRISPR knockout screen using the drug tunicamycin (Tun), a potent inhibitor of DPAGT1 function, on Drosophila S2R+ cells. Loss of DPAGT1 impairs N-glycosylation and causes massive protein misfolding, leading to reduction of cell surface glycoproteins and endoplasmic reticulum (ER) stress. We introduced a whole genome guide RNA library into S2R+ cells expressing constitutive Cas9. Cells were grown under either vehicle or Tun selection. Final populations were sequenced to determine candidate genes causing resistance or sensitivity to DPAGT1 inhibition. Gene Ontology of these genes related to the hexosamine pathway and glycolysis, two pathways important for N-glycosylation. We also performed a Concanavalin A (ConA) screen to assay cell surface glycoproteins in an identical pool of cells. This assay found loss of genes in GPI anchor biosynthesis (e.g. PIG-A and PIG-H) could rescue cell surface glycoproteins under DPAGT1 inhibition.

We created an in vivo Drosophila DPAGT1 model where Alg7/DPAGT1 RNAi is driven in the eye to cause a degraded eye phenotype. Using this and another ER stress model, we validated candidate genes via RNAi knockdown. RNAi in these models revealed that loss of the mannosyltransferase Dpm1, involved in mannnose addition in all downstream glycosylation pathways, could strongly rescue DPAGT1 inhibition and ER stress phenotypes. Testing its downstream pathways (O- and C-mannosylation [rt, CG6659], N-glycosylation [Alg3], GPI biosynthesis [PIG-M]), we found loss of O-mannosylation, N-glycosylation, and GPI biosynthesis partially recapitulates the Dpm1 RNAi rescue of DPAGT1 inhibition and ER stress. These findings suggest impairment of Dpm1, or its downstream pathways, as key targets for potential therapeutics for DPAGT1 loss.

183 The Protein Phosphatase-1 regulatory subunit dPPP1R15 controls collective cell migration via the eIF2-alpha-ATF4-dependent ER stress pathway Yujun Chen, Jocelyn McDonald Kansas State University

Collective cell migration is essential for many developmental and pathological processes. Drosophila border cells migrate as a cohesive cluster during oogenesis and provide an excellent genetic model to elucidate conserved mechanisms of collective cell movement. Previously, we discovered that Protein Phosphatase 1 (Pp1) catalytic subunits are critical molecular regulators of border cell collective versus single cell behaviors. We now show that the conserved Pp1 regulatory subunit dPPP1R15 (also known as GADD34) plays a crucial role in border cell migration. Overexpression of dPPP1R15 causes border cells to round up and completely dissociate from the cluster during migration. These defects are rescued by overexpression of Pp1 catalytic subunits (Pp1c). Moreover, overexpressing PPP1R15 mutants that cannot bind to Pp1c no longer show migration or cluster-dissociation phenotypes. These data together indicate that dPPP1R15 functions through Pp1c in border cells. In contrast, RNAi-mediated knockdown of dPPP1R15 prevents border cell delamination from the follicular epithelium. Live imaging reveals that, compared to control border cells, dPPP1R15-RNAi border cells also extend much smaller protrusions that are misdirected, thus contributing to the failure of border cells to migrate. The Pp1c/PPP1R15 complex is best known as a phosphatase complex that dephosphorylates eIF2alpha in the endoplasmic reticulum (ER) stress pathway. Indeed, loss of dPPP1R15 elevates phosphorylation and activation of eIF2alpha and increases expression of the known downstream transcription factor, ATF4, in border cells. In addition, upregulation of PERK, a major eIF2alpha kinase, blocks border cell delamination and migration. Interestingly, depleting PERK by RNAi also delays border cell migration. Overexpression of a non-phosphorylatable eIF2alpha mutant, or knockdown of ATF4, significantly rescues PPP1R15-RNAi induced migration defects. Overexpression of a phosphomimetic eIF2alpha mutant, or of ATF4, impairs migration. Together, our data support the idea that restraining the PERK-eIF2alpha-ATF4 ER stress pathway via the dPPP1R15 phosphatase is critical for proper border cell migration. Our studies identify a new role for dPPP1R15 as a key regulator of collective cell behaviors through modulation of the ER stress response.

184 Increased intracellular pH promotes cell death in the developing Drosophila eye Joanne Mendez, Juan Pacheco, Daniel Orozco, Bree Grillo-Hill San Jose State University

Constitutively increased intracellular pH (pHi) is common to most cancers regardless of their tissue of origin or genetic background. Cancer research has traditionally focused on cancer-associated mutations and dysregulated signaling pathways. However much less is known about how changes in cellular chemistry, including pHi, regulate cancer cell behaviors. Our lab developed tools to increase pHi in the absence of other transforming mutations by overexpressing the Drosophila sodium-proton exchanger, DNhe2, in the Drosophila eye (GMR>DNhe2). We have found that GMR>DNhe2 flies have a rough eye phenotype in adult flies. To determine the underlying cause of this rough eye,
we performed phenotypic analysis in larval and pupal eyes. We found the GMR>DNhe2 flies have increased proliferation in larval eyes discs. Paradoxically, we found fewer interommatidial cells at the end of patterning in pupal eyes (11.4 compared to 15 cells per counting area in control). One aim of our current work is to perform a time course analysis of cell number in GMR>DNhe2 flies to determine when in development these cells die, and to determine the mode of cell death. We next tested for genetic interactions between DNhe2 and cell death genes. We found that the pH- dependent cell death is p53-dependent but caspase independent, which is inconsistent with apoptosis, but suggests autophagy. Second, we want to identify the genes that mediate this increased cell death at higher pH. A dominant modifier screen identified overexpression of the oncogene Myc as a strong suppressor of the GMR>DNhe2 rough eye. We tested whether this suppression is due to Myc attenuating the hyperproliferation effects of over-expressing DNhe2, but we saw no effect. Currently, we are quantifying the effects of co-expressing Myc and DNhe2 on cell number in pupal eyes. Together, our findings will elucidate mechanisms for pH-regulation of conserved, critical developmental processes and provide evidence for new paradigms in growth control.

185 Lamp1 mediates lipid transport, but is dispensable for autophagy in Drosophila Norin Chaudhry1, Margaux Sica2, Satya Surabhi3, David Sanchez Hernandez2, Ana Mesquita2, Adem Selimovic1, Ayesha Riaz2, Laury Lescat2, Hua Bai1, Gustavo C. MacIntosh3, Andreas Jenny2 1) Iowa State University, Ames, IA, USA; 2) Albert Einstein College of Medicine, New York, NY, USA

The endolysosomal system not only is an integral part of the cellular catabolic machinery that processes and recycles nutrients for synthesis of biomaterials, but also acts as signaling hub to sense and coordinate the energy state of cells with growth and differentiation. Lysosomal dysfunction adversely influences vesicular transport-dependent macromolecular degradation and thus causes serious problems for human health. In mammalian cells, loss of the lysosome associated membrane proteins LAMP1/2 strongly impacts autophagy and cholesterol trafficking. Here we show that the previously uncharacterized Drosophila Lamp1 is a bona fide ortholog of vertebrate LAMP1/2. Surprisingly and in contrast to Lamp1/2 double mutant mice, Drosophila Lamp1 is not required for viability or autophagy, suggesting that fly and vertebrate LAMP proteins acquired distinct functions, or that autophagy defects in Lamp1/2 mutants may have indirect causes. However, Lamp1 deficiency results in an expansion of the acidic compartment in flies. Furthermore, we find that Lamp1 mutant larvae have defects in lipid metabolism as they show elevated levels of sterols and diacylglycerols (DAGs). Since DAGs are the main lipid species used for transport through the hemolymph (blood) in insects, our results indicate broader functions of Lamp1 in lipid transport. Our findings make Drosophila an ideal model to study the role of LAMP proteins in lipid assimilation without the confounding effects of their storage and without interfering with autophagic processes.

192 Enabling recombination on the 4th chromosome: FRT101F and Bloom syndrome helicase Stuart Newfeld1, Mary Jane O’Connor2, Samuel Goldsmith1, Petra Tauscher1, Samantha Daly1, Osamu Shimm14, Michael O’Connor2 1) School of Life Sciences, Arizona State University, Tempe AZ; 2) Dept. Genetics, Cell Biology and Development, Univ. Minnesota, Minneapolis, MN; 3) Institute of Biotechnology, University of Helsinki, Helsinki, Finland; 4) Institute of Molecular and Cell Biology, University of Tartu, Tartu, Estonia

Genes on the long arm of the Drosophila 4th chromosome are difficult to study because the chromosome lacks mitotic and meiotic recombination. Without recombination it is not possible to employ clonal analysis – the creation of a single mutant cell at a specific time and place followed by the phenotypic examination of its descendants. Here we report new resources for the 4th chromosome. For mitotic recombination we generated a chromosome with an FRT very near the centromere in 101F and a derivative that carries FRT101F with a distal ubiquitously expressed GAL80 transgene. This pair of chromosomes enables both unmarked and MARCM clones. An example from the IPC neurons of the brain is shown. For meiotic recombination we demonstrate that a Bloom syndrome helicase and recombination defective double mutant genotype can create recombinant 4th chromosomes via female meiosis. An example creating a w+ and y+ chromosome is shown. An efficient method for X to 4th transposition is also described. These chromosomes and strategies should accelerate the genetic analysis of protein coding genes on the 4th chromosome, including the 44 genes with no demonstrated function. Studies of these well conserved genes will close gaps in our knowledge of development and physiology.

193 Temperature-Inducible precision guided Sterile Insect Technique, Tl-pgSIT Nikolay Kandul, Junru Liu, Omar Akbari Division of Biological Sciences, Section of Cell and Developmental Biology, UC San Diego, La Jolla, California, 92092 USA

Releases of sterile males are the gold standard for many insect population control programs, and precise sex sorting to remove females before male releases is essential to the success of these operations. To advance traditional methods for scaling the generation of sterile, fit, and competitive males, we previously described a CRISPR-mediated precision-guided Sterile Insect Technique (pgSIT). PgSIT functions by exploiting the precision and accuracy of CRISPR to simultaneously disrupt genes essential for female viability and male fertility. It utilizes a simple breeding scheme requiring two strains: one expressing Cas9 and the other expressing multiple guide RNAs (gRNAs). A single mating between these strains
mechanistically results in synchronous Cas9/RNA-mediated dominant biallelic knockouts of target genes throughout development, resulting in the complete penetrance of desired phenotypes in all progeny. This two-locus pgSIT design was used to engineer pgSIT systems in Drosophila melanogaster, Drosophila suzukii, a global pest of economically important soft-skinned fruits, and Aedes aegypti, the principal vector for arboviruses including dengue/yellow fever, chikungunya, and Zika virus. While effective at generating F1 sterile males, the two-locus pgSIT depends on a genetic cross between the two parental strains which requires maintenance and sexing of two strains in a factory. Therefore, to further advance pgSIT by removing this crossing step, we describe a one-locus Temperature-inducible pgSIT (TI-pgSIT) technology and demonstrate its proof-of-concept in D.melanogaster. Importantly, we were able to develop a true-breeding strain for TI-pgSIT that eliminates the requirement for sex sorting, a feature that may help further automate production at scale.

194 Seamless genetic engineering via CRISPR-triggered SSA allows spatio-temporal control of gene labelling Gustavo Aguilar, Milena Bauer, Alessandra Vigano, Carlos Jiménez-Jiménez, Isabel Guerrero, Markus Affolter 1) University of Basel, Switzerland; 2) Centro de Biología Molecular Severo Ochoa (CSIC-UAM), Spain

Precise genome engineering is essential for both basic and applied research, permitting the manipulation of genes and gene products in predictable ways. The improvement of CRISPR/Cas technology rocketed the speed and ease by which exogenous sequences are integrated into specific loci. To this day, a number of strategies permit gene manipulation. Nevertheless, knock-in generation in multicellular animals remains challenging, partially due to the complexity of insertion screening. Even when achieved, the analysis of protein localization can still be unfeasible in highly packed tissues, where spatial control of gene labelling would be ideal. Here, we propose a method based on Homologous Directed Repair (HDR) and Single Stranded Annealing (SSA) repair pathways. HDR mediates the integration of a switchable cassette. Upon a subsequent CRISPR-triggered repair event, resolved by SSA, the cassette is seamlessly removed. We named the technique SEED/harvest. By engineering the Hedgehog pathway components, we demonstrated fast and robust knock-in generation with both fluorescent proteins and short protein tags in tandem. The implementation of short homology arms, further simplified and cheapen the process. Seamless knock-in generation can be achieved in as fast as one and a half months.

The use of SEED cassettes is not restricted the germ line. SSA can also be triggered in somatic cells, permitting conditional gene labelling. This is the first time CRISPR is used to trigger endogenous tissue-specific gene labelling. Since the technology is solely based on CRISPR, it can be easily used in combination with other tissue-specific CRISPR tools, as membrane labelling and generation of knock-outs. We provide evidence of cell-type specific endogenous labeling as well as temporal control of gene labeling, mediated by conditional expression of guideRNAs.

While our tools permit control of protein labeling with fluorescent proteins, the in vivo visualization and manipulation of proteins with short tags in tandem still represents a challenge. To complement the SEED technology, we have developed a toolkit based on rational nanobody engineering and functionalization that permits the visualization and manipulation of proteins tagged with short tags.

195 SpyChIP identifies genome-wide and cell type-specific transcription factor occupancy Siqian Feng, Richard Mann Jerome L. Greene Science Center, Columbia University, New York, NY

A major drawback of chromatin immunoprecipitation (ChIP) based techniques is the lack of cell type-specificity. To overcome this limitation, we developed SpyChIP to identify sites of cell type-specific transcription factor (TF) occupancy in native physiological contexts without tissue dissociation or nuclei sorting. SpyChIP takes advantage of a specific covalent isopeptide bond that rapidly forms between the 15 amino acid SpyTag and its binding partner, the 17 kD protein SpyCatcher. In SpyChIP, the target TF is fused with SpyTag by genome engineering, and an epitope tagged SpyCatcher is expressed by the Gal4/UAS system in cell populations of interest, where it covalently links to SpyTag-TF. Cell type-specific ChIP results are obtained by performing ChIP against the epitope on SpyCatcher using chromatin prepared from whole tissue.

Using SpyChIP, we characterized the genome-wide binding profiles of the Hox protein Ultrabithorax (Ubx) in two non-overlapping domains of the Drosophila haltere imaginal disc. Our results revealed extensive region-specific Ubx-DNA binding events, thus highlighting the significance of cell type-specific ChIP results and the limitations of whole tissue ChIP approaches. Analysis of SpyChIP results for Ubx provided novel insights into the relationship between chromatin accessibility and Ubx-DNA binding, as well as different mechanisms Ubx employs to regulate different downstream cis-regulatory modules (CRMs). We envision that the covalent bond between SpyTag and SpyCatcher will have widespread in vivo applications, and our demonstration of SpyChIP sets the stage for carrying out many other cell type-specific characterizations and manipulations in vivo that were previously unachievable.

196 The continuum of Drosophila embryonic development at single cell resolution Xingfan Huang, Ronnie Blecher-Gonen, Diego Calderon, Riza Daza, Stefano Secchia, Beth Martin, Baekgyu Kim, Alessandro Dulja, Cole Trapnell, Eileen Furlong, Jay Shendure 1) The Crown Genomics institute of the Nancy and Stephen Grand Israel National Center for Personalized Medicine, Weizmann Institute of Science, Israel; 2) Department of Genome Sciences,
Single cell technologies are a powerful new means to study metazoan development, enabling comprehensive surveys of cellular diversity at profiled timepoints, and shedding light on the dynamics of regulatory element activity and gene expression changes during the in vivo emergence of each cell type. However, nearly all such atlases of embryogenesis remain limited by sampling density, i.e. the number of discrete time points at which individual embryos are harvested. Given the rapidity with which molecular and cellular programs unfold, this limits the resolution at which regulatory transitions can be characterized.

To construct a continuous representation of embryogenesis in vivo, we would ideally sample embryos continuously. Although not possible with most model organisms, it is potentially possible in Drosophila melanogaster, where collections of timed and yet somewhat asynchronous embryos are easy to obtain. Drosophila could therefore serve as a test case to develop a framework for the inference of continuous regulatory and cellular trajectories as embryogenesis progresses. Of course, as Drosophila is a preeminent model organism that has yielded many advances in the biological and biomedical sciences, obtaining a single cell atlas of Drosophila embryogenesis is also an important goal in itself.

In this study, we report a continuous, single cell atlas of chromatin accessibility and gene expression that spans Drosophila embryogenesis. We profiled chromatin accessibility in almost one million, and gene expression in half a million, nuclei from eleven tightly staged, overlapping windows of 0 through 20 hours of embryogenesis. Leveraging the asynchronicity of embryos within each collection window, we developed a statistical model to estimate the age of each nucleus more precisely, resulting in continuous views of molecular and cellular transitions throughout embryonic development. From these data, we identify cell types, infer their developmental relationships, and link cell type-specific changes in transcription factor expression to changes in the accessibility of their cognate motifs. Looking forward, this strategy may facilitate future investigations of in vivo gene regulation throughout Drosophila embryogenesis at arbitrarily high temporal resolution.

**Optogenetic manipulation of endogenous proteins in Drosophila by light-inducible trapping** Yineng Xu, Bei Wang, Chun han Weill Institute for Cell and Molecular Biology and Department of Molecular Biology and Genetics, Cornell University

During animal development, the same molecular event can result in drastically different consequences depending on when and where it occurs. Thus, the spatial and temporal properties of signaling inputs and gene activity largely determine the biological outcomes. Spatiotemporal regulation is particularly important for neuronal development and function: Spatially, neurons occupy broad domains and may incur distinct signaling events at different cellular compartments as a result of unique interactions with the surrounding microenvironment. Temporally, neurons develop through distinct stages and require series of molecular events that occur in particular sequences. However, widely used methods for studying gene function, such as mutations, RNAi, and CRISPR, lack the spatiotemporal regulation required for dissecting fine developmental mechanisms. In contrast, light-controllable modules hold great promise for precise and instant manipulation of molecular events in cells. Towards this goal, we developed OptoTrap, a light-dependent protein trapping system, which builds on two light controllable systems—Cry2olig and magnets. This system forms large protein aggregates under blue light and can trap proteins that are tagged by GFP or split GFP. We generated several variants of this system for diverse applications in neurons and epithelial cells and characterized their association and dissociation properties. We demonstrate that this system can effectively trap GFP-tagged endogenous proteins of diverse sizes, subcellular locations, and functions. Functionally, light-dependent protein trapping of the septate junction protein Nrg in epithelial cells caused gain-of-function (GOF) and cell shrinkage, suggesting Nrg activation through clustering. In contrast, optogenetic trapping of kinesin heav chain (Khc) in somatosensory neurons caused effective loss-of-function (LOF), resulting in disruption of microtubule-dependent transport and dendrite reduction. Similarly, trapping of Nmnat in neurons caused dendrite reduction, likely due to disruptions of Nmnat’s chaperon function. OptoTrap allows us to fine tune protein manipulation for graded phenotypes by changing illumination conditions, and thus offers great flexibility for dissecting protein functions in neuronal development. This toolkit should be applicable to broader developmental stages and diverse cell types for studying development mechanisms.

**Spying on the dynamics of neuropeptides by the GRAB sensors in Drosophila xiju xia** Yulong Li, 1,2,3, Yulong Li, 1,2,3,4, 1) State Key Laboratory of Membrane Biology, Peking University School of Life Sciences, Beijing, China; 2) PKU-IDG/ McGovern Institute for Brain Research, Beijing, China; 3) Peking-Tsinghua Center for Life Sciences, Academy for Advanced Interdisciplinary Studies, Peking University, Beijing, China; 4) Chinese Institute for Brain Research, Beijing, China

Neuropeptides are essential signaling molecules transported and secreted by large dense-core vesicles (DCVs). Neuropeptides play a critical role in many physiological processes, including brain development, sleep, circadian rhythm,
and feeding behaviors, alterations in neuropeptide levels have been associated to many pathological states, such as anxiety, stress and addiction related disorders. For a precise understanding of neuropeptides’ functions, it is important to probe when, where and how they are released in brain. Here, we developed a series of genetically-encoded green GRAB (GPCR activation based) sensors for detecting neuropeptides in living flies, including sNPF, CCha1, DH31 and FMRFa. These peptide sensors exhibited nanomolar to micromolar EC\textsubscript{50} affinity and exquisite selectivity for cognate peptide ligands. Preliminary results suggested that transgenic expressing sNPF sensors are capable of detecting in vivo peptide release events triggered by high K\textsuperscript+ and optogenetic stimulation. Furthermore, calyx and horizontal lobe of mushroom body neurons (Kenyon cells) shows distinct release probabilities. Thus, these GRAB sensors provide sensitive and specific molecular probes to unravel the in vivo dynamics and molecular mechanisms of neuropeptide release in Drosophila.

**Key words**: neuropeptides, GRAB sensors, sNPF

199 **Cryopreservation method for Drosophila melanogaster embryos** Li Zhan, Min-gang Li, Amanda Neisch, Thomas Hays, John Bischof University of Minnesota, Minneapolis, MN

The fruit fly (Drosophila melanogaster), a foundational genetic model organism in the past century, has driven important biomedical science breakthroughs including six Nobel Prizes. However, more than 160,000 unique genotypes worldwide must be manually maintained through frequent and costly transfer of breeding adults to fresh food. In comparison, cryopreservation provides enormous advantages including protection against genetic drift, decreasing stock maintenance costs, and reducing the risk of stock loss caused by contamination or accidental mixing of precious stocks. Here we report an easily implemented and robust cryopreservation protocol of *Drosophila melanogaster* embryos, demonstrating potential for wide adoption. We present innovations for embryo permeabilization, cryoprotectant agent (i.e., anti-freezer) loading, and rewarining. After embryo collection, our protocol takes ~35 min to freeze and store the embryos in liquid nitrogen, ~25 min to thaw the embryo and remove the intra-embryonic cryoprotectant agent. We showcase its implementation in 25 distinct strains including wildtype and mutants. Using this protocol, >50% embryos hatch and >25% of the resulting larvae develop into adults (i.e., survival normalized to untreated embryos) after cryopreservation for most strains. For strains with initially low survival after cryopreservation, we show that the survival can be substantially improved by outcrossing to mitigate the effect of genetic background. After successive cryopreservation of 5 generations and 12-month storage in liquid nitrogen, the flies retain normal sex ratio, fertility and original mutation. Importantly, we find that non-specialists are able to use this protocol to obtain consistent results after training.

201 **Minute mutations, cell competition, and cellular surveillance** Nick Baker Albert Einstein College of Medicine

I will be describing my lab’s work on cell competition and its physiological functions in Drosophila. Cell competition is the elimination of cells within an organ that only occurs when other cells are available to replace them. The term was originally coined to describe the elimination of ‘Minute’ (ie Rp/+ cells from mosaic imaginal discs containing wild type cells, which are progressively lost even though entirely Rp/+ animals are viable. Cell competition was originally envisaged as a homeostatic process responding to variations in cellular growth rate, but is now thought to provide a selection against abnormal cells that may arise in developing tissues. We used genetic screens to isolate mutations preventing cell competition. Analyzing these mutations changed how we think about the Minute phenotype, which is now understood mostly to reflect a transcriptional response mediated by Xrp1, the key transcription factor that is expressed in Minute mutants. Remarkably, mutating this single transcription factor prevents cell competition and even restores normal overall translation to Minute cells. Minutes reduce translation though Xrp1-dependent phosphorylation of eIF2a by PERK, a kinase that also responds to ER stress, and not by reduction in ribosome numbers. Cell competition mutations do not affect organ size or reproducibility, instead they prevent certain kinds of abnormal cells from being eliminated. For example, aneuploid cells can be eliminated by cell competition on the basis of their abnormal Rp gene dose. Xrp1 seems also to be a common target of other cellular defects that trigger cell competition, including defects in ubiquitylation and protein turnover. Curiously, Xrp1 is a very rapidly evolving gene, which suggests it is involved in some evolutionary arms race, eg with pathogens. Interestingly, Xrp1 was first identified as a transcriptional target of p53 in the DNA damage response. Consistent with this, cells with different p53 activity levels compete with one another in imaginal discs. Notably, in mammalian cells Rp mutations lead to p53 activation, and there are multiple examples where different p53 activity levels lead to competition between mammalian cells. This suggests that aspects of mammalian p53 function may have been acquired by a p53 target gene in *Drosophila*, and leads us to speculate about how cell competition might contribute to cancer surveillance and tumor suppression.

202 **Innate immune signaling sculpts neuron-glia interactions across lifespan** Heather Broihier Case Western Reserve University, Cleveland, OH

Abstract is not available at the time of print.

203 **Coping with mechanical stress: tissue dynamics in homeostasis and repair** Yanlan Mao University College London
During growth and development, tissue dynamics, such as tissue folding, cell intercalations and oriented cell divisions, are critical for shaping tissues and organs. However, less is known about how tissues regulate their dynamics during tissue homeostasis and repair, to maintain their shape after development and upon wounding. In this talk we will discuss how tissues respond to mechanical perturbations, such as stretching or wounding, by altering their actomyosin contractile structures, to change tissue dynamics and cell shape, and thus preserve tissue shape and patterning. We will present new data on how changes in the 3D shape of cells upon wounding promotes efficient wound repair. We combine genetics, biophysics and computational modelling to study these processes.

204 The evolution of morphological novelties at the cellular and gene regulatory levels Mark Rebeiz1, Shodja Donya2, Rice Gavin3, Vincent Ben4, Smith Sarah1,2, McQueen Eden1,3 1) University of Pittsburgh; 2) present address, Yale University; 3) present address, University of Michigan

The evolutionary origins of complex anatomical structures such as the eye or wing remain a major puzzle in evolutionary developmental biology. The development of morphology is controlled by gene regulatory networks (GRNs) composed of transcription factors, signaling pathways, and the regulatory sequences (enhancers) they control to regulate expression of structural genes that ultimately confer physical properties upon a tissue. To understand how new morphologies evolve, we strive determine the structure of their GRNs, localize evolutionary changes within these GRNs, and unravel how the network governs cellular behavior. I will present our work on the GRNs and cellular processes underlying new morphological structures in the genitalia of Drosophila melanogaster and its close relatives.

205 What long-term quantitative Imaging of stem-cells in their natural environment can tell us about the way they are born, differentiate, and talk to eachother Guy Tanentzapf University of British Columbia, Canada

Abstract is not available at the time of print.

206 Becoming an oocyte: demise of the germ cell program and new beginnings Prashanth Rangan1, Kahini Sarkar2, Noor Kotb 2, Shane Breznak2 1) Icahn School of Medicine at Mount Sinai; 2) University at Albany, SUNY

The continuity of sexually reproducing organisms depends on germ cells. Differentiating germ cells undergo meiosis to give rise to oocytes in females, which upon fertilization launches the next generation. Once the oocyte fate is specified, oocytes synthesize mRNAs and proteins, called the maternal contribution, that regulate development of the early embryo. My lab utilizes the Drosophila female germline to ask the following questions: How is germ cell entry into meiosis controlled? And what mechanisms eliminate the mitotic germ cell-specific programs upon acquisition of oocyte fate? My lab has discovered that production of viable oocytes depends on a stepwise progression through previously unidentified programmatic steps. First, that coordination of female germ cell differentiation and meiotic entry is governed by the male-specific lethal (Msl) complex member, Msl3, which licenses the expression of genes required for these programs. Further, we found that once germ cells have differentiated and committed to meiosis-I, genes critical for promoting differentiation are transcriptionally silenced by a feedback loop between heterochromatin and the Nucleopore complex. In addition, a cohort of perduring RNAs that promote germ cell differentiation and meiosis-I are removed by the No-Go Decay RNA surveillance pathway members. Finally, loss of transcriptional silencing or failure to remove perduring RNAs that promote differentiation and meiosis-I disrupts acquisition of the maternal contribution and causes loss of the oocyte fate. I will share our recent findings on the mechanisms regulating translation of meiotic proteins and the trigger that initiates silencing of the germ cell program.

207 Pioneers, settlers, and life on the OregonR trail: Transcriptional regulation during development Melissa Harrison, Meghan Freund, Marissa Gaskill, Tyler Gibson, Sam Krabbenhoft, Elizabeth Larson, Audrey Marsh, Eliana Torres-Zelada, Katherine Vietor University of Wisconsin-Madison

Coordinated changes in gene expression allow a single fertilized oocyte to develop into a complex multicellular organism. These changes in expression are controlled by transcription factors that gain access to discrete elements in the genome, allowing them to activate gene expression. While nucleosomes act as a barrier to transcription-factor occupancy, pioneer transcription factors have specialized functions that enable them to access binding sites within nucleosomes, establish accessible chromatin and facilitate binding of additional factors. My lab is particularly focused on how these pioneer factors act at the top of gene regulatory networks to control cell fate, including early embryonic reprogramming and stem-cell maintenance. By studying conserved developmental transitions using the powerful toolbox available for studies in Drosophila, we have begun to uncover shared properties of these pioneer transcription factors and identify barriers to their reprogramming functions. Our investigation of the pioneer factors Zelda, GAGA factor (GAF) and Grainy head and their roles in diverse tissues provides fundamental insights into how cell-fate transitions are controlled over development and how, when mis-regulated, can lead to disease.

208 Temporally dynamic antagonism between transcription and chromatin compaction controls stochastic photoreceptor specification Robert Johnston JOHNS HOPKINS UNIVERSITY
Nervous systems are extremely complex, with a myriad of neuronal subtypes defined by distinct functions, morphologies, and connectivities. Neuronal fate specification is driven by lineage, signaling, and stochastic regulatory inputs. The mechanisms controlling stochastic fate specification, in which a cell randomly chooses between two or more fates, are poorly understood. Studying the Drosophila retina, we characterized how natural variation in transcriptional regulation alters stochastic photoreceptor specification and color perception. We determined a mechanism in which simple interactions between transcription and chromatin compaction underlie this stochastic cell fate choice. Our findings suggest fundamental mechanistic principles underlying stochastic fate specification across species. Our work is revealing how cells harness molecular noise to control cell fate decisions in a metazoan.

209A Identifying potential caspase substrates involved in spermatid terminal differentiation in Drosophila Tsili Braun, Shmuel Pietrokovski, Eli Arama Department of Molecular Genetics, Weizmann Institute of Science, Rehovot, Israel

Caspases are the executioners of apoptotic cell death, but these proteases also promote a variety of non-lethal cellular processes. The requirement of caspase activity for spermatid terminal differentiation in Drosophila, a process also known as spermatid individualization, as well as how these cells avoid death in the presence of active caspases, has been well established by our group and others in the past two decades. However, the precise role of caspases during the process of spermatid individualization is still unknown. Moreover, although dozens of caspase-dependent non-lethal cellular processes (CDPs) have been characterized so far, identifying the precise functions of caspases during these cellular processes remained largely elusive and has been highly challenging. Perhaps the main reason for the slow progress has been the need to identify specific caspase protein substrates and the consequence of their cleavage; in particular that caspase activity is restricted during CDPs and thus might only partially cleave some substrates. To start addressing this question in spermatids, we first compiled a list of about 2,000 potential caspases substrates, mainly identified in several large proteomics studies of mammalian cultured cells undergoing apoptosis. We identified the Drosophila orthologs of these proteins and focused on those that are known to mediate cytoskeletal dynamics, a main feature of spermatid individualization, and in which the caspase cleavage site has been conserved. We then downregulated these genes in the male germ cells both in a wild type background and in the background of downregulation of the main effector caspase Drice. We then subjected the male flies to sterility tests. Of those, we mainly focused on the genes that produced synthetic sterility when downregulated together with drice, and which exhibited grossly normal spermatid maturation but failed to individualize. This approach yielded several potential caspase substrate candidates, which are now being further studied in depth for their roles during spermatid individualization.

210B Non-apoptotic activation of Drosophila Caspase-2/9 limits the growth of open-wound-like tumours by modulating JNK signalling and the tumour microenvironment Luis Alberto Lopez1, Derek Xu1, Kenneth Yamada2 1) University of Oxford; 2) National Institute of Dental and Craniofacial Research, NIH.

Personalised cancer therapy requires a deep molecular understanding of tumour features and the interplay between transformed cells and the immunological microenvironment. We have capitalized on a clinically relevant Drosophila tumour model, which simultaneously upregulates the EGFR and JAK/STAT signalling pathways (EJS tumours), to investigate the role of caspases in regulating tumour properties. Our results indicate that widespread non-apoptotic activation of initiator caspases limits JNK signalling and ensures cell fate commitment in EJS tumours. Additionally, caspase activation lowers the remote recruitment and in situ proliferation of EJS tumour-associated macrophages that fuel tumour growth. These findings uncover unrecognised tumour-suppressor activities of caspases that prevent exacerbation of cellular malignancy and tumour overgrowth without inducing apoptosis. Furthermore, they encourage the exploration of caspase-based therapeutic approaches against EGFR/JAK-STAT-activating tumours.

211C Differential sensitivity to cell death cues in long-lived, non-regenerative cells in the Drosophila hindgut Jessica Sawyer, Ruth Montague, Dongwon Lee, Don Fox Duke University

Quiescent long-lived cells often resist many types of cellular insults, which is an advantage in tissues with limited regenerative capacity. However, the molecular underpinnings that promote cellular longevity within a tissue are still largely unclear. We have established the adult hindgut of Drosophila melanogaster as a model to understand responses to cell death in a long-lived, low cell turnover organ. The Drosophila hindgut is comprised of three distinct segments: the diploid pylorus, the polyploid ileum, and the polyploid rectum. We have previously shown that ectopic expression of the pro-apoptotic genes hid and rpr causes cell death and regenerative hypertrophy in the pylorus. Here, we show that expression of hid and/or rpr does not activate a caspase sensor nor lead to cell death in the adult hindgut ileum or rectum. Polyploidy does not protect against cell death cues in these cell types, as ectopically expressing the caspase dronc (Caspase-9), often a downstream hid/rpr target, results in polyploid cell death. Surprisingly, the pylorus mostly resists cell death following dronc overexpression, which suggests that dronc expression alone is insufficient to kill pyloric cells. However, cell death in the pylorus depends on the canonical hid/rpr/dronc cell death pathway, as reducing the amount of dronc or increasing the levels of the dronc inhibitor diap1 reduces hid-mediated cell death and regeneration in the pylorus. Collectively, these data suggest that canonical death signaling is active in the regenerative
pylorus, but distinct sensitivities to specific nodes of apoptotic regulation are present in each hindgut segment. We then compared our results to apoptotic gene expression in two other regenerative tissues: the larval eye and the adult midgut (both polyplody enterocytes and diploid enteroblasts). Both of these tissues behave like the regenerative pylorus. Specifically, hid/rpr overexpression robustly kills eye and midgut cells, while dronc overexpression does not. These data further underscore the distinct cell death signaling responses in the long-lived hindgut ileum and rectum. Finally, we find that the long-lived, hid/rpr-insensitive ileum and rectum lack a robust regenerative response to dronc killing, which may explain their distinct apoptotic responses. Our results argue that the Drosophila hindgut is an excellent model to reveal unique apoptotic regulation of long-lived epithelial tissues.

212A  Knockdown of CG6191 (Mary Shelley) results in compensatory apoptosis in the imaginal wing disc mediated through JNK signaling  Jacob Kagey1, Razan El Yaman1,2, Ali Zamati2 1) Biology Department, University of Detroit Mercy; 2) ReBUILDetroit, University of Detroit Mercy

Drosophila share developmental pathways with humans and other organisms, and many of these pathways are disrupted in human tumorigenesis. We are studying the developmental expression patterns and knock-down phenotypes of the Drosophila gene CG6191, which we have named, Mary Shelley (MS). The human homolog of MS is Cables-1 and has been shown to be frequently downregulated in a number of human cancers, such as ovarian and endometrial. Using the MiMIC system, we find that MS expresses in a distinct band along the dorsal/ventral boundary compartment of the wing disc during larval development. The expression of MS is similar to the well-established expression pattern of the Notch and Wingless cell signaling pathway in the wing. To understand the developmental role that MS is playing in wing development, we utilized RNAi knockdown in the posterior compartment. We find that despite the RNAi confined to the posterior compartment, both the posterior and anterior compartments have a reduction in tissue size. This is due to increased apoptosis at the larval stage both autonomously (posterior) and non-autonomously (anterior). This pattern of non-autonomous apoptosis has previously been shown to be facilitated by the JNK signaling pathway. In MS knockdown cells we see an autonomous increase in the expression of the JNK ligand Eiger and a non-autonomous increase in Puckered, suggesting that the non-autonomous apoptosis is driven by aberrant JNK signaling. Currently we are utilizing a dominant-negative Basket allele to determine if the non-autonomous apoptosis observed is dependent upon JNK signaling. Understanding the downstream consequences of MS knockdown will help to better elucidate the developmental role of MS in Drosophila development.

212B  Ribosome protein mutant cells rely on the GR64 cluster of gustatory receptors for survival and proteostasis in Drosophila  Michael Baumgartner1,3, Alex Mastrogiannopoulos1, Iwo Kucinski2,4, Eugenia Piddini1 1) School of Cellular and Molecular Medicine, University of Bristol, Biomedical Sciences Building, University Walk, Bristol BS8 1TD, UK.; 2) The Wellcome Trust/Cancer Research UK Gurdon Institute and Zoology Department, University of Cambridge, Tennis Court Road, Cambridge CB2 1QW, UK; 3) Perelman School of Medicine at the University of Pennsylvania, 340 Civic Center Blvd, Philadelphia, PA, 19104; 4) Wellcome & MRC Cambridge Stem Cell Institute and Department 17 of Haematology, University of Cambridge, UK

Mutations in ribosome protein (Rp) genes and ribosome biogenesis factors result in debilitating diseases known as ribosomopathies. Recent studies in Drosophila have shown that cells heterozygous mutant for Rp genes (Rp/+) exhibit proteotoxic stress and aggregates, which drive stress pathway activation and apoptosis. Understanding how Rp/+ cells fend off proteotoxic stress could suggest mechanisms to ameliorate these and other conditions caused by proteotoxic stress. Here we find that Rp/+ epithelial cells express all six Gustatory Receptor 64 (Gr64) genes, a cluster of sugar receptors involved in taste sensation. We show that Rp/+ cells depend on Gr64 for survival and that loss of Gr64 autonomously exacerbates stress pathway activation and proteotoxic stress by negatively effecting autophagy and proteasome function in Rp/+ cells. This work identifies a non-canonical role in proteostasis maintenance for a family of gustatory receptors known for their function in neuronal sensation.

212C  BMP-gated cell cycle progression drives anoikis during mesenchymal collective migration  Frank Macabenta, Hsuan-Te Sun, Angelike Stathopoulos California Institute of Technology, Pasadena, CA

Organ integrity requires the elimination of abnormal cells to avoid compromised patterning and function. While cell competition has been characterized as a way to eliminate abnormal cells in epithelial tissues, little is known about how collectively-migrating mesenchymal cells maintain quality control. Here we investigate how lost cells within a migratory cohort are eliminated prior to organogenesis using the Drosophila embryonic caudal visceral mesoderm (CVM), migratory cells that ultimately form the midgut musculature. Through a combination of genetic manipulation, immunostaining, and confocal live imaging, we find that FGF signaling via heartless (htl) is critical to maintaining cell survival by antagonizing the cell death gene head involution defective (hid) specifically in the CVM, and that expression of hid is regulated in a cell-cycle dependent manner prior to myoblast fusion, ensuring that unfit cells are cleared before they can contribute to assembly. We additionally use a cell cycle-coupled degron system to show that precise coordination of the cell cycle is mediated by BMP signaling supported by CVM-expressed Tolkin (Tok) metalloprotease processing secreted Decapentaplegic (Dpp) ligand in the extracellular milieu. Mechanistically, we identified a BMP-
responsive enhancer that controls expression of the mitotic activator string (stg) in the visceral mesoderm. Through this study, we have demonstrated a system by which collectively migrating mesenchymal cells can control for migration errors via crosstalk between FGF and BMP signaling, and how errors in this system can result in significant consequences, including inappropriate invasion of other tissues.

215A Role of M1BP, a transcriptional pausing factor in JNK-mediated cell death during eye development

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In all multicellular organisms, transcriptional regulation is crucial to regulate differential gene expression, which is important during development and growth. Transcriptional pausing is one such mechanism used to control gene expression. Recently, we have shown that M1BP, a transcriptional pausing factor, promotes eye development by suppressing wingless (wg) expression. We showed that loss of M1BP induces ectopic caspase-mediated cell death that is triggered by wg induction. Blocking caspase-dependent cell death using p35 showed significant but incomplete rescue. M1BP is the functional homolog of ZKSCAN3, an autophagy repressor in humans. Jun-amino-terminal-(NH2)-Kinase (JNK) signaling is known to activate both caspase-dependent and -independent cell death. We hypothesized that M1BP could have a role in mediating multiple forms of cell death via JNK signaling during eye development. In our study, we have used the Drosophila melanogaster (Fruit fly) model to study the role of JNK pathway modulation during M1BP mediated eye suppression. Using the GAL4-UAS system, we modulated JNK signaling components along with downregulating M1BP. Here we present data that shows that the absence of M1BP results in activation of autophagic marker and JNK signaling. We also show that activation of JNK signaling enhances M1BPRNAi phenotype and downregulation of JNK signaling rescues the M1BPRNAi no eye phenotype.

216B PDZD8 promotes autophagy at ER-Lysosome contact sites to regulate synaptic growth

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Sites of apposition between organelles, referred to as membrane contact sites (MCSs), are hotspots for intracellular signaling, lipid metabolism, and organelle biogenesis/dynamics in eukaryotic cells. The endoplasmic reticulum (ER) forms an extensive and dynamic network of MCSs with almost all organelles. MCSs between the ER and endo-lysosomes are particularly abundant, suggesting important physiological roles. PDZD8 is an intrinsic ER transmembrane protein with a synaptotagmin-like mitochondrial lipid-binding protein (SMP) domain that has been reported to localize to ER-late endosome/lysosome and ER-mitochondria MCSs. PDZD8 is enriched in neurons. However, its role in the nervous system remains poorly understood. We identified Drosophila PDZD8 in a candidate screen for uncharacterized conserved regulators of synapse formation and function. We used the CRISPR-Cas9 system to generate null alleles and endogenously tag PDZD8. Interestingly, we find that PDZD8 is expressed at synapses throughout the central nervous system and the larval neuromuscular junction (NMJ), where it localizes to ER-lysosome MCSs. We show that activity-induced synaptic growth, neurotransmission, and locomotion are dysregulated in PDZD8 mutants, indicating important roles in nervous system development and function. We further show that PDZD8 regulates synaptic growth via autophagy. In PDZD8 mutants we see accumulation of autophagic proteins as well as autolysosomes. Our analyses suggest that PDZD8 is required for autolysosome maturation to promote autophagic flux in neurons. Overall, we propose that PDZD8-mediated ER-lysosome membrane interactions promote autophagy to regulate synaptic growth.

217C Investigating the contributions of Rab11 and the UPR in amyloid-β load at the Drosophila neuromuscular junction

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Accumulation of abnormal aggregates formed by amyloid-β (Aβ) peptides and synaptic dysfunction are characteristics of Alzheimer’s disease (AD). The accumulation of Aβ fragments during AD results in endoplasmic reticulum stress activating unfolded protein response (UPR) to restore proteostasis and can lead to synapse dysfunction and neurodegeneration. UPR is initiated by activation of stress sensors including the protein kinase R-like ER kinase (PERK), inositol-requiring enzyme 1 (IRE1), and activating transcription factor (ATF). UPR-inhibiting drugs are currently under investigation as a possible approach to modulate the metabolism of the Aβ precursor, amyloid precursor protein, and neuroplasticity. To assess the contribution of the UPR to Aβ load, we pharmacologically inhibited the UPR in a Drosophila AD model. Aβ levels were measured in AD model flies raised on STF-083010 and MCK-3946 (IRE1 inhibitors), GSK2656157 (a PERK kinase inhibitor), and the chemical chaperone Tauroursodeoxycholic acid. We find that AD model animals raised on 10 µM STF-083010 and 50 µM STF-083010 show decreased Aβ loads compared to control AD model animals. In another approach, we examined the effects of Rab11 on Aβ load. Rab11 is a GTPase that facilitates vesicle trafficking from the recycling endosome to the plasma membrane. Vesicle trafficking defects are characteristic of late-onset Alzheimer’s disease. Our collective results will provide insight on the contribution of both Rab11 and the UPR to Aβ load and synaptic dysfunction.
The stress response transcription factor ATF4 regulates oocyte maturation

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Metazoans have evolved various stress response mechanisms to cope with cellular stress inflicted by external and physiological conditions. The Integrated Stress Response (ISR) is an evolutionarily conserved pathway that mediates adaptation to cellular stress via the transcription factor, ATF4. Here we describe a previously unknown role for Drosophila ATF4, encoded by the gene cryptochrome (crc), in oocyte maturation. We found that crc mutants have decreased egg laying and increased frequency of dorsal appendage defects in their eggs, with no apparent effect on fertilization rates. We observed substantial arrest in mid-oogenesis in crc mutant ovaries accompanied by premature apoptosis in early-stage egg chambers. Careful examination of ISR reporters revealed no detectable expression of ATF4 in the ovary, so we propose a non-autonomous role for ATF4 in the regulation of oogenesis.

Using a GFP insertion in the crc locus, we observed high levels of ATF4 expression in fat tissues surrounding the ovary. RNAi-mediated depletion of crc in fat tissues impaired oocyte maturation and decreased fertility similar to crc mutants. High levels of ATF4 expression in fat tissues suggest that this tissue has elevated basal levels of cellular stress. ATF4 activation in Drosophila can be mediated by one of two upstream ISR kinases: PERK, which responds to endoplasmic reticulum stress, or GCN2, which responds to amino acid deprivation. We are currently examining which of these kinases drives ATF4 expression in adult fat tissues to regulate oogenesis. Recent work has demonstrated that ATF4 acts as a coactivator for the Ecdysone receptor isoform EcR-B2 to regulate molting behavior. Since ecdysone signaling is a prominent regulator of oogenesis, we are currently parsing the role of ATF4-EcR-B2 signaling in fat tissues and its non-autonomous effects on oocyte maturation.

The crc 5’ leader has multiple cis-regulatory elements called upstream open reading frames (uORFs) that typically influence translation of the main ATF4-encoding ORF. We previously demonstrated that crc translation is regulated by the translation reinitiation factors eIF2D and the DENR-MCTS1 heterodimer. Consistently, loss of these factors yields similar fertility defects, implicating translation reinitiation in fat tissues as a regulator of oogenesis. Together, our data indicate that stress signaling in fat tissues non-autonomously regulates oocyte maturation.

A Drosophila screen identified a role of histone methylation in ER stress preconditioning

Katie Owings, Clement Chow University of Utah

Organisms face many stressors, and an ongoing challenge is understanding how an individual can respond to numerous insults over a lifetime. The accumulation of misfolded proteins results in cellular stresses, including endoplasmic reticulum (ER) stress. Many studies examine the ER stress response in isolation. In reality, cellular stresses rarely occur in isolation but often in the context of other stresses. Little is known about ER stress preconditioning, whereby conditioning with low levels of stress alters the ability to withstand subsequent ER stress. This project aims to use natural genetic variation to characterize ER stress preconditioning and its underlying mechanisms.

I began with an ER stress preconditioning screen that utilized the 200 Drosophila Genetic Reference Panel (DGRP) strains. Flies were subjected to heat shock (or no heat shock control), allowed to recover, placed on tunicamycin to induce ER stress until death, and survival was measured. Different genetic backgrounds led to a striking range in phenotypic responses to ER stress preconditioning, ranging from dying half as fast to 4.5 times faster with preconditioning than with no preconditioning. A genome-wide association study revealed that histone H3-K4 methylation is a solid potential mechanism of ER stress preconditioning. Several candidate modifiers have known roles in histone methylation. H3-K4 methylation marks promoters at transcribed genes and may play a role in transcriptional regulation. Two top modifiers, Pdp1 and TfIIA-L, are general transcription factors associated with RNA polymerase II. These hits solidify a potential role of transcriptional regulation underlying ER stress preconditioning. RNAseq was performed in the phenotypically extreme DGRP strains at different points in the preconditioning protocol to identify potential predictive gene expression signatures. Differentially expressed genes indicate a potential role of immune genes in ER stress preconditioning.

An effective ER stress response is critical for healthy development and aging. Disruptions in this response have been implicated in multiple human diseases, from diabetes to neurodegeneration. Understanding how previous stress events influence the ER stress response will provide insight into this pathway’s fundamental biology and have important implications for therapeutic development.

Deciphering an unrecognized role of bZIP transcription factor IRBP18 during unfolded protein response (UPR) in Drosophila

Sahana Mitra, Hyung Don Ryoo New York University

In eukaryotes, the endoplasmic reticulum (ER) serves as an essential organelle where membrane and secretory proteins undergo folding and maturation. Any physiological conditions that overwhelm the protein folding capacity of the ER could cause its dysfunction, widely referred to as “ER stress”. The ‘Unfolded protein response (UPR)” is a signaling mechanism that regulates gene expression in response to such ER stress. There are three classical UPR signaling
pathways mediated by ER stress sensors IRE1, PERK, and ATF6 respectively, which are conserved in *Drosophila*. The best-characterized effector of the PERK pathway is a bZIP transcription factor ATF4. The 5′ UTR of ATF4 contains regulatory upstream open reading frames (uORFs) that block ATF4 expression in unstressed cells, but stimulates ATF4 translation specifically in response to PERK activation. We recently found that Xrp1 (bZIP transcription factor) is another effector of PERK that mediates an ATF4-independent branch of PERK signaling. Xrp1 mediates the induction of the anti-oxidant gene expression reporter, gstD-GFP, in response PERK activation. We found that Xrp1 also contains a regulatory 5′ UTR with uORFs. Mutation of the last uORF causes a de-repression of the main ORF expression even in an unstressed cells. We further examined the role of *IRBP18*, a *Drosophila* homolog of *CEBPG* that is known to form a heterodimer with Xrp1 during the regulation cell competition and repair of DNA breaks. We find that the *IRBP18* is required for ER stress-induced gstD-GFP reporter expression. These results indicate that IRBP18 mediates an ATF4-independent branch of the unfolded protein response that may help maintain the redox equilibrium in cells under ER stress.

221A  **ER stress-induced JNK promotes stress granule formation via epigenetic modifications in C9orf72 mediated ALS/FTD** *Sahana TG*, Ke Zhang Mayo Clinic, Jacksonville, FL

Amyotrophic Lateral Sclerosis is a neurodegenerative disease affecting upper and lower motor neurons. A GGGGCC hexanucleotide repeat expansion (HRE) in the gene *C9orf72* is the most common genetic cause of ALS. Interestingly, *C9orf72* mutation also causes Frontotemporal Dementia (FTD), one of the most common early-onset dementia (collectively referred to as c9ALS/FTD). The HRE generates dipeptide protein repeats (DPRs) via noncanonical translation, resulting in five different DPR species. Among these species, the arginine-rich DPRs, poly(glycine-arginine, or GR), and poly(proline-arginine, or PR) are highly toxic and play a critical role in c9ALS/FTD pathogenesis. Stress granules (SGs) are RNA-protein condensates formed in eukaryotic cells upon stress. In c9ALS/FTD, these condensates recruit several RNA binding proteins namely TDP-43, FUS, hnRNP which can potentially form solid aggregates. However, how stress granules cause neurodegeneration is incompletely understood.

In an RNAi screen in a *Drosophila* model of c9ALS/FTD, we identified loss of *basket*, the fly homolog of c-Jun N-terminal kinase (JNK) to suppress neurodegeneration. JNK is a member of the mitogen-activated protein kinase pathway (MAPK). We found that the *basket*/JNK activity is upregulated in fly or cell models of c9ALS/FTD via the activation of ER-stress response protein IRE1/TRAF2. Furthermore, JNK hyperactivation transactivates G3BP1, a key SG assembly factor. Our results show that JNK promotes G3BP1 expression via epigenetic modulation of histone protein 3 (H3). Moreover, IRE1 or JNK inhibitors suppress H3 modification, G3BP1 protein levels, SG assembly, and survival defects in cells expressing poly(GR) or poly(PR). Hence, our data shows a unified connection between key ALS physiologies including ER stress, JNK/ MAPK activation, and SG assembly which contributes to neurodegeneration.

221B  **Adenosine receptor and its downstream targets, Mod(mdg4) and Hsp70, work as a signaling pathway modulating cytotoxic damage in Drosophila** *Michal Zurovec*1,2, Yu-Hsien Lin1,2, Houda Ouns Maaroufi1,2, Lucie Kucerova1, Lenka Rouhova1,2 1) Biology Centre, Inst Entomology, Ceske Budejovice, Czech Republic; 2) Faculty of Sciences, University of South Bohemia, Ceske Budejovice, Czech Republic

Extracellular adenosine (Ado) is an important signaling molecule involved in stress responses. Studies in various model organisms have shown that Ado is involved in “danger recognition” and tissue protection. However, the mechanism of Ado signaling is not well understood. In this study, we observed that Ado signaling involves the mutant huntingtin (Q93-mHTT) protein have reduced extracellular adenosine concentration in hemolymph. Using genetic methods, we altered Ado concentration, suppressed Ado receptor and Ado transporter function, and observed the effects of these changes on Q93-Htt fly viability. We found that overexpression of enzymes of Ado metabolism and suppression of Ado receptor (AdoR) or Ado transporter (ENT) can minimize mHTT-induced mortality. We also identified downstream targets of the AdoR pathway, the modifier of mdg4 (Mod(mdg4)) and heat shock protein 70 (Hsp70), which are able to modulate the formation of mHTT aggregates. Finally, we have shown that reduction of Ado signaling modulates some other responses to stress, including parautax and heat shock treatments. Our study provides important insights into how Ado regulates stress responses in *Drosophila*.

223C  **Investigating neuronal survival from caspase activity in neurodegeneration** *Morgan Mutch*, Jiaqi Chen, Denise Montell University of California, Santa Barbara

The regulation of cell death and survival is critical for tissue development and function. Our lab has characterized a process named ‘anastasis,’ which is defined as cell survival following executioner caspase activation, previously thought to be the irreversible marker of apoptosis. Anastasis has been found to function during development, oncogenesis, and cell survival following injury in a variety of animal and cell culture models. Most anastasis research has focused on mammalian cancer cell lines and fly epithelial tissue, both highly proliferative cell types. However, the significance of anastasis has yet to be determined in the nervous system. Post-mitotic neurons have been shown to initiate apoptosis and activate executioner caspasases without executing apoptosis in a neurodegenerative disease model. We propose that anastasis might enhance survival of neurons following stress like human Tau expression where it might function to slow neurodegeneration. I performed a loss of function screen in *D. melanogaster* photoreceptor neurons to test the functions...
of genes that affect anastasis in proliferating wing imaginal discs for effects on neurodegeneration. The results suggest that the Insulin Receptor/Akt pathway, which drives cell proliferation, enhances Tau toxicity in the Tau-induced rough-eye model. I am currently investigating how Tau interacts with the Insulin Receptor/Akt pathway in fly photoreceptors, and how expression of signaling components changes Akt activity in a Tau background. In addition, I am exploring the role of Insulin Receptor signaling in regulating anastasis and apoptosis in a neurodegeneration model. I will present progress in understanding anastasis and genetic regulators of neurodegeneration in the fly nervous system.

224A Xrp1 and Irbp18 trigger a feed-forward loop of proteotoxic stress to induce the loser status Paul Langton1, Michael Baumgartner2, Remi Logeay1, Eugenia Piddini1 1) School of Cellular and Molecular Medicine, University of Bristol, Bristol, UK; 2) Perelman School of Medicine at the University of Pennsylvania, Philadelphia, PA, USA

Cell competition induces the elimination of less-fit “loser” cells by fitter “winner” cells. In Drosophila, cells heterozygous mutant in ribosome genes, Rp+/+, known as Minutes, are outcompeted by wild-type cells. Rp+/+ cells display proteotoxic stress and the oxidative stress response, which drive the loser status. Minute cell competition also requires the transcription factors Irbp18 and Xrp1, but how these contribute to the loser status is partially understood. Here we provide evidence that initial proteotoxic stress in RpS3/+ cells is Xrp1-independent. However, Xrp1 is sufficient to induce proteotoxic stress in otherwise wild-type cells and is necessary for the high levels of proteotoxic stress found in RpS3/+ cells. Surprisingly, Xrp1 is also induced downstream of proteotoxic stress, and is required for the competitive elimination of cells suffering from proteotoxic stress or overexpressing Nrf2. Our data suggests that a feed-forward loop between Xrp1, proteotoxic stress, and Nrf2 drives Minute cells to become losers.

225B Mechanisms for culling of Drosophila wing disc cells with loss-of-heterozygosity after irradiation Jeremy Brown1, Inle Bush1, Justine Bozon1, Tin Tin Su1,2 1) Department of Molecular, Cellular and Developmental Biology, University of Colorado Boulder, Boulder, CO; 2) University of Colorado Cancer Center, Anschutz Medical Campus, Denver, CO

The ability of ionizing radiation (IR) to induce cell death is the basis for radiotherapy which is used to treat approximately half of cancer patients. IR causes DNA damage and events such as loss of heterozygosity (LOH). Undesirable effects such as oncogenesis can be promoted by functional loss of a tumor suppressor due to IR induced LOH. To study LOH events we use a cell-autonomous fluorescence-based system that uses the QF/QS transcriptional module. This allows our studies to expand into larval, pupal, and adult stages of Drosophila wing development. In parallel with QF/QS, we use the GAL4/UAS system to knockdown or overexpress genes of interest and we use engrailed-GAL4 to restrict UAS expression to the posterior compartment of the wing disc. In our studies, we compare the number and area of LOH clones found in the posterior to those of the anterior compartment after IR. Previously with these tools, we identified two distinct phases of LOH cell culling during development with p53-dependent as well as p53-independent mechanisms (Brown et al., PLoS Genetics, 2020, PMID: 33075096). Today, we will present recent data on LOH culling regarding genes involved with various cellular processes including Apoptosis-induced-Proliferation, ROS signaling, JNK and p53 signaling, and DNA repair.

226C An in vivo Drosophila screen to identify new regulators of ATF4 signaling Kristoffer Walsh NYU Vilcek Institute, New York, NY

In response to various physiological or external stresses, eukaryotic cells activate the Integrated Stress Response (ISR) to regulate protein synthesis and gene expression. ISR signaling begins when stress-activated kinases phosphorylate the translational initiation factor eIF2a, inhibiting general mRNA translation. Under these conditions, a small number of specially-regulated mRNAs – including that of transcription factor ATF4 – are translationally upregulated to activate a stress-induced gene expression program.

Our lab has previously demonstrated that cap-dependent translation inhibitor 4EBP is induced by ATF4 in the Drosophila fat body through the amino acid deprivation-activated ISR kinase GCN2. The 4EBP intron contains ATF4-binding sites that not only respond to environmental stress but also show physiological ATF4 activity during the development of larval fat body tissue. We developed a Drosophila-based ATF4 reporter that relies upon the 4EBP-intron ATF4-binding sites to drive the expression of dsRed, which we designated 4EBP-dsRed.

Using 4EBP-dsRed, I performed an in vivo RNAi screen to find new regulators of ATF4 in the Drosophila larval fat body. Among over 200 RNAI lines, 20 showed a decrease in the 4EBP-dsRed signal relative to wild-type larvae, implicating them as candidate positive regulators of ATF4. We have proceeded to validate these candidates through confocal microscopy and qPCR of endogenous 4EBP. By performing additional experiments in Drosophila tissue and cell culture, we aim to determine how these candidates regulate ATF4 and the ISR.

ISR dysregulation has been implicated in various diseases, including neurodegenerative disorders such as Charcot-Marie-Tooth disease. Identifying new positive regulators of ATF4 could provide a greater understanding of the ISR, potentially
227A Characterizing the landscape of alternative splicing events regulating the clearance of nurse cells by non-professional phagocytes in Drosophila melanogaster oogenesis Shruthi Bandyadka, Diane PV Lebo, Ana Fiszbein, Kimberly McCall Boston University

The death and clearance of nurse cells marks an important developmental milestone in Drosophila melanogaster oogenesis. A subset of the Main Body Follicle Cells (MBFCs) that encapsulate the oocyte and the nurse cells within the egg chamber undergo marked cytoskeletal remodeling to become Stretch Follicle Cells (SFCs). Towards the final stages of oogenesis, the nurse cells dump their contents into the oocyte and are subsequently induced to die by the phagocytosis machinery of SFCs. This non-autonomous, phagocyte-dependent form of cell death is accomplished by the SFCs migrating into the spaces between the nurse cells and acidifying them externally by employing proton-pumping Vacuolar-ATPases. In this study, we characterize the co-transcriptional events that precipitate the transition of columnar, non-phagocytic MBFCs into squamous, phagocytic SFCs, including those that orchestrate the engulfment and processing of nurse cells. We focus on differential expression at the transcript level arising from alternative splicing, which significantly amplifies the transcriptomic repertoire, and thereby the proteomic diversity of tissues.

To capture the incidence and diversity of splicing events between MBFCs and SFCs at the genome-level, we sequenced the mRNAs undergoing translation by immunoprecipitating GFP-tagged ribosomes using Translating Ribosome Affinity Purification (TRAP-seq). We analyzed the short-read RNA-seq data using HIIndex to identify terminal exons that could act as hybrid internal exons, and LeafCutter to identify intron excision events and novel splice sites.

We observed that hundreds of terminal exons in MBFCs were classified as hybrid internal exons in SFCs. Gene ontology analysis revealed that a plurality of genes with hybrid first exons had important physiological functions such as cytoskeletal, trans-membrane, or secreted proteins and were involved in major signaling pathways, including the immune-related Imd and TNF-alpha pathways. Specifically, we discovered that the first exons of Vacuolar-ATPases subunit Vha100-2 and kinase Src42A in MBFCs were used as internal exons in SFCs. Additionally, we found statistically significant exon skipping events in the calcium-binding protein Mlc1 and in Sec23, which is involved in vesicle trafficking in autophagy.

In summary, we have identified novel regulatory events orchestrating nurse cell clearance that are not captured at the gene-level, indicating that co-transcriptional regulation of gene expression could play a more prominent role in cell death and clearance than was previously appreciated.

228B Wdr59 regulates the interaction of GATOR1 with RagA to inhibit TORC1 activity in the Drosophila ovary Yingbiao Zhang, Chun-Yuan Ting, Shu Yang, Mary Lilly National Institute of Child Health and Human Development, NIH, Bethesda, MD

TORC1, a master regulator of metabolism, is dysregulated in a wide array of human diseases. The GATOR complex is an upstream regulator that controls the activation of TORC1 in response to nutrient stress. The GATOR complex is comprised of two subcomplexes, the GATOR1 complex (Iml1, Nprl2, Nprl3), inhibits TORC1 activity by serving as a GAP for the TORC1 activator RagA, a component of the lysosomally located Rag GTPase. Conversely, the GATOR2 complex opposes the activity of GATOR1, thus serving to promote TORC1 activity. Current models developed from studies in mammalian tissue culture cells suggest that the GATOR2 complex contains 5 components, Mio, Wdr24, Seh1, Sec13 and Wdr59, TORC1 activity is down regulated by loss function of any one of these components. In Drosophila, loss function mutants of mio, seh1 and wdr24 exhibit decreased TORC1 activity and cell growth in the female germline. Surprisingly, we find that deletion of the Wdr59 gene lead to the opposite phenotype. Specifically, wdr59 mutants have increased TORC1 activity and ovarian growth and decreased levels of autophagy. Our data support a model in which Wdr59 harbors neither GATOR2 nor GATOR1 function, but act as a positive regulator of GATOR1 by inhibiting the binding affinity of RagA to the GATOR1 complex in Drosophila germ cells.

229V Toxicological study and genetic basis of BTEX susceptibility in Drosophila melanogaster Temitope Adebambo1,2,3, Don Fox2, Adebayo Otitolouju1 1) Emory University School of Medicine; 2) Duke University, Durham, NC; 3) University of Lagos, Nigeria

Drilling and marketing operations in the oil and gas sector has led to the environmental release of benzene, toluene, ethylbenzene and xylene (BTEX) causing harmful effects to man and other environmental components. Drosophila melanogaster has established protocols that can be used to study the effect of these chemicals and the genome wide mechanism underlying their toxic actions. In this study, the toxicological profile of benzene, toluene, ethylbenzene, p-xylene, m-xylene, and o-xylene in Drosophila melanogaster was evaluated in adult flies and first instar larvae of white w1118 strain. The impact of fixed concentrations of benzene and xylene on apoptosis and mitosis were also investigated in adult progenitor tissues (imaginal discs) found in the third instar larvae. In addition, Genome Wide Association Screening (GWAS) of the Drosophila Genetic Reference Panel (DGRP) was conducted to identify genes that are critical for toxicological responses in Drosophila melanogaster for p-xylene. The results of the toxicity tests showed that toluene and p-xylene are the most toxic chemicals to adult flies with LC50 ≥ 0.166 mM, while a significant and dose-dependent
decrease in fly eclosion was observed with benzene, p-xylene, and o-xylene. A significant increase in apoptotic markers and Phospho-Histone-3 (PH3) activity was also observed in the imaginal wing discs of larvae exposed to benzene and p-xylene. The genome-wide analyses revealed few single nucleotide polymorphisms (SNPs) with very low p-values (p<10-5) in 38 regions of Drosophila melanogaster genome as critical for responses to p-xylene. This study reveals the strength of Drosophila melanogaster genetics and apoptotic-proliferative ability of the imaginal disc as an accessible approach to study BTEX compounds.

230V Symptoms following Traumatic Brain Injury in a Drosophila melanogaster CTE Model are Ameliorated by Ketone Body Enantiomers Katelyn Mooney1,2, Geoffrey Tanner1,2, Jeremy Balsbaugh1, Kate Gavilanes1, Dariana Mota1, Joseph Mooney1, Mirsha Pierre1, Alana Grant1, Hanna Val Pelt1, Yves Thelusma1 1) University of Connecticut; 2) The Connecticut Institute for Brain and Cognitive Sciences

Traumatic Brain Injury (TBI) events—which are increasing in both concern and incidence in contact sports—are defined as brain injuries sustained from a blow or force to the head that disrupts normal brain function. Following TBI, individuals may experience headaches, confusion and disorientation, and difficulty speaking. Multiple TBI events may lead to Chronic Traumatic Encephalopathy (CTE), a progressive neurodegenerative condition characterized by impaired judgment and learning and memory, and personality changes including anxiety, depression, and hyper-aggressive behavior. Progressively more young athletes in contact sports are experiencing repeated TBI, making research into TBI and CTE critical.

The ketogenic diet (KD) is a low-carbohydrate, adequate-protein, and high-fat diet that has been used effectively in the clinic to treat neurological disorders such as intractable pediatric epilepsy. We have shown that directly supplementing standard high-carbohydrate Drosophila melanogaster diets with ketone bodies (K Bs; specifically, beta-hydroxybutyrate, BHB) increases lifespan and decreases male-on-male aggression following TBI events. Our results have suggested that different enantiomeric forms of exogenous BHB may vary in their effectiveness at reducing aggression and increasing lifespan. Compared with a control diet (CD) alone, dietary supplementation with either pure R-BHB (the main endogenous form), or a BHB racemic mixture, both significantly decrease number of post-TBI aggressive events in male-male pairs; racemic and pure-S-BHB lengthen latency to the first aggressive event. The mechanism by which BHB supplementation ameliorates elevated aggression and increases lifespan following TBI is poorly understood. BHB’s primary metabolic function is mitochondrial ATP production under conditions of low circulating glucose levels leading to elevated brain-wide ATP/ADP ratios. However, BHB can also act as a GPCR ligand, a gene-expression regulator (via histone deacetylase inhibition), and a sirtuin protein activator (via increased cytoplasmic [NAD+]).Sir2 activation promotes cell survival through a variety of mechanisms, including activation of autophagy proteins (e.g. autophagy-related protein 8a; Atg8a) which can mediate clearance of damaged cellular components. We hypothesize that BHB works to decrease neuronal cell death following TBI by improving neuronal metabolic stability and by activating and upregulating levels of Sir2 and Atg8a. Future experiments will include confirming reduced neuronal cell death, studying the ability of BHB to rescue learning following TBI, and determining NAD+/NADH ratios in whole brains from TBI-subjected flies receiving exogenous BHB.

231V Roles of Hippo and Ecdysone Receptor Signaling in the regulation of dronc Karishma Gangwani1, Amit Singh1,2, Madhuri Kango-Singh1,2 1) Department of Biology, University of Dayton, Dayton OH; 2) Integrative Science and Engineering Center, University of Dayton

The Hippo pathway coordinately controls the essential growth regulatory processes of cell proliferation and death, and is required for regulating organ size in both flies and humans. Downregulation of Hippo signaling results in tissue overgrowth and tumorigenesis, whereas hyperactivation of Hippo signaling results in caspase-mediated cell death. Caspases are major regulators of apoptosis and suppression of caspase dependent apoptosis is often linked to aberrant growth. Further, JNK activation or hyperactivation of Hippo pathway by Mst1 (Drosophila Hippo) overexpression or YAP (Drosophila Yorkie, Yki) knockdown causes mitochondrial damage that induces Caspase-9 mediated apoptosis in cancer cell lines. Thus, caspases play a role in maintaining tissue homeostasis and in diseases like cancer. Previously, we reported that the initiator caspase Drosophila Nedd-2 like caspase (dronc), Caspase-9 homolog, is a transcriptional target of the Hippo pathway in Drosophila. We showed that dronc transcription requires the growth inhibitory Hippo component Warts (Wts). Developmental apoptosis is controlled by small lipophilic hormones like ecdysone. Earlier studies have shown that dronc expression is regulated by the Ecdysone receptor (EcR) signaling pathway and an EcR regulatory element has been identified on the dronc promoter. Recently, Wts was shown to regulate Ecdysone production through Yki and the microRNA Bantam, suggesting a feedback mechanism involving Hippo and Ecdysone pathways. We found that depletion of Scalloped (Sd, TEAD in mammals) or Ecdysone receptor (EcR) or their corepressors resulted in derepression of dronc expression suggesting that dronc may be regulated by a default repression mechanism. Other preliminary data suggest that dronc regulation may involve cooperative interactions between Sd and EcR. Although an EcR binding site is mapped to the dronc promoter, the role of Sd has not been characterized. Using qRT-PCR based approaches we confirmed that dronc expression is regulated by Yki/Sd and EcR. Further, we identified a potential Sd site on the dronc promoter. Here, we present our work on the regulation of dronc by the Hippo and EcR signaling pathways,
and its implications on development.

232V  JNK-independent Eiger/TNFR signaling during cell competition  Aditi Sharma Singh, Laura A. Johnston  Columbia University Medical Center, New York, NY

Cell competition is a surveillance mechanism that promotes tissue/organ fitness by allowing healthy cells to contribute to organ development at the expense of cells that are relatively less fit. Such competitive interactions can benefit the tissue by eliminating unhealthy cells, but can also be detrimental when oncogenic cells take over an otherwise healthy tissue in a process known as super-competition. We have developed a model of super-competition using Drosophila wing imaginal discs, wherein cells over-expressing the proto-oncogene Myc (“winners”) compete for space with wild-type (WT) cells (“losers”) during the rapid growth phase of larval development. In this model, the death and elimination of WT loser cells from the wing disc is mediated by cell-cell interactions that activate a cell competition signaling module (CCSM) in loser cells, consisting of Toll-related receptors, the secreted ligand Spatzle, and NF-kappaB proteins. Our recent genetic evidence indicates that Eiger (Egr), the sole TNF in Drosophila, and its cognate receptor, Grindelwald (Grnd)/TNFR, also contribute to the elimination of WT loser cells. Egr/Grnd signaling requires the TNFR adaptors Traf4/6 to kill the loser cells, but not Tak1 or other canonical downstream JNK effectors, indicating that in the competitive context, Egr functions independently of JNK activity. Moreover, although Grnd is expressed in all wing disc cells, in-vivo studies with Egr reporters and GFP fusion proteins reveal no evidence of Egr expression in either loser or winner cells. However, Egr, produced as both a secreted and transmembrane protein, is expressed in multiple tissues in growing larvae, so could remotely contribute to loser cell elimination. To determine which cells/tissues produce Egr during cell competition, we are selectively depleting egr from each expressing tissue while concomitantly inducing competitive interactions in wing discs. In addition, we are carrying out experiments to determine the mechanism by which Egr/Grnd activity intersects with the CCSM. Our experiments will provide insight into how local and remote signals converge to influence cell-cell interactions in the growing wing disc during Myc induced super-competition.

233V  Utilizing Live Cell Imaging in Drosophila melanogaster Salivary Glands to Determine if Resveratrol Treatment Activates Heat Shock Factor DNA Binding  Martin Buckley, Nichole Webb, Riley Bricker, Tyra Skalos, Stacy Hrizo  Slippery Rock University of Pennsylvania

Oxidative stress is a hallmark of many aging diseases. There is great interest in understanding the mechanism of action of anti-oxidant treatments and their effects on the cell. One major stress response pathway is the heat shock response (HSR) that is mediated by the transcription factor, heat shock factor (HSF). The HSR is activated in cells exposed to conditions that induce protein misfolding such as: high heat, oxidants, reductants and other chemical stresses. HSF activates expression of the Hsp70 chaperone, which helps cells deal with protein folding stress. We examined the effect of feeding the anti-oxidant, resveratrol on the ability of wildtype Drosophila to withstand heat stress. Treatment with 100 and 400uM resveratrol increased the ability of the flies to withstand heat stress. One possible reason for this is the flies had increased HSF activity due to the resveratrol treatment. To examine this hypothesis, Drosophila larvae expressing HSF-GFP were dissected to obtain the salivary glands. These glands contain polytene chromosomes that allow for visualization of HSF binding onto the chromosome using confocal microscopy. The major binding site that is easily visualized is a doublet of HSF binding at the Hsp70 loci on the chromosome. Salivary glands at room temperature function as a non-heat shock control and exhibit no binding of HSF-GFP at the Hsp70 loci. Salivary glands heated to 37C for 10 minutes function as the positive control and exhibit the expected Hsp70 doublet from HSF-GFP binding of the DNA. We are testing variable concentrations of resveratrol to determine if it activates HSF-GFP binding of the DNA in salivary glands under non-heat shock conditions. Future experiments may examine if the HSF-GFP is transcriptionally active when cells are treated with resveratrol.

234V  Same tissue, different responses: How do different cells in the Drosophila wing imaginal disc respond to ionizing radiation and contribute to tissue homeostasis?  Joyner Cruz, Andreana Gomez, Iswar Hariharan  University of California Berkeley

Tissues consist of highly organized cell populations which must mount coordinated responses to damaging stressors in order to maintain tissue viability. In the Drosophila wing imaginal disc, observations of coordinated homeostatic responses to diverse damaging events are abundant- however, the exact mechanisms of this coordination are not fully understood. Here we show that cells in the notum of the wing disc are resistant to ionizing radiation, and that they lose this resistance through developmental time. We also show that apoptosis in this region occurs independently of the JNK stress-signaling pathway, in contrast to other regions of the disc. Our results provide further evidence that different cell subpopulations in the wing disc, which are canonically considered to be of the same cell type, have strikingly heterogeneous responses to ionizing radiation and may contribute to total tissue response in specialized ways.

235C  Characterization and functional analysis of diverse reactive Oxygen species produced during the immune response to bacterial infection  Alva Duenas, Caitlin Harris, Catherine Brennan  Cal State University Fullerton
Reactive Oxygen Species (ROS) are a class of highly reactive Oxygen-containing ions and molecules that can be produced in cells by direct enzyme-mediated catalysis, as byproducts of mitochondrial respiration, and as secondary metabolites produced by the reaction of ROS with each other and other molecules in the cell. Identifying these short-lived molecules can be challenging due to their short-lived nature, as well as their similar chemistries and interactions with commonly used fluorescent ROS probes. In the immune response, ROS are produced in both fly and mammalian macrophages, where they play roles in both direct killing of engulfed microbes, as well as regulatory roles in macrophage activation and inflammation. We are dissecting the fly macrophage ROS response using a combination of genetic and biochemical tools, to better characterize ROS produced at different stages of the immune response, and to determine their distinct functions. We find that reactive Nitrogen species (RNS) produced by Nitric Oxide Synthase modulate the ROS response, and are responsible for some inflammatory processes previously ascribed to ROS.

### 236A Identification and characterisation of functionally distinct macrophage subpopulations in Drosophila

**Martin Zeidler**, Jonathon Coates, Elliot Brooks, Amy Brittle, Emma Armitage, Iwan Evans

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*Drosophila* blood is dominated by a macrophage-like lineage of cells termed plasmatocytes which differentiate in two waves during embryonic and L3 larval development and are maintained throughout adult life. Throughout this time, plasmatocytes fulfill multiple roles, including wound responses, clearance of apoptotic debris and the provision of an innate immune response to infection. However, until very recently, the plasmatocyte lineage was thought to represent a homogeneous population — a stark contrast to vertebrate macrophages which assume a range of diverse functional and gene expression states.

Here we present our identification of a number of enhancer elements labelling plasmatocyte subpopulations, which we show to vary in abundance across development. We show that some of these subpopulations exhibit functional differences compared to the overall blood cell population, including more potent responses to wounds, altered apoptotic cell clearance and differential localisation / dynamics in pupae and adults. Genes located adjacent to enhancers showing altered plasmatocyte behaviour include Calnexin14D, which, when over expressed in all plasmatocytes, improves wound responses, causing the overall population to more closely resemble the marked subpopulation. These results suggest that differential gene expression not only labels, but also drives changes in subset behaviour. Finally, we show that exposure to increased levels of apoptotic cell death is sufficient to modulate the number of cells within certain subpopulations — further suggesting that subpopulations are dynamic and respond to environmental cues and stresses.

Taken together, our work demonstrates a degree of adaptive and functional macrophage heterogeneity in *Drosophila* that has not previously been described. It has identified mechanisms involved in subpopulation specification and function and provides a starting point for future studies into functional in vivo *Drosophila* macrophage heterogeneity.

### 237B Diptericin A protects flies from opportunistic gut infections in a sex dependent manner

**Sarah Mullinax**, Robert Unckless University of Kansas

Living things must be able to balance their immune defenses to control potentially pathogenic microbes while also fostering beneficial ones. This is especially important when considering gut immunity. Most animals are constantly ingesting bacteria from their food and their gut immunity must be able to rid the animal of potentially harmful bacteria while maintaining beneficial bacteria. Several have shown that the humoral immune system influences the gut microbiome in *Drosophila*, but there has been much less investigation of the role of specific effector genotypes in shaping the microbiome and microbiome-related phenotypes. In this this study we focused on the *Drosophila* antimicrobial peptide Diptericin A. Diptericin A is especially important in the defense against a systemic infection by the Gram- bacteria *Providencia rettgeri* and the survival probability is highly dependent on the genotype of diptericin A. Diptericin A is expressed in the gut under normal rearing conditions. We tested the effect of the presence/ absence and genotype of diptericin A in the gut after oral infection using gnotobiotically reared flies. We assessed the effect of mono- and poly-association with common gut bacteria including *Acetobacter* and *Lactobacillus* species as well as with *P. rettgeri*. We found remarkable sex-specific, microbe-specific and sex-by-microbe effects on bacterial load and fitness-related traits. Most markedly we found a mono-association with *Lactobacillus plantarum* greatly reduced the lifespan of female diptericin A null flies. This decrease in lifespan in diptericin A null flies was also observed in poly-associations that included *L. plantarum*. Surprisingly these results were not seen in male flies, lending support to the need to include both sexes in microbiome studies. Furthermore we are seeing a potential diptericin genotype effect when female flies are poly-associated with *L. plantarum* and *A. tropicalis* which is interesting in the context of there also being genotype effects after systemic infections. Overall this research characterizes the importance of having functional diptericin A in the gut for fly fitness.
With World Health Organization (WHO) scientists recently confirming an outbreak of Zika virus (ZIKV) in India, the characterization and understanding of the molecular mechanisms underlying immune responses against ZIKV have, once again, become urgent. ZIKV, a member of the Flaviviridae family, is transmitted to humans primarily by the infected Aedes mosquito species, causing neurologic complications that range from sensory neuropathy and seizures to congenital Zika syndrome (microcephaly) in infants born to mothers infected during pregnancy. Previous research presents compelling evidence that establishes the fruit fly Drosophila melanogaster as a reliable model for studying arboviruses, as many of the signaling pathways identified are evolutionarily conserved amongst insect species. Particularly, the large conservation between Drosophila and mosquitoes has paved the way for our study to provide novel insights into ZIKV pathogenesis and host antiviral immune function using a brain tumor Drosophila model. By examining gene expression in adult Drosophila, we show that ZIKV can replicate efficiently in the brain and activate various immune responses. More specifically, our results confirm that RNA interference (RNAi) plays a key role in limiting ZIKV replication in the brain, thus forming a potent antiviral defense in Drosophila and presenting a potentially powerful tool to combat the host brain tumor. Behavioral analysis also shows that ZIKV causes severe locomotor impairment in infected flies, a common phenotype of neurodegenerative diseases such as Zika. Most importantly, our observations show significantly higher ZIKV rates and effects in female flies than males, indicating possible differences in the rates of infection and susceptibility to the development of disease. These findings make it imperative to continue uncovering the specifics of the complex host-virus interactions and provide additional insights that can potentially be harnessed for the control of different viral pathogens and define the full spectrum of antiviral immunity across various hosts.

An organism’s diet is a critical factor for its ability to survive an infection, and overnutrition of dietary sugar in particular has been demonstrated to increase susceptibility to infection across animal models and humans. Adult Drosophila melanogaster reared on high sugar diets experience higher bacterial burdens and higher mortality post infection, although the genetic and physiological mechanisms that lead to this outcome are not well understood. Prior studies that have investigated the impact of high dietary sugar on survival of infection have been performed using flies that that fed on high-sugar diets throughout their entire life, thus making it difficult to distinguish developmental consequences from acute metabolic effects on immunity. We hypothesized that even transient exposure to high-sugar diet might cause metabolic dysregulation with adverse consequence for immune function. To test this hypothesis, we reared D. melanogaster on a standard 4% (w/v) sucrose diet, then transferred them to one of six experimental diets varying from 0% - 24% sucrose upon eclosion. After 3-5 days of feeding, we assayed their infection survival over five days against systemic infection with two Gram-negative bacteria, Providencia rettgeri and Serratia marcescens, and one Gram-positive bacteria, Enterococcus faecalis. I found that even three days of exposure to elevated sugar in the diet was sufficient to significantly reduce the probability of surviving infection. However, this effect was specific to the Gram-negative bacteria assayed, suggesting that the immediate metabolic effects of high sugar may have disproportionate impact on the immune pathways that are required for combating Gram-negative infection. In ongoing work, we are testing infection survival in other Gram-positive bacteria, like Lactococcus lactis, to determine whether the sugar content affect is specific to Gram-negative bacteria. Additionally, we are testing whether elevated dietary sucrose specifically impairs IMD pathway signaling, and the extent to which different pathogens may themselves have different sensitivities to the excess sugar within their host. These results are foundational for further understanding of the genetic and physiological mechanisms by which high sugar impacts infection survival.

The respiratory burst is a powerful weapon utilized by leukocytes to protect the host from dangerous pathogens and degrade cellular debris. Nevertheless, Reactive Oxygen Species (ROS) are exceptionally reactive molecules that could cause serious bystander damage to both, the phagocyte itself and the surrounding environment. Therefore, to sustain full functionality under these hostile conditions, immune cells must trigger a complex and robust self-protection program. This cytoprotective system must be carefully orchestrated to ensure leukocytes limit the undesirable side-effects of the constant oxidative challenge, while allowing low levels of ROS for crucial cellular signalling. Here, we discuss the molecular nature of these self-protective strategies, dissecting in vivo both their precise mechanism of activation and physiological relevance. We provide evidence that Drosophila macrophages robustly activate the master redox-sensitive transcription factor Nrf2 upon apoptotic corpse engulfment and this activation is driven by calcium and PI3K-dependent release of ROS by NOX oxidases at the phagosome. We find that this Nrf2-mediated antioxidant response is vital to promote Drosophila macrophage tolerance to oxidative stress during normal homeostatic behaviour, preserving efficient basal motility and allowing a robust and timely detection of epithelial wounds. Moreover, by reducing the oxidative burden, Nrf2 not only delays the onset of immune-senescence but supports host longevity by...
limiting collateral damage to surrounding tissues. Overall, we find that by preserving a healthy immune system, leukocyte Nrf2 plays important paracrine effects by delaying premature systemic aging and improving organismal lifespan. We propose that future therapeutic interventions aimed to boost immune self-protection may be beneficial to delay immune aging and alleviate correlated morbidities.

241C Identifying Candidate Genes and Genetic Networks that Influence the Age-specific Ability to Clear an Infection Using Genome Wide Association Tests (GWAs) Shonda Campbell, Jeff Leips University of Maryland Baltimore County

The innate immune response is an evolutionarily conserved process that is essential for host survival in all multicellular organisms. As individuals age, immune functions begin to decline, or immunosenesce, reducing one's ability to fight infections, posing a serious risk to human health. The way that age affects the immune response can vary greatly among individuals, and although this variation has a significant heritable component, the genes responsible for this variation are not known. Previous data using 12 inbred *Drosophila melanogaster* lines from the *Drosophila* Genetic Reference Panel (DGRP) found that the genes implicated in clearing an infection were different between young and aged flies. The goal of this project is to identify genes that regulate age-specific immune responses to better understand the mechanisms that give rise to immunosenescence. Currently, standard bacterial clearance assays and genome wide association tests (GWAs) are being performed using the 192 DGRP lines available. To assess the ability of each line to clear an infection, one and five week old virgin flies are injected with 18.4nL of an *E. coli* solution at an OD$_{600}$ of 1.0+ 0.01. After 24-hours, the surviving flies are individually homogenized and plated on streptomycin LB/agar plates. The colony forming units per milliliter (CFU/mL) represents the phenotype of that individual and reflects the remaining bacteria in the fly. A GWAS will then be performed at both ages to identify candidate genes involved in clearance across ages. Those candidate genes will be the focus of future studies where the results could lead to improved therapeutic treatments in an aging population, providing age-appropriate drug targets to restore the immune function.

242A Modified binding site of IDGF proteins is important for their function Vaclav Broz1, Houda Ouns Maaroufi1,2, Michal Zurovec1,2, Lucie Kucerova1 1) Biology Centre of the Czech Academy of Sciences, Institute of Entomology, Ceske Budejovice, Czechia; 2) University of South Bohemia, Faculty of Science, Ceske Budejovice, Czechia

IDGFs (Imaginal Dish Growth Factors) are a family of chitinase-like proteins secreted into the insect hemolymph from the fat body and hemocytes, they also represent important component secreted from salivary glands and silk glands of insects. IDGFs lost the hydrolytic activity of true chitinases but they are still able to bind carbohydrate moieties. They are implicated in multiple functions, including extracellular matrix reorganization, cell growth stimulation, wound healing, insect hemolymph clotting, antimicrobial activity, detoxification and also insect silk spinning. With its crystal structure determined, *Drosophila* IDGF2 is the best characterized member of the IDGF family. IDGF2 possess a unique shaped carbohydrate binding domain which is different from chitinases and other chitinase-like proteins.

We conducted a glycan array screen to find the binding partner of IDGF2. We established a functional in viv0 assay based on the ability of IDGF2 to induce genes involved in innate immunity and inflammation. We compared the induction of wild-type and binding domain mutant forms of IDGF2. Moreover, we also examined the mutant protein distribution in tissue.

In the present study, we have identified Galα1-3(Fucα1-2)Galβ1 carbohydrate moiety as the preferred binding partner of IDGF2. Unlike other known chitinase-like proteins, IDGF2 does not bind to N-acetylgalcosamine. Furthermore, we proved that the function of IDGF2 depends on its binding ability.

Our work brings new insights about insect chitinase-like proteins binding abilities, which are crucial for their function as modulators of host response in insects as well as mammals.

243B Phagocytic defects lead to or exacerbate neurodegeneration through increased immune signaling Guangmei Liu1, Johnny Elguero1, Katie Tiemeyer1, Heena Gandevia1, Iker Etchegaray1, Mel Feany2, Kim McCall1 1) Department of Biology, Boston University, Boston, MA; 2) Department of Pathology, Harvard Medical School, Brigham and Women's Hospital, Boston, MA

In nervous system development, as well as in disease and injury, neurons die through programmed cell death, leaving behind cell corpses which must be removed. The clearance of these corpses is accomplished through phagocytosis. In the nervous system, glial cells act as phagocytes, engulfing dead neurons and debris to ensure proper morphology and tissue homeostasis. Glial phagocytosis has been implicated in several neurological diseases. In humans, increased numbers of phagocytic glia are observed in conditions like Alzheimer’s disease, Parkinson’s disease, and traumatic brain injury. In vitro, glia have been shown to clear protein aggregates like those found in neurodegenerative disease. Moreover, variants of genes implicated in glial phagocytosis have been identified as risk factors for neurodegenerative diseases. However, how phagocytosis defects might cause or worsen neurodegeneration remains unknown. We are using the *Drosophila melanogaster*, whose complex nervous system harbors phagocytic glia analogous to those in humans, to tackle this question. Our lab previously found that mutant flies lacking the phagocytic receptor Draper show an accumulation of neuronal cell corpses, which result from developmental programmed cell death and persist throughout the organism’s life. And flies lacking glial Draper display age-dependent neurodegeneration.
To determine how phagocytic defects lead to neurodegeneration in the *draper* mutant, we investigated the hypothesis that persisting cell corpses in the brain lead to chronic increased immunity, resulting in neurodegeneration. We measured activation of the immune pathway Imd in aging *draper* mutants and found that the antimicrobial peptide attacin A is highly overexpressed in fat body. We then suppressed the Imd pathway by knocking down *Relish* in glia and fat body in *draper* mutants and found that neurodegeneration was reduced, indicating that immune activation promotes the neurodegeneration in *draper* mutants. Taken together, these findings indicate that phagocytic defects lead to or exacerbate neurodegeneration through increased immune signaling, both systemically and in the brain.

To define the causality of neurodegeneration in *draper* mutants, we are testing two hypotheses that the neurodegenerative phenotype results from persisting corpses or from defective phagocytic function of glia. We are comparing the severity of neurodegeneration in aging flies with either only persisting corpses or only defective phagocytic function. To further define the consequences of neurodegeneration caused by knocking down *draper* in fly glia, we are characterizing the cellular identities of dead neurons in these aging flies and examining related behavioral deficits such as locomotion activity and learning and memory function.

**244C Peroxisomes regulate the Imd amyloid fibril formation and subsequent Relish signaling pathway** Yizhu Mu1,2, Anni Kleino3, Neal silverman2, Francesca Dicara1,3 1) Dalhousie university, Halifax, NS, Canada; 2) University of Massachusetts Medical School, Worcester, MA, USA; 3) IWK heath center, Halifax, NS, Canada

The Imd pathway plays a pivotal role in the Drosophila defense response against bacterial infection. Gram-negative bacterial infection activates Imd protein which promotes the activation of the NF-kB homolog Relish for the robust antimicrobial peptide gene expression. Although Imd activation is finely regulated, our knowledge of the cellular mechanisms modulating the Imd pathway is still in its infancy. Understanding of the Imd pathway is further complicated by the involvement of Imd protein in the formation and resolution of amyloid-like structures which are indispensable for the activation of the Imd pathway. Peroxisomes are essential lipid metabolic organelles known to play an important role in innate immunity through regulation of cellular lipids milieu in the activated immune cells. Moreover, it is known that defects in peroxisome metabolism inhibits the activation of the Imd pathway. Therefore, we hypothesize that changes in the lipid milieu at the plasma membrane might contribute the Imd amyloid fibril formation and subsequent signaling. Here we demonstrated that peroxisomes regulate Imd amyloid-like structure formation and the activation of the Imd signaling. Using immunofluorescence experiments of *Drosophila* S2 cells, we found that Gram-negative bacterial infection results in the robust amyloid formation in wild type (WT) S2 cells whereas in the absence of functional peroxisomes (peroxisome depleted cells) amyloid structures are not detected. Furthermore, bacterial infection results in the formation of distinct Imd punctate structures that co-localize with amyloid structure in the WT S2 cells suggesting the activation of the Imd pathway. In contrast, peroxisome deficient S2 cells show impairment in the formation of distinct Imd punctate structures in response to bacterial infection. Furthermore, lipids such as PIP2 and PA (whose biosynthesis are known to require peroxisomes) were found to interact with Imd protein in vitro. Altogether, our initial study shows that peroxisomes play a functional role in the activation of the Imd pathway by regulating IMD amyloid structures formation.

**245A Identification of Enhancers of the Drosophila Innate Immune System** Lianne Cohen, Zeba Wunderlich Boston University, Boston MA

An organism’s ability to launch an appropriate and robust response to environmental stimuli is critical for its survival. Infection from a variety of microorganisms is a common and potentially lethal threat that host organisms combat by initiating an immune response. Much of this response is controlled by regulatory DNA, specifically promoters and enhancers. Enhancers are *cis*-regulatory sequences containing transcription factor binding sites (TFBS) that can direct gene expression at a distance along the chromosome from the transcriptional start site. While enhancers and TFBS have been identified for several immune responsive genes in *Drosophila*, the vast majority of enhancers that regulate immune induced genes are unknown and uncharacterized. Without knowledge of these regulatory elements, our understanding of how immune signal pathways interact at the *cis*-regulatory level and the grammar needed for this type of gene expression is limited. To identify immune responsive enhancers, we stimulated the major immune pathways, Toll and IMD, both individually and together in S2 cells and measured enhancer activity via STARR-seq (Self transcribing active regulatory-region sequencing). This activity-based assay identifies sequences that can drive their own expression upon activation of immune signaling pathways in a high-throughput manner. A large-scale analysis of immune enhancers permits us to characterize the regulatory grammar found in these sequences, as well as unknown immune TFBS. Additionally, by stimulating two immune pathways in this experiment we can investigate the role enhancers play in producing synergetic gene expression. The identification of immune enhancers will provide insights into how animals use regulatory sequences control their gene expression response to a rapidly changing environment and thus survive infection from pathogens.

**246B Title: Exploring transcriptional signatures of Anti-Microbial Peptides early in infection to predict infection outcomes** Radhika R1, Brian Lazzaro2,3 1) Cornell University; 2) Cornell Institute of Host-Microbe Interactions and Disease (CIHMID)

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#Dros22 | 73
Production of Anti-Microbial Peptides (AMP) is an integral part of the anti-bacterial infection response in insects. Controlled primarily by the Toll and Imd signaling pathways, a wide array of AMP genes are upregulated within four to six hours of hosts encountering bacterial pathogens. A model of infection progression developed by Duneau et al. 2017 suggests that this early phase of infection response, lasting up to about twelve hours, is very important for effective pathogen control. Hosts that survive infection do so by restricting pathogen load to a low, sustained residual burden that they retain for the rest of their life, while those that allow pathogens to grow unchecked succumb to infection. In this study, using Drosophila melanogaster as our model system, we test whether differences in AMP induction kinetics in the early phase of infection predict how well hosts curtail pathogen growth and thereby survive infections. Our hypothesis is that faster upregulation of the AMP(s) will correlate with increased probability of host survival and lower residual bacterial loads among surviving hosts. We first systematically tested which AMP(s) are the most important for controlling infection by each of four different bacteria using a panel of mutants that are deficient for different combinations of AMP genes. We then measured ten lines from the Drosophila Genome Reference Panel (DGRP) for their ability to survive infection with the four bacteria. In ongoing studies, we are testing whether the kinetics of AMP upregulation predict the probability of host survival of infection, specifically predicting that the strongest correlation will be seen with the expression of AMPs that are disproportionately important for controlling an infection by a given bacteria while other co-regulated AMPs may correlate more weakly. If the data support this hypothesis, that would provide key evidence that the rapid induction of AMP gene expression is essential for control of infection.

247C Metchnikowin alleles are associated with both immune and life history phenotypes Jessamyn Perlmutter1, Joanne Chapman2, Robert Unckless1 1) Department of Molecular Biosciences, University of Kansas, Lawrence, KS; 2) Enteric, Environmental and Food Virology Laboratory, Institute of Environmental and Scientific Research, New Zealand

Antimicrobial peptides (AMPs) are at the interface of interactions between hosts and microbes and are therefore expected to be fast evolving to keep pace with pathogens. However, production of these peptides is also costly to animals and the host must balance its own energetic and antimicrobial interests. Previous work identified distinct peptide alleles that are repeatedly maintained not only within populations of Drosophila melanogaster, but also across at least three other fly species. The maintenance of these polymorphisms across both geography and more than 10 million years of divergence, coupled with evidence showing balancing selection on AMPs in flies, suggest there is a distinct functional importance to each allele. Is each allele more effective against specific pathogens (specificity hypothesis), or is one allele more effective generally, but also more costly to produce (autoimmune hypothesis)? One AMP, Metchnikowin (Mtk), has a single residue that segregates as either proline (P) or arginine (R) in populations of four Drosophila species. To assess the functional difference between the alleles, we created D. melanogaster lines with the P allele, the R allele, or a Mtk null mutation using CRISPR/Cas9 genome editing. Here, we report results from experiments assessing the specificity versus autoimmune hypotheses using these lines. Testing of the flies against a repertoire of bacteria and fungi demonstrated differences in survival rates across allele or null mutation lines with some pathogens. In addition, measurements of various life history traits in these lines reveal differences in longevity in adulthood, suggesting differential fitness costs to producing the alleles. In summary, there are various measurable functional differences between Mtk alleles in Drosophila melanogaster, providing several lines of support for adaptive maintenance of Mtk polymorphisms in contrast with expectations of rapid AMP evolution.

248A JAK/STAT mediated metabolic reprogramming during immune response Ellen McMullen, Lukas Strych, Tomas Dolezal University of South Bohemia in České Budějovice

JAK/STAT is a highly conserved pathway, which plays a key role in immune response. In the case of Drosophila larvae, JAK/STAT signaling is involved in differentiation of hemocytes into lamellocyte to fight against parasitoid wasp infection (Leptipilina boulardi). This, in part, is mediated by the secretion of cytokines Upd2 and Upd3 from hemocytes, which activates JAK/STAT signaling in skeletal muscles; required for the efficient encapsulation and melanization of the wasp egg. Deletion of Upd2 and Upd3 leads to significant reduction in lamellocyte number, and therefore efficient immune response.

In times of food scarcity, or high metabolic demand tissues compete for the nutrients available. Parasitoid wasp infection leads to a redistribution of nutrients from surrounding tissues, such as muscles and fat body, to provide sufficient energy for immune response. This supports the ‘selfish’ or ‘privileged’ immune system hypothesis, as the immune cells are prioritized over other tissues to aid the survival of the animal. Reduction in carbohydrate uptake in peripheral tissues during infection is thought to be mediated by suppression of insulin signaling (IS) in these non-immune organs. Our findings show lower levels of pAkt in the larval fat body and skeletal muscles during infection, indicating IS suppression. We propose the reduction in IS observed is a result of Insulin resistance (IR) in these tissues. Furthermore, we suggest this IR occurs via Upd2 and Upd3 mediated activation of JAK/STAT in muscles leading to a systemic metabolic reprogramming during infection.

249B The Role of Professional Phagocytes during Cell Death in the Ovary of Drosophila melanogaster Alexandra Chasse, Shruthi Bandyadka, Max Werthemier, Kim McCall Boston University, Boston, MA
Regulated cell death is a key process necessary for maintenance of homeostasis and removing toxic cells from the organism and involves signaling responses among several cell types that are not well understood. Apoptosis is the best understood type of regulated cell death and once cells begin apoptosis, they produce multiple signaling cues to entice phagocytes. Macrophages are the professional phagocytes (PPs) that clear apoptotic cells in many tissues and organs, however other cells, such as tissue resident epithelial cells, act as non-professional phagocytes (NPPs). While there has been extensive research on mechanisms used to engulf dying cells for both PPs and NPPs, there is little research done to identify mechanisms used for communication. The Drosophila ovary is a powerful model for cell death mechanisms despite being “immune-privileged” meaning that the ovary is protected from an uncontrolled immune response by denying hemocytes access to the organ. However small numbers of hemocytes have been detected in the ovary and previous work has shown that the cytokine Unpaired 3 (Upd3, IL-6 like) is highly expressed in NPP Follicle Cells (FCs) during engulfment of apoptotic Nurse Cells (NCs) but is not required for engulfment. Another pro-inflammatory cytokine, Eiger (TNF-like), was shown to be required for cell clearance but it was not determined whether it is required by the FCs or the NCs. Further, the fate of follicle cells after engulfing nurse cells is still unknown. We are characterizing the response of hemocytes when cell death is induced in the ovary as well as investigating the signaling pathways used by professional and non-professional phagocytes to maintain homeostasis during the complete removal of dying cells.

250C The role and regulation of metabolic enzymes astray and Nmdmc during infection Krista Grimes1, Esteban Beckwith1,3, Marc Dionne1 1) MRC Centre for Molecular Bacteriology and Infection, and Department of Life Sciences, Imperial College London; 2) Instituto de Fisiología, Biología Molecular y Neurociencias (IFIBYNE), UBA-CONICET, Buenos Aires

Immunity and metabolism are closely interconnected processes. Mounting an immune response is energetically costly, and therefore relies on metabolism adapting to provide the necessary energy and raw materials. The aim of this work was to further characterise the metabolic changes that occur in Drosophila during bacterial infection. Using Targeted DamID, we found that two metabolic genes with linked roles in amino acid metabolism, astray and Nmdmc, are upregulated in the fat body during infection. astray is a phosphoserine phosphatase that catalyses the final reaction in the de novo production of serine from glycolytic intermediates. Serine can be converted into glycine using a tetrahydrofolate cofactor, which is produced by a cycle of reactions involving Nmdmc. We found that in flies infected with Gram positive bacteria, the upregulation of astray and Nmdmc is dependent on both Dif and FOXO, key components of the Toll and insulin signalling pathways, which may act in a co-ordinated manner. Further, fat body knockdown of the transcription factor MEF2 leads to an increase in the expression of astray and Nmdmc in uninfected flies, suggesting that MEF2 may contribute to the repression of both genes in the absence of infection. We next sought to establish whether astray and Nmdmc play functional roles in supporting the immune response. Fat body specific astray knockdown results in a significant survival defect during Staphylococcus aureus infection, whilst bacterial numbers remain unchanged, indicating that astray contributes to host tolerance. In contrast, knockdown of Nmdmc in the fat body leads to a modest survival increase. Together, these data demonstrate that metabolism is transcriptionally fine-tuned in a manner that has complex impacts on the course and outcome of infection and indicate that regulation of amino acid metabolism via FOXO is critical to support host health during immune responses.

251A Endocrine regulation of metabolism and immunity in response to commensal and pathogenic bacteria Scott Keith1,2, Brian Lazzaro1,2, Brooke McCartney1 1) Department of Entomology, Cornell University, Ithaca, NY; 2) Cornell Institute of Host-Microbe Interactions and Disease, Cornell University, Ithaca, NY; 3) Department of Biological Sciences, Carnegie Mellon University, Pittsburgh, PA

Animals are continuously exposed to diverse populations of environmental microbes ranging from beneficial bacteria to harmful pathogens and must rapidly adjust energy metabolism and immune reactions in response. Circulating hormones can modulate the balance between interrelated physiological processes through effects on gene expression and cell biology. In Drosophila, ecdysone and insulin signaling sustain metabolic homeostasis, developmental growth, and immunity in the context of both commensal microbiota association and infection with pathogens. However we know little about the mechanisms of this regulation, including the extent to which these hormones facilitate immuno-metabolic crosstalk and how interactions with microbes shape that crosstalk. We have identified the neuronal synaptic plasticity factor Arc1, which is homologous to mammalian Arc/Arg3.1 proteins, as a novel host gene that supports larval metabolism and growth in response to microbiota perturbation. We found that Arc1 can function in the insulin-producing cells of the larval brain and the ecdysone-synthesizing prothoracic gland to support development in germ-free (GF) Drosophila, and that hallmarks of insulin and ecdysone signaling were perturbed in GF Arc1 mutants. We additionally discovered that flies lacking Arc1 exhibit altered antimicrobial peptide induction levels, increased microbial loads, and increased mortality after systemic infection with the bacterial pathogens Erwinia (Pectinobacter) carotovora and Providencia rettgeri. These data reveal an unexpected new role for Arc proteins, which have been previously studied exclusively in the context of the nervous system. Future work will test whether Arc1 regulation of ecdysone and insulin signaling affects immune responses to systemic infection. Ongoing research will directly investigate how the interplay between ecdysone and insulin signaling in the fat body regulates IMD
pathway function and restructures energy metabolism to ensure survival of infection, potentially at the cost of resistance to other stressors like nutritional challenge. Through this work, we will better understand the mechanistic principles of hormone-mediated metabolic and immune system regulation in the context of host-microbe interactions.

252B Domestication of a phage-encoded DNAse I by Drosophila Rebecca Tarnopol, Jaden Ha, Susan Bernstein, Kirsten Verster, Noah Whiteman University of California, Berkeley

Advances in genomics have illuminated a potential role of horizontal gene transfer (HGT) as a source of evolutionary novelty in animals. We discovered the horizontal transfer of the cytolethal distending toxin B (cdtB) gene from bacteriophage (phage) donors into the nuclear genomes of several insects, including drosophilid flies. cdtB is a major virulence factor found in Proteobacteria and Actinobacteria that encodes a DNAse I-type nuclease. CdtB homologs encoded by bacterial pathogens of mammals typically induce cell cycle arrest and apoptosis of eukaryotic host cells. In secondary endosymbiotic bacteria of pea aphids (Ca. Hamiltonella defensa), phage-encoded cdtB may protect the insect host against parasitoid wasp attack. We hypothesize that HGT-derived cdtB is playing a similar protective role in the Drosophila lineages that harbor it. How insects are able to deploy HGT-derived CdtB to potentially kill parasitoids without harming their own cells remains unclear. Addressing this outstanding question is essential to understanding how a virulence gene that targets eukaryotic cells became functionalized within a eukaryotic genome. We hypothesize that drosophilid hosts may mitigate CdtB toxicity using a number of strategies, such as limiting its expression to particular tissues, or that the protein has evolved to not target the host genome (i.e., through modified substrate specificity or interactions with native host proteins). Here, we used the UAS-GAL4 system to test toxicity of cdtB expression in Drosophila melanogaster, which does not natively encode cdtB. We conducted in vitro nuclease assays to determine the specificity of drosophilid CdtB towards methylated vs. unmethylated DNA. We further investigated mechanisms by which drosophilid flies may evade CdtB toxicity by expressing the toxin in yeast. Overall, our work illuminates how prokaryotic genes whose products evolved to kill eukaryotic cells are integrated into animal gene networks.

253C ShKT-domain-containing protein from parasitic nematode is toxic to Drosophila melanogaster Aklima Khanam Lima, Harpal Dhillon, Adler Dillman University of California Riverside

Entomopathogenic nematodes (EPNs) are insect-parasitic nematodes which kill insects quickly on infection. EPNs from the genus Steinernema release excretory/secretory proteins (ESPs) into the insect host at the early stages of infection and these ESPs are toxic to insects. These proteins actively participate in tissue damage and immunomodulation of the host during infection. One of these proteins is an ShKT-domain-containing protein. ShKT-domain-containing proteins are named after the venom of sea anemone Stichodactyla helianthus. The ShKT domain is found in diverse groups of animal venoms and forms a large group of family called the ShK superfamily. They are known as a potent K+ channel blocker and perform many critical roles in animal venoms e.g., immunomodulator, neurotoxic, paralytic, hemolytic, etc. Our goal is to study the function of the ShKT protein from S. carpocapsae nematode and characterize its role in host-parasite interactions. We generated a recombinant version of ShKT protein using yeast expression system. To characterize the activity of the protein, we used the fruit fly (Drosophila melanogaster) model system. We injected 100 ng of purified ShKT protein into fruit flies and observed ~50% mortality. In addition to mortality, we evaluated additional health metrics of the fruit flies using behavioral assays (negative geotaxis and chill coma recovery). We found that the health of fruit flies was significantly affected by the ShKT protein.

254A Nematode secreted PLA, displays toxicity and immunosuppression in Drosophila melanogaster Ogadinma Okakpu, Sophia Parks, Harpal Dhillon, Adler Dillman University of California Riverside

Parasitic nematodes are an ongoing problem for human health and agriculture. Key to their success in infection is the ability to evade or suppress host immunity. This immunomodulatory ability is likely due to the release of hundreds of excretory/secretory proteins (ESPs) upon infection. While ESPs have been shown to display immunosuppressive effects on a host, there are still gaps in understanding about the molecular interactions between individual proteins released and host immunity. Our lab has recently identified a secreted phospholipase A, (sPLA,) released from the entomopathogenic nematode (EPN) Steinernema carpocapsae, which has immunosuppressive and toxic capabilities in insects. Immunosuppression by Sc-sPLA, is characterized by its ability to suppress the toll pathway and phagocytosis which lead to a decrease in survival against bacterial infections, and an increase in pathogenic bacterial growth in an infection model. Toxicity by Sc-sPLA, is characterized by increased mortality, with the severity being both dose and time dependent. EPNs have homology to vertebrate-parasitic nematodes, and thus can be used as model systems for understanding how nematode ESPs alter host immunity. sPLA, enzymes specifically, are known to be modulators of host immunity via lipid signaling. Understanding how Sc-sPLA, suppresses D. melanogaster host immunity will help elucidate EPN immunomodulation at the molecular level and may increase our understanding of the role of lipid signaling in insect immunity, which is an understudied aspect of the insect immune response.
Galbut virus is a double-stranded RNA virus in the family *Partitiviridae* that is found in 100% of *D. melanogaster* populations but only in ~60% of individuals within those populations. Galbut virus infection produces no obvious phenotype, however, a small fitness cost could be sufficient to select for resistance, and multiple lines of evidence support the hypothesis that some flies are resistant to infection. Endogenous viral elements (EVE) are integrated viral sequences that represent potential mechanisms of viral resistance. Recently, a galbut virus EVE has been identified in European *D. melanogaster* populations. I hypothesized that the galbut virus EVE might confer resistance to galbut virus infection. To test this hypothesis, I created a line of flies homozygous for the EVE by crossing an inbred line containing the EVE, (Global Diversity Line N14) and a robust inbred line from the Drosophila Genetic Reference Panel (DGRP strain 517). To test whether flies with the EVE can become infected, I microinjected galbut virus EVE+ individuals and found that only a small frequency (less than 1%) of offspring of injected parents were infected. To test whether the EVE is responsible for resistance, I am using CRISPR/Cas9 to remove the EVE. By removing the EVE, experiments can be done in parallel with EVE+ and non-EVE flies that are identical otherwise to test whether the EVE is actually conferring resistance.

**256C Age-dependent antiviral immunity in Wolbachia-infected Drosophila melanogaster** Brian Kmicik, Casey Goltz, Lakhira Sheffield, Stanislava Chtarbanova The University of Alabama, Tuscaloosa, AL

*Wolbachia* (*Wolb*) is an obligate, maternally-transmitted, intracellular bacterium known to infect ~50% of all insect species on Earth, including *Drosophila*. One aspect of *Wolbachia*-*Drosophila* interactions is a phenomenon called “pathogen blocking,” which corresponds to *Wolbachia*-mediated protection of *Drosophila* against pathogenic viral infections. The genetic and molecular mechanisms underlying this protection are not fully understood and are thus an ongoing area of investigation. Moreover, a gap in knowledge exists about the nature of the *Wolbachia*-*Drosophila* relationship as a function of host age. This is important to address as the aging process itself affects host physiology, including immunity to viral infections.

We have previously shown that aging impairs the ability of *Drosophila* to withstand the pathological consequences of infection (the viral ‘tolerance’ mechanism) with the RNA-containing Flock House Virus (FHV). Here, using 149 (66 Wolb-free and 83 Wolb-positive) lines from the *Drosophila* Genetic Reference Panel (DGRP), we demonstrate that the age-dependent survival of FHV infection is a continuous trait, and is significantly modulated by the presence of Wolb independently of underlying lifespan. For several of the Wolb-positive DGRP lines, we established Wolb-free stocks using standard tetracycline treatment. Our analysis of one line so far, DGRP320, indicates that age-dependent decrease in survival of FHV infection observed in Wolb-free flies is suppressed when Wolb is present in the stock. Using RT-qPCR, we show no significant changes in Wolb 16S gene expression, nor in FHV load between young and aged, Wolb-free and Wolb-positive hosts. Our results indicate that Wolb possibly modulates disease tolerance to FHV with aging, independently of Wolb load.

**257A The Evolutionary Genetic Basis of Bacterial-Mediated Embryonic Lethality** Dylan Shropshire¹, Mahip Kalra², Seth Bordenstein² 1) University of Montana; 2) Vanderbilt University

*Wolbachia* are maternally-transmitted, intracellular bacteria that occur in approximately half of arthropod species worldwide. They can spread rapidly through host populations via the cytoplasmic incompatibility (CI) drive system. CI causes embryonic death when infected males mate with uninfected females, but offspring of infected females are rescued. Two proteins, CifA and CifB, underlie the genetic basis of CI and rescue, but how amino acid sites across these proteins contribute to CI and/or rescue remain unknown. Here, we employed evolution-guided, substitution mutagenesis on conserved amino acids to understand their relative contributions to CI and rescue. We report that amino acids in CifA’s N-terminal unannotated region and annotated catalase-related domain are important for both complete CI and rescue, whereas C-terminal residues in CifA’s putative domain of unknown function are solely important for CI. Moreover, conserved CifB amino acids in the predicted nucleases, peptidase, and unannotated regions are essential for CI. Taken together, these findings indicate that (i) all CifA amino acids determined to be crucial in rescue are correspondingly crucial in CI, (ii) an additional set of CifA amino acids are uniquely important in CI, and (iii) CifB amino acids across the protein, rather than in one particular domain, are all crucial for CI.

**258B Microbial Influence on Drosophila sechellia Fitness on Octanoic Acid** Jake Erley, Zachary Drum, Caroline Pitton, Joseph Coolon Wesleyan University

The fruit fly *Drosophila sechellia* has evolved to become a dietary specialist for the fruit of *Morinda citrifolia*, also known as noni. This specialization is unexpected because the fruit contains toxic volatile compounds, namely octanoic acid (OA) that kills other *Drosophila* species. Our prior research found that adult *D. sechellia* flies exposed to OA had widespread down-regulation of genes with immune system functions. This suggests possible interaction between the *D. sechellia* microbial community and toxin resistance. Previous studies in insects have demonstrated important roles of both gut
microbes and endosymbionts like Wolbachia on organismal traits including toxin resistance. In order to determine if gut microbes and/or Wolbachia contribute to D. sechellia OA resistance we made three fly lines: D. sechellia with normal microbe assemblage (gut microbes and Wolbachia), D. sechellia with normal gut microbes but no Wolbachia and D. sechellia maintained on antibiotics that have very few if any associated microbes. Ongoing experiments are testing OA resistance in these lines, identifying genome-wide gene expression differences among the lines with different microbe assemblages and 16S metagenomics analysis of the microbial community present in each of the different lines generated for this study.

259C  
**Kismet affects gut biomechanics, the gut microbiome, and gut-brain axis in Drosophila melanogaster** 
Chloe Welch1, Angelo Niosi1, Henry Vo1, Punithavathi Sundar2, Aliyah Penn3, Jeffrey Cavanaugh4, Prince Yadav1, Wendy Lee, PhD2, Mikkel Herholdt Jensen, PhD2, Eliza J. Morris, PhD2, Kimberly Mulligan, PhD1  
1) California State University, Sacramento; 2) San Jose State University

The gut microbiome may contribute to the pathophysiology of neurodevelopmental disorders (NDDs), yet it is unclear how NDD risk genes affect gut physiology in a manner that may alter bacterial colonization. We addressed this question using Drosophila melanogaster with a mutation in kismet (kis), the ortholog of the human autism risk gene Chromodomain Helicase DNA Binding Protein 8 (CHD8). We used kis mutant flies to examine gut biomechanics, gut microbiota, and to explore the connection between gut microbiota and behavior. To quantify changes in gut tissue mechanics, we affixed whole guts between two clips mounted on a high-precision force transducer and length controller, capable of measuring forces to micro-Newton precision. Our measurements revealed significant changes in the mechanics of kis mutant guts compared to wild-type, in terms of elasticity, strain stiffening, and ultimate tensile strength. To characterize the gut microbiomes, we used 16S metagenomic sequencing and found that loss of kis profoundly impacted the abundance of many bacterial taxa in the midgut. To investigate the putative connection between the gut microbiome and behavior, we treated kis mutant and control flies with an antibiotic and then evaluated courtship behavior. Depletion of the gut microbiome rescued courtship defects of kis mutant flies, indicating a connection between the mutant gut microbiome and behavior. In striking contrast, depletion of the gut microbiome in the control strain induced courtship defects. This result demonstrates that antibiotic treatment can have either positive or negative impacts on behavior, depending on the status of gut dysbiosis prior to treatment.

260A  
**The relationship between natural diet, microbiome, and life history in Drosophila melanogaster** 
Brittany Burnside, Sarah Gottfredson Brigham Young University

My project focuses on the relationship between natural diets, the microbiome, and *Drosophila melanogaster* life history traits. I am studying the intersection between these traits in order to better understand how dietary impacts on host microbiota alters *D. melanogaster* life history. As a starting point, I have measured the development rate and starvation resistance of a *D. melanogaster* CantonS stock with an unmanipulated microbiota (‘conventional’) when reared on 22 different, autoclaved diets (12 vegetables and 8 fruits). Eclosion time for each fly vial was measured every 6 hours until eclosion of all eggs was complete. Then, 5–10 day-old female flies were transferred in groups of 10 into vials containing 1% agarose, and their survival was measured every 4 hours until all flies in a vial were dead. There was significant variation in the starvation resistance and development rates of flies reared on the different diets. A tradeoff between these two traits was observed in flies reared on most of the diets, except for carrots, spaghetti squash, and limes. At the time of transfer, I also stored some flies for microbiome analysis, to determine the conventional microbiota composition of flies reared on each diet. Currently, I am working to sequence the microbiomes of these flies to guide my creation of a gnotobiotic bacterial mixture to use in follow-up experiments that will compare the development rate and starvation resistance of axenic (‘bacteria-free’) and gnotobiotic flies on these same diets. These follow-up experiments will allow me to test how the diet affects the life history of the flies, both dependent upon and independent of the fly microbiota. Diet-dependent life history variation in axenic flies will reveal which diets sustain fly development and maturity independent of the microbiota. Life history values and microbiota composition in gnotobiotic flies will reveal interactions between the diet and the microbiota, including which combinations maximize particular life history traits. Together, these findings help establish how and why natural variation in the diets and microbiota composition of *D. melanogaster* can contribute to life history variation in locally adapted fly populations. More broadly, understanding these factors will help establish an explanatory framework for how host-microbe interactions influence animal evolution.

261B  
**Molecular and transcriptional characterization of a physical niche mediating symbiotic gut microbiome colonization in Drosophila melanogaster** 
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A mechanistic understanding of gut microbiome colonization is fundamental to understanding the microbiome’s functions and its physiological relevance. Specific associations with certain microbiome species in a host’s gastrointestinal tract have been consistently observed among many animal species including fruit flies, mice, and humans. However, little
is known about how these specific microbiome species are chosen for stable colonization and how such specificity is determined.

We hypothesize that the host gut actively promotes stable microbiome colonization with appropriate species via the molecular production of the appropriate environment, termed the “commensal niche.” Maintaining the commensal niche for certain microbiome species can be physiologically important for effective food digestion, efficient nutrition absorption, and enhanced immunity.

Previously, we isolated the fly gut microbiome strain, *L. plantarum (LpWF)*, from a wild fruit fly. The strain preferentially colonizes the fly foregut region, in particular the proventriculus inner lumen, with high stability, suggesting a commensal niche for *LpWF* is produced by the fly. We took advantage of this strong colonization phenotype to ask how the commensal niche for *LpWF* is developed and maintained at the cellular and molecular levels. We applied bulk and single-cell RNAseq to profile the transcriptional changes in the spatially defined commensal niche over time to identify specific cellular and molecular components essential for appropriate microbiome colonization. We are evaluating the identified candidates through loss-of-function and gain-of-function experiments.

In summary, our study seeks to decipher the molecular basis of gut microbiome colonization and how it benefits the host. Such a mechanistic understanding is likely analogous to other host-microbiome interactions as a general paradigm and could enable the design of intervening strategies for a probiotic gut microbiome that benefits human health.

262C  Microbiota effects on climbing abilities in *w*1118 flies  
Tanner B. Call1, Paige E. Bonnette2, John M. Chaston3, Shaleen B. Korch4, Gerald B. Call4  
1 Arizona College of Osteopathic Medicine, Midwestern University, Glendale, AZ; 2 Biomedical Sciences Program, College of Graduate Studies, Midwestern University, Glendale, AZ; 3 Department of Plant and Wildlife Sciences, College of Life Sciences, Brigham Young University, Provo, UT; 4 Department of Pharmacology, College of Graduate Studies, Midwestern University, Glendale, AZ

It is known that the microbiota can affect motor function in *Drosophila melanogaster*. We undertook an experiment to test the effects of different bacterial mono- and combination-associations in *w*1118 in our novel climbing assay. This assay uses the TriKinetics Multibeam Monitor to record the location of 16 individual flies every second over a 20-minute period. Analysis of the data can reveal multiple climbing and motor parameters. Using the standard bleaching protocol, *w*1118 embryos were made axenic (germ-free) and were then mono-associated with *Acetobacter pomorum*, *Acetobacter tropicalis*, *Lactobacillus brevis*, or *Lactiplantibacillus plantarum*. Another group of *w*1118 flies were associated with a combination of all four bacteria (Combo). Interestingly, the Combo group had the most significant effects on climbing metrics compared to the axenic control group, including increases in average height climbed, number of climbs, and total height climbed, which were unchanged by any of the mono-associations. The amount of time spent by the flies in the lower tube portion was decreased in both the Combo and *A. tropicalis* groups. These two groups also had an increased amount of time spent in the upper tube portion, suggesting that *A. tropicalis* increases the ability and or motivation of the flies to climb to the top of the tube and stay there longer. Both of the *Acetobacter* mono-associations and the Combo group had increased total movements and passes through the middle of the tube, indicating that the *Acetobacter* mono-associations promote increased locomotion in general. Interestingly, the *L. brevis* and *L. plantarum* mono-associations had no effects on climbing. This detailed analysis of climbing behavior in this study reveals some interesting findings. First off, the combination of all four bacteria had the most effects on climbing, indicating that the mixed microbiota in conventional flies likely promotes substantial influences on climbing. This finding shows that the host climbing phenotype is determined by community interactions in the microbiota and not by one or more community members acting individually. Second, the *Acetobacter* species appeared to be the most significant contributors to the effects observed with the Combo group, while the *Lactobacillus* species had no effects on climbing.

263A  Effects of host genetic feeding preferences in shaping microbiota composition in *D. melanogaster*  
Caroline Massey, John Chaston, Maggie Johnson Brigham Young University, Provo, Utah

The microorganisms within a host, commonly referred to as the microbiota, play an important role in the development of an organism and their life history traits, including fecundity and lifespan. In the model organism *Drosophila melanogaster* host genotype can significantly alter the microbiota composition, meaning that host genotype can determine at least in part which microorganisms are present and how abundant they are. What is not fully understood are the mechanisms by which host genotype selects the microbiota composition. In this work, I seek to better understand how the genetic feeding preferences of *D. melanogaster* help determine host microbiota composition. I have been using an assay where I will measure the variation in the microbiota composition of different fly populations when they are given a choice of different microbes in their diet; or when no choice is provided. This work will contribute to our understanding of how host genotype influences the microbiota variation observed between genetically distinct organisms.

264B  The Influence of Lab Manipulated Fermented Fruit and Maternally Inherited Microbiota on Metabolic Phenotype  
Oluwatobi Fijabi1, Graham Jones1, Derek Maas4, Andrei Bombin2, Laura Reed1  
1 University of Alabama; 2)
The infectious and pandemic nature of obesity is known to have a multifaceted root cause and must be addressed in a multivariate manner. Recent efforts have implicated the gut microbiome to be correlated with obesity-related phenotypes in Drosophila. Our previous research explored the influence of the natural microbiota of rotten peach fruit consumed by Drosophila in the wild in comparison to the conventional corn-based lab diet. In this present study, we are particularly interested in peach (P) and strawberry (S) diets because of the presence of health-enhancing phytochemicals, dietary fibers, and unsaturated fatty acids. Therefore, we investigated the influence of microbiota in shaping obesity-related phenotypes in Drosophila by testing the influence of genotype, maternal, and dietary microbiota in rotten P and S diets in shaping life history and metabolic traits. Organic P and S fruits were naturally fermented for six days and assessed for microbial and phytochemical composition through 16S rRNA sequencing and GC-LS/MS, respectively. We sterilized embryos with hypochlorite and ethanol to eliminate maternal bacteria on the chorion before larval emergence and treated the fermented P and S diets with autoclaving and antibiotics. Autoclaving is a thermal and pressure treatment that transforms and denatures nutrients but eliminates all microbes present. Conversely, the antibiotics eliminate 99% of microbes without heat application, keeping the nutritional integrity of the food. Hence, we adopted a novel strategy of rearing larvae of three DGRP lines on the naturally fermented P and S diets simulating the wild environment and on autoclaved and antibiotic P and S diets, giving us insight into the interactions between these treatments and our measured phenotypes. We anticipate that our results will reveal similar microbial compositions in P and S diets in beta diversity and major taxa. We also predict that larvae raised on P and S diets will have significantly higher survival, weight, triglyceride, and faster developmental time which are obesity-related phenotypes compared to the autoclaved and antibiotic P and S diets. In sum, we expect to see significant interactive effects between diet, parental microbiome and fly genotype in shaping metabolic phenotypes of larvae.

**265C The influence of environmental factors on the composition of fruit fly microbiota.** Reese Hunsaker, John Chaston Brigham Young University, Provo, Utah

The microorganisms that are associated with a host are called the microbiota and can provide key benefits to their hosts. For example, the microbiota of the fruit fly Drosophila melanogaster strongly influences fly life history traits and evolution. Variation in the composition of fruit fly microbiota can influence or determine fly fecundity, life span, and starvation resistance. Importantly, we have observed variation in D. melanogaster microbiota composition in flies that naturally adopt different life history strategies, establishing a link between these fly characteristics. However, one key gap is that we do not know which processes determine variation in microbiota composition in wild fly populations. The goal of this research is to test if three possible environmental factors can influence the abundance of bacterial species in the D. melanogaster microbiota. To do this, I am rearing gnotobiotic flies, inoculated from birth with six bacterial species, to wild flies in conditions that vary by one of three major environmental parameters: temperature, humidity, and photoperiod. Then, I measure changes in whole body microbiota composition of adult flies. I have already performed several temperature replicates and these data showed a strong influence on fly microbiota composition. I still need to perform humidity and photo period replicates. Together, these approaches will show which environmental conditions have the greatest effect on fly microbiota composition.

**266A Evaluating Approaches for Bacterial Mono-association in Parkinson’s disease Model Drosophila melanogaster** Paige E Bonnette1, Gerald B Call2 1) Department of Biomedical Sciences, College of Graduate Studies, Midwestern University, Glendale, AZ; 2) Department of Pharmacology, College of Graduate Studies, Midwestern University, Glendale, AZ

Parkinson’s disease (PD) is a neurodegenerative disease commonly associated with motor symptoms. Additionally, PD patients often suffer from non-motor symptoms, including gastrointestinal issues such as constipation and gut dysbiosis. These gut manifestations in a neurodegenerative disease support the hypothesis of a gut-brain-axis, a bi-directional communication pathway between the central nervous system and the gut. To investigate this relationship, our lab has been performing bacterial mono-associations in a PD model (*park*25 Drosophila melanogaster). The *park*25 flies are an excellent PD model as they possess many similar phenotypes to PD, such as dopaminergic neuron loss and impaired motor function. To perform a mono-association experiment, the flies must first be made germ-free, or axenic. The most frequently used method of rearing axenic *Drosophila* is by embryo dechorionation. This process generally consists of rinsing embryos with a 0.6% bleach solution followed by sterile water rinses and placement onto a sterile diet. Following this, our approach to collect homozygous *park*25 axenic flies involved transferring sterile, homozygous pupae (identified by the absence of the Tubby phenotype) to new sterile diet tubes. To create gnotobiotic adult flies, we performed two bacterial inoculations: the first with the embryos and the second onto the sterile diet that the pupae were transferred on to. While this method did produce successfully mono-associated *park*25 flies, the adult survival rate was low. Therefore, we wanted to see if there was a more efficient and effective method to perform mono-association experiments. We began by maintaining axenic fly stocks of both our control (w1118) and *park*25 flies. Axenic status of the stocks was checked periodically throughout the experiments. To perform a mono-association from the axenic stocks, adult axenic flies were allowed to deposit embryos for three days on new, sterile food. When the adults were removed, the embryos and larvae...
present on the food were inoculated with *Lactiplantibacillus plantarum* or maintained axenic. Pupae were transferred from these tubes and placed into new sterile diet tubes without a second inoculation. Approximately 5-6 days post-eclosion, flies were homogenized, and bacterial colonies were cultured and counted to determine the average CFUs/fly. Despite there being one less inoculation event compared to the standard protocol, this method showed that the flies colonized at a similar level to previous experiments in our lab. Compared to our initial protocol, the axenic stock method with the single inoculation event colonized equally in 25% of flies (96.4%), or even better in *park* flies (266%). Preliminary results suggest that this new method is similar to the traditional dechorionation method with regard to eclosion, adult survival, and motor function (climbing ability), which will be reported at the meeting.

267V Transcriptional Profiling of Immune Priming in *Drosophila melanogaster* Kevin Cabrera1,2, Duncan Hoard1, Zeba Wunderlich1,2 1) University of California, Irvine; 2) Boston University

Studying the way organisms fight off infections is a universally useful endeavor: anything that is alive has the possibility of getting sick. The historical paradigm in immunology focused on the binary distinction between the innate and adaptive immune systems. Increasingly, this binary convention has been challenged by the observations primed innate immune responses in organisms with and without an adaptive immune system. This so-called “immune priming” helps organisms more effectively fight off a second infection after survival of an initial infection. Despite the wide prevalence of immune priming across many clades from plants to insects to mammals, much of the mechanistic work on this phenomenon has focused on cell-based mammalian models and lacked transcriptional characterization in insects. Here, we use *Drosophila melanogaster* to create a powerful, *in vivo* model for studying epigenetic control of immune priming. By infecting flies with an insect-derived strain of the Gram-positive bacterium *Enterococcus faecalis*, we modeled infection response and quantified enhanced survival in immune primed cohorts. We also tracked bacterial load over time and found preliminary evidence showing that the enhanced survival in primed cohorts does not correlate with differences in bacterial load. Using RNA-seq, we have tracked transcriptomic changes associated with immune priming in the primary immune organs of *D. melanogaster*, the fat body and hemocytes. Using differential expression analysis, we classified families of genes that remain activated throughout experiment, more efficiently re-activate upon re-infection, or are qualitatively unique to a primed immune response. Delineating the relative contributions of each of these mechanisms not only reveals the drivers of infection survival, but also suggests epigenetic mechanisms of gene regulation and tradeoffs between the immune response and other biological processes. By integrating gene expression with chromatin remodeling events and the effect of gene deletion on priming ability, we begin shortlisting regulatory elements that may be driving primed immune response. In this way, we unveil a concerted mechanism explaining immune priming in the fly.

268V Establishing the feasibility of *Drosophila melanogaster* as a model system for *Acinetobacter baumannii* infection Parvin Shahrestani, Maria Soledad Ramirez, Melanie Garcia California State University, Fullerton

*Acinetobacter baumannii* is an antibiotic-resistant bacterium of public-health concern. Current model systems used to study *A. baumannii* infection are limited. My objective is to establish the feasibility of the common genetic model organism, the fruit fly (*Drosophila melanogaster*), as an alternative for studying *A. baumannii* infection. Preliminary data revealed that *D. melanogaster* inoculated in the thorax with a pin-needle dipped in *A. baumannii* suspension had statistically lower survival than controls. Although, survival was still approximately 95%. Using a nanoinjector to apply higher infection doses resulted in further reductions in post-infection survival in a dose-specific manner. A strain of *A. baumannii*, AMA, thought to be more pathogenic in humans in comparison to the model strain A118 was seen to be more pathogenic against *D. melanogaster*. These results support the feasibility of using *D. melanogaster* to study this dangerous human pathogen. In upcoming work, we will distinguish between tolerance to and resistance against *A. baumannii* and test the impacts of antibiotics on post-infection host condition. Establishing *D. melanogaster* as a model to study *A. baumannii* infection will advance current understanding of antibiotic resistant mechanisms, immune defense evasion, as well as assist with drug development.

269V The role of host microbiota in aging of *Drosophila melanogaster* Courtney Mueller, Parvin Shahrestani California State University, Fullerton

Laboratory selection can cause large phenotypic differences in eukaryotic populations without the introduction of new mutations. A key potential contributor to an animal’s evolution could be the host’s microbiota, however this element has been largely ignored in previous laboratory studies. Indeed, many host traits that can evolve in response to laboratory selection, can also be influenced by the microbiota. Moreover, when populations undergo laboratory selection for divergence in health-related traits, they also become differentiated in their microbiota. What evolves first? Do host traits evolve before the host microbiota changes or does the microbiota change as a result of the evolution of the host phenotype? I hypothesize that if the host phenotype evolves before the microbiota, then populations that are selected for fast development will evolve before changes to the microbiota. Using laboratory selection, I will evolve fast-developing populations from slow-developing populations. Each generation, for at least 10 generations, I will monitor development time and host genetic control of the microbiota. I predict that when populations
of Drosophila melanogaster are selected for fast development, changes to development time evolve before changes to the microbiota. In other words, the microbial changes are a consequence of host evolution rather than a contributor to host evolution. Efforts to uncover the mechanisms of fast evolution are widely studied without the microbial component. Here is one of the first looks into how microbiota affect development time of D. melanogaster.

270V Metabolic regulation of blood progenitor homeostasis and heterogeneity by TCA cycle in development and immune response in Drosophila larvae

Immunity as well as metabolism are quite old and extensively focused and sort after fields, but their inter-dependency, as Immune- metabolism, is being advocated very recently. This crosstalk infers how under immune compromised state, metabolic shift occurs and metabolites (a-KG, Succinate, Fumarate, etc.) takes on the tasks of proliferation, differentiation and activation of the concerned immune progenitors/cells. Recent work from our lab also highlighted GABA (released from brain) can elicit distinct immune cell population, under wasp infestation, which is nowhere to be seen in homeostasis (Madhwal et. al., 2020). Through our work we would like to switch the conventional paradigm of looking at TCA cycle as just an “intermediary step in the glucose metabolism for energy production” to the “reservoir of cardinal immune metabolites”. Metabolites are well known to get exchanged between the cellular compartments, which brands them competent for being signalling molecules. And what could be the better place for studying metabolites than TCA cycle, which is source as well as sink for all the three macromolecules of life. Conventionally TCA cycle, as the name suggest, always considered to be a cycle, but is it actually a cycle or an important junction of various cycles, so as to facilitate the exchange as well as maintenance of constant concentration of each metabolite. If the later is true, then each of the metabolites generated here can be an independent signalling molecules and directs the heterogeneity of different system. Here with our thorough analysis of TCA cycle, we would like to shed some light onto the crosstalk between the intermediary metabolites, development of blood progenitors and the heterogeneity produced within.

271V Amyloid Beta Peptide Plays an Immune Role in Alzheimer’s Disease Pathogenesis

Alzheimer’s Disease (AD) is a neurodegenerative disease characterized by severe memory decline and cognitive impairments. In AD patients’ brains, aggregates of amyloid beta (Aβ) peptides, called amyloid plaques, are hypothesized to trigger innate immune responses that contribute to AD pathogenesis. Interestingly, recent studies have demonstrated that an increased level of Aβ protects the host against pathogen infections. Using Drosophila as a model organism, we find that the loss of the Drosophila APP homolog, APPL, leads to immune deficits against parasite infection, and that overexpression of Drosophila Aβ produces an inflammatory phenotype. These findings suggest that Aβ might be required for a successful immune response. We additionally aim to examine whether there is a correlation between Aβ aggregation and inflammatory responses and whether Aβ-mediated inflammation is associated with cognitive defects. We will produce flies that develop different levels of Aβ aggregation and examine the inflammation responses of these flies during a pathogen infection. In addition, a fly oviposition assay will be used to examine the cognitive functions of the Aβ-expressing flies. This study will help to shed light on how innate immunity and infection contribute to the development of AD.

272V Immune role of Drosophila melanogaster Kazal-type serine protease inhibitor CG14933

Serine protease inhibitors (serpins) exhibit major regulatory functions in the proteolytic cascades of both arthropods and mammals. In Drosophila melanogaster, two of the major immune responses, the Toll pathway and the melanization cascade, are tightly regulated by serpins. Despite their known function in immunity, only a few of the 30 genes predicted to encode for serpins have been characterized, and knowledge about their exact mechanism in which they control the immune response to prevent excessive activation is still scarce. In this study, we investigate the immune role of the previously uncharacterized Drosophila putative serpin CG14933 containing a Kazal domain. By using CG14933<sup>SK1</sup> flies, a mutant devoid of any CG14933 expression, as well as RNAi knockdown and deficiency models, we show that flies lacking the gene show an increased susceptibility to Pseudomonas entomophila infection but not to any other infections that we tested. This clearly indicates an involvement of CG14933 in Drosophila immunity, although CG14933 itself is not induced by infection. We demonstrate that CG14933 has no role in the regulation of major immune pathways and does not act as a neutralizer of bacterial proteases. Additionally, we found that in adult CG14933<sup>SK1</sup> flies, active phenoloxidase levels are significantly increased compared to wild type flies. We were able to rescue the sensitivity phenotype to P. entomophila infection by employing a double mutant devoid of both CG14933 and PPO1, the prophenoloxidase activated in early stages of infection. Moreover, by using tissue-specific RNAi-mediated silencing of CG14933 we reveal a significant role of the gene in the fat body of Drosophila. Altogether, we propose that CG14933 plays a role in negatively regulating the melanization response in Drosophila. Furthermore, we provide evidence of a possible link between excessive
melanization and increased susceptibility to *P. entomophila* infection.

**273V**  
**Gut barrier defect and hyperactivation of innate immune response in a *Drosophila* model of NGLY1 deficiency**  
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Mutations in human *N*-glycanase 1 (NGLY1) cause a congenital disorder of de-glycosylation (CDDG) known as NGLY1 deficiency. It is a multisystem disorder with symptoms including global developmental delay, lack of tears, movement disorders, and chronic constipation. However, the biological roles of NGLY1 and pathophysiology of NGLY1 deficiency are not well understood. The *Drosophila* homolog of human NGLY1 is encoded by *Pngl*. Loss of *Pngl* results in semi-lethality. Previous reports from our group have shown developmental and functional defects in *Pngl* mutant intestine due to impaired BMP signaling and reduced AMPKα level, respectively. However, the lethality in *Pngl* mutants is only partially explained by these affected pathways (BMP and AMPK). This suggests the potential contribution of other biological pathways to the lethality in *Pngl* mutants. To investigate this, we performed RNAseq on the midgut tissue of *Pngl* mutant larvae and observed mis-regulation of a number of gene categories. In agreement with previous studies, proteasomal genes constituted one of the top down-regulated gene categories. Notably, major up-regulated gene categories were related to immune response, leading to the hypothesis that hyperactivation of innate immune response might contribute to the lethality in *Pngl* mutants. Reducing the innate immune signaling by decreasing Toll and immune deficiency (IMD) pathways rescued the lethality of *Pngl* mutants by 20-23%. Decreasing the gene dosage of the forkhead box O family transcription factor Foxo (reported to induce immune genes expression in *Drosophila* midgut upon stress) rescued ~40% of lethality. We observed increased activation of Foxo in the *Pngl* mutant larval enterocytes. Fluorescently labeled dextran feeding assay showed gut barrier defects in *Pngl* mutant larvae. Pharmacological disruption of peritrophic matrix (a major component of the gut barrier in flies) by feeding polyoxin D to wild-type larvae showed mild lethality and increased expression of innate immune response genes. Taken together, our data present evidence for gut barrier defects and hyperactivation of innate immune response in a *Drosophila* model of NGLY1 deficiency.

**274V**  
**The Impact of Increasing Concentrations of Ragweed Pollen on the Innate Immune System and Allergic Response of *Drosophila melanogaster***  
Shaila Sachdev Princeton High School, Princeton, NJ

The global warming crisis, caused by human activities, not only affects the environment but also human health. The increase in average annual temperature leads to longer growing seasons and higher pollen concentrations leading to an increase in the development of asthma and atopic allergies. Ragweed pollen happens to be the primary allergen trigger around the globe and affects more than 25 million Americans. *Drosophila melanogaster* exhibits a similar inflammatory response as humans making it a useful replicate asthma model. Although fruit flies only possess an innate immune system, they are outstanding invertebrate model organisms because they have comparatively simple physiology which allows for easy manipulation while possessing the major organs relevant to asthma. For instance, their trachea is made entirely of epithelial cells and directly exposed to high oxygen pressure and environmentally produced reactive oxygen species (ROS). It is believed that these reactive oxygen species may be generated by pollen which plays a potentially major role in influencing inflammatory responses in the airway epithelium. The purpose of this research is important to understand the extent of ragweed pollen’s ability to induce an allergic reaction in a live biological species. To model this experiment, I will be varying the concentration of ragweed pollen that the fruit flies are exposed to in their food media during their third instar larvae stage. Healthy larvae are usually buried in their food, but upon oxygen deprivation -- due to increased pollen concentration -- they move to the surface, which suggests impairments of the tracheal system. This behavior will be quantified by measuring the percentage of larvae showing this behavior, which will provide information about the availability of oxygen to the epithelial tissue. If *Drosophila melanogaster* are exposed to ragweed pollen, then they should exhibit oxygen deprivation by remaining at the surface of the food media because ragweed pollen is a reactive oxygen species and presents major mediators of inflammatory responses in the airway epithelium.

**275V**  
**In vivo demonstration of polymorphisms in antimicrobial peptides shaping host-pathogen interactions***  
Mark Hanson, Lena Grollmus, Bruno Lemaitre EPFL

Antimicrobial peptides (AMPs) are key players in innate defence against infection. In *Drosophila*, a large array of immune peptides contribute to host defence downstream of the Toll and Imd NF-κB pathways. Our recent work using single and compound AMP mutations confirmed that AMPs can additively or synergistically contribute to combat pathogens in vivo. For instance, Drosocin, Attacin, and Diptericins collectively contribute to defence against *Providencia burhodogranariea*. However we also revealed a high degree of specificity wherein one AMP can play a major role in combatting a specific pathogen. We found a specific importance of Drosocin for defence against *Enterobacter cloacae*, and previous work has shown a primary role for Diptericins in defence against *Providencia rettgeri*. We also recently showed how AMPs shape the fly microbiome over aging, wherein AMP-deficient flies suffered increased microbiota load and community diversity.
mimicking flies lacking Imd immune signalling. Finally, we recently described how furin cleavage enables a polypeptide AMP gene to produce multiple distinct peptides under the regulation of a single promoter. Here we take advantage of newly-available mutations to dissect the specificity of these interactions in an isogenic genetic background. We confirm the Drosocin peptide specifically confers defence against Enterobacter cloacae, but also investigate a previously uncharacterized Drosocin peptide downstream of its furin cleavage site. This underspecified peptide specifically contributes to defence against P. burhodogranariea, while the classic Drosocin peptide does not in fact contribute to this defence. Moreover a Threonine/Alanine polymorphism in this underspecified Drosocin peptide mirrors the effect of gene deletion in defence against P. burhodogranariea. Thus the Drosocin gene confers defence against different pathogens using two distinct peptide products. At the same time, we dissect the contribution of the two Diptericin genes in defence against P. rettgeri and other bacteria. We confirmed previous findings that Diptericin A is specifically important in defence against P. rettgeri. Surprisingly, we also uncovered a highly specific interaction between Diptericin B and systemic infection by a member of the gut microbiome. This finding of alternate specificities for the two Drosocin genes in defence against environmentally-relevant microbes paints a picture for how the immune system encodes innate defences. We show not only how AMPs and their alleles are critical to defence against specific natural enemies, but also provide a needed example for how specificity in defence can be derived from the evolution of AMP gene sequence and structure.

276V Does varying investment in egg production modify immune defense in mated female Drosophila melanogaster? Kathleen Gordon1; Mariana Wolfner2; Brian Lazzaro1,2 1) Department of Entomology, Cornell University, Ithaca, NY; 2) Department of Molecular Biology and Genetics, Cornell University, Ithaca, NY; 3) Cornell Institute of Host-Microbe Interactions and Disease, Cornell University, Ithaca, NY

In D. melanogaster and many other species, female reproductive investment comes at a cost to immunity and resistance to infection. In previous work, we have shown that D. melanogaster females become more susceptible to bacterial infection within hours of mating and for at least ten days. During mating, the male transfers seminal fluid proteins that change many aspects of female physiology and behavior, including inducing a rapid and sustained increase in egg production. One of these seminal proteins, Sex Peptide (SP), is involved in reducing female immune capacity. At least some of this effect may be due to stimulation of egg production by SP, as female flies without a germline, and who therefore produce no eggs, retain high immune capacity upon mating. Production of yolk protein to provision eggs is one of the major demands of reproduction, and we hypothesized that the level of investment in yolk protein production might directly or indirectly trade off with immune capacity. First, we ask whether high levels of production of yolk proteins (YPs) in the fat body, occasioned by increased egg production, interferes with that same tissue’s ability to produce antimicrobial peptides (AMPs), the main effector molecules of the Drosophila immune system. In ongoing experiments, we are testing fertility and immune phenotypes of unmated or mated females with missense mutants in the three YP genes (Tanaka et al. 2021) and of null mutations in YP genes that we are generating with CRISPR-Cas9 genome editing. If removing YP production allows a higher immune response, we will test whether the SP-mediated increase in juvenile hormone (JH) levels in mated females is the basis for this tradeoff, as JH is known to stimulate YP production and decrease immune response. We will additionally test whether post-mating immune capacity is rescued by mutations that block early or late oogenesis. Together, these results can inform the mechanisms through which egg provisioning affects female immune response and generates a reproduction-immunity tradeoff.

277V Not quite FedEx: How are venom proteins packaged for delivery by the parasitoid wasp Ganaspis hookeri? Nicholas Bretz, Chris Lark, Nathan Mortimer Illinois State University

Parasitoid wasps are common pathogens of Drosophila melanogaster, and use venom proteins in order to overcome fly immunity. Venom proteins derived from the parasitoid wasp species Ganaspis hookeri alter the immune response mounted by immune cells known as plasmatocytes within infected D. melanogaster larvae. This venom activity is mediated by a unique venom-specific isofrom of the SERCA (Sarco/endoplasmic reticulum Ca2+-ATPase) calcium pump. Venom SERCA activity inhibits the calcium burst normally mounted by plasmatocytes following infection, rendering them unable to melanize the foreign wasp egg. The mechanism by which SERCA and other venom proteins are transported into the host is not completely understood, but preliminary evidence suggests that venom proteins are packaged into venom-specific vesicles known as venosomes. These venosomes allow venom proteins to gain access to plasmatocytes likely via the interaction between virulence factors present on venosomes and host factors on the surface of plasmatocytes. Ultracentrifugation of G. hookeri venom separates venom proteins into unique fractions. Experiments utilizing nanoparticle tracking analysis and dynamic light scattering suggest that two of these fractions are composed of a heterogenous mix of vesicles, further supporting the idea that venoms can be stored as cargo within venosomes. With this, we are using fluorescence labeling of putative venosomes, mass spectrometry and SERCA activity assays to further characterize the venom transport mechanism. Vesicle packaging and specified targeting is a burgeoning field of research with potential applications in drug delivery for disease treatment.

278V Microbiome remodeling influences Drosophila immune response across generations Krystal Maya-Maldonado, Nichole A. Broderick Johns Hopkins University
In insects, including *Drosophila melanogaster*, the microbiome can exert intergenerational effects on host physiological processes through impacts on reproduction and development. In addition, specific microbiota members have been shown to influence mating, reproductive behavior, and development across generations.

Relatively, previous studies in *Drosophila* and other insects have shown that immune status can be transmitted to progeny, a process referred to as transgenerational immunity. While several mechanisms of transgenerational immunity have been proposed, these have focused mainly on the transfer of an immunological experience to the offspring.

Immune status is also known to be influenced by the microbiome, but the role of microbiota in transgenerational immunity has not been as well studied. To address this knowledge gap, here we examined how interactions between a host, its’ microbiome, and gut pathogen (s) modulate host immunity and microbiome composition across generations. First, we analyzed how the parental microbiome influences outcome in offspring immunity. We used the well-standardized protocols in the *Drosophila* model to get germ-free flies (axenic (AX) flies), and we compared them with conventional reared (CR) flies, which contain a native microbiota. Parents from each condition (CR or AX) were challenged orally with *Pseudomonas entomophila*, and the offspring were reared to the next generation. In the offspring, females were challenged with *P. entomophila* orally and we recorded survival, pathogen bacterial load, microbiome composition, and gene expression to analyze how the offspring respond to the challenge. This strategy was applied across two generations (offspring adults in F1, and F2). We found that *P. entomophila* persist in food where F1 develops, causing changes in microbiome composition of both the food and in the offspring. Interestingly, when we challenge F1 adults with *P. entomophila* we observed a better survival in the offspring from parents that were challenged, although pathogen bacterial load is the same between the offspring from parents’ challenge and unchallenged. Our results suggest an immune tolerance effect is mediated by microbiome alterations across generations. Future studies will further characterize this mechanism of tolerance, as well as examine whether there have been genetic or transcriptional changes in the microbiome or pathogen.

**279V  Role of Juvenile hormone in mediating trade-offs between immunity and reproduction** Vanika Gupta, Brian Lazzaro Cornell University

Immunity and reproduction are processes that reciprocally limit each other in a wide diversity of organisms. Mating stimulates the production of juvenile hormone (JH) by *Drosophila melanogaster* females, and JH is immunosuppressive. Thus, *D. melanogaster* females suffer reduced immunocompetence as a consequence of mating. JH additionally promotes fecundity and also plays a vital part in nutrient mobilisation in insects. The *D. melanogaster* genome encodes two JH receptors, *germ cell expressed (gce)* and *methoprene-tolerant (met)*, which have different expression patterns and distinct functions. Previous reports indicate that *met* regulates fecundity while *gce* controls post-mating immunosuppression. Using tissue-specific knockdown of the two receptors *gce* and *met*, we further define the role of these receptors in determining the interaction between reproduction and immunity as mediated by the immunosuppressive properties of JH.

**280C  Comparative sex chromosome evolution in Drosophila robusta species group** Kamalakar chatla, Doris Bachtrog University of California Berkeley

Neo-sex chromosomes frequently arise within the Drosophila genus through fusions between the ancestral X and Y chromosomes with autosomes. Typically, newly formed neo-Y chromosomes degenerate and become heterochromatinized, and the gene loss creates selective pressure for the homologue (neo-X) to acquire dosage compensation mechanisms to maintain proper gene dosage in males. Systematic, comparative studies are needed to study the dynamics of sex chromosome evolution, including heterochromatinization of neo-Y and dosage compensation on neo-X. In order to dissect the molecular mechanisms and evolutionary pressures driving the differentiation of these neo-sex chromosomes, we generated high-quality reference genomes, including Y chromosomes, using Nanopore single molecule sequencing combined with Hi-C scaffolding for several species with neo-sex chromosomes of varying age. We sequenced *D. nigromelanica, D. melanica, D. robusta* and *D. lacertosa* whose neo-sex chromosomes formed between 4.6 MY to 15 MY ago. The assembled Y chromosomes sizes varied from 10 Mb to 65 Mb which is inversely proportional to the age of neo-Y chromosomes. Homology to the neo-X and gene density decreased with Y age. *D. nigromelanica* harbors the youngest neo-Y chromosome maintaining 59% of genes, while the oldest neo-Y chromosome in *D. lacertosa* has less than 5% of genes remaining. All four species showed gene decay with accumulation of repetitive DNA on the neo-Y chromosome, and tandem amplification of some protein-coding genes. The neo-Y chromosome of *D. nigromelanica* is almost double the size of the ancestral autosome due to massive acquisition of repetitive DNA and tandem amplification of protein-coding genes. The three species carrying the older neo-sex chromosomes have fully dosage compensated neo-Xs that acquired male-specific lethal (MSL) complex binding sites. In contrast, *D. nigromelanica’s* neo-X is not fully dosage compensated yet. Overall, our high-quality genome assemblies help to dissect the evolutionary trajectory of neo-sex chromosomes differentiation.

**281A  Effects of epigenetic silencing of transposable elements on local recombination rate** Yuheng Huang, Grace Yuh Chwen Lee UC-Irvine
A wide range of taxa shows a negative association between transposable elements (TEs) density and meiotic recombination rates across genomes. Such association is commonly thought to be caused by reduced efficacy of selection against TEs in regions of low recombination or direct modification of recombination rates by TEs. Much of the research on TE evolution has focused on the selection explanation, leaving the latter possibility largely unexplored. One possible mechanism by which TEs impact local recombination rates is through the modifications of chromatin environments. To reduce the selfish replication of TEs, hosts have evolved mechanisms to epigenetically silence them through the enrichment of heterochromatic marks at TEs. And it has been demonstrated that the enrichment of heterochromatic marks suppresses recombination initiation. Indeed, across the genomes, the heterochromatin environment is positively correlated with TE density and negatively correlated with recombination rates among different taxa. Therefore, we hypothesize that the TE-mediated heterochromatin enrichment can suppress the local recombination rate. To test the hypothesis, we aim to identify the associations between the distribution of recombination and epigenetically silenced TEs in three inbred strains with distinct TE insertion profiles. To measure recombination rate at a fine scale, we developed a novel approach that uses long-read sequencing to identify recombinant haplotypes in pooled individuals. We collected F1 offspring from two crosses and backcrossed them to the shared paternal line. For each cross, we collected more than 6000 F2 offspring and sequenced them with PacBio in a pool, aiming to identify crossover events using the long reads. This approach mitigates the need of sequencing individual flies in the traditional approach for constructing recombination maps. To benchmark this approach, we also sequenced 192 F2 flies individually with Illumina short-reads as well as in a single pool using PacBio long-reads to assess the sensitivity and specificity of our proposed new approach. We will then measure the enrichment of a repressive heterochromatic mark (H3K9me3) around the TEs and test their associations with recombination rates.

282B Evolution of Drosophila glue adhesiveness Manon Monier, Virginie Courtier Institut Jacques Monod, Université de Paris, CNRS, Paris, France

Bioadhesives display remarkable physico-chemical properties that allow living organisms to stick to a great variety of surfaces. Drosophila larvae produce a glue to attach themselves to a substrate for several days during metamorphosis. This glue is mainly composed of 7 Salivary gland secretion (Sgs) proteins and Eig71Ee protein. Alignments of Sgs1, Sgs3, Sgs7, Sgs8 and Sgs3bis genomic regions across diverse Drosophila species reveal that Sgs3 has a high rate of duplication and deletion. Contrary to most of Drosophila species that attach to wood, leaves, or dry fruit skin, Drosophila suzukii larvae attach to fresh fruit skin or into the soil. Drosophila suzukii is an invasive pest species and represents a threat for fruit crops worldwide. We found that D. suzukii and it’s closely-related species D. biarmipes adhere poorly to glass compared to D. melanogaster. Our current analysis suggests that this weak adhesion is due to a lower amount of glue produced by salivary glands. Our work paves the way for a better understanding of the genetic basis of glue adhesion and for future industrial applications.

Keywords: bioadhesives, Drosophila suzukii, adhesion, pupa, genes synteny

283C Reconstructing the evolutionary history and neofunctionalization of the ZAD-Znf chromatin regulator dwg Jack Jurmu1,2, Andrew Arsham1,2 1) Bemidji State University; 2) North Hennepin Community College

The ZAD-Znf family of genes is evolutionarily dynamic, expanding sporadically across insect lineages via duplication and retrotransposition. The family is characterized by the presence of a conserved N-terminal zinc-coordinating ZAD domain upstream of anywhere from one to over a dozen grouped C2H2 zinc finger domains, connected by a poorly defined “linker” domain. New or fast-evolving ZAD-Znf genes can become essential through neofunctionalization. For example, nicknack and oddjob, two genes within a cluster of five ZAD-Znf paralogs, are essential in Drosophila melanogaster and have recently gained a heterochromatic regulatory function. Deformed wings (dwg, aka zeste-white5 or zw5) and CG2712 are neighboring and paralogous ZAD-Znf genes on the X chromosome of D. melanogaster. Protein sequence analysis suggests that both have acquired new functions within the last 5 million years. While dwg plays an essential role regulating insulators and enhancers in development, little is known about its non-essential near neighbor CG2712. In collaboration with the Genomics Education Partnership, we annotated the coding sequences of the two genes in D. melanogaster and other Drosophilid lineages diverging up to 40 million years ago. We compared domain and sequence architecture in order to reconstruct their evolutionary history and investigate their recent neofunctionalization.

284A Horizontal transfer of an apoptosis-inducing toxin gene in an agriculturally destructive fruit fly genus Saron Akalu, Kirsten Verster, Noah Whiteman University of California - Berkeley, Berkeley, California

Horizontal Gene Transfer (HGT) is the movement of genetic material between species, as opposed to via transmission from parent to offspring (i.e. vertical gene transfer). A growing body of evidence underscores the role of HGT in animal evolution. We previously discovered HGT of an apoptosis-inducing gene, aip56, into the nuclear genomes of the tephritid fruit fly genus Bactrocera, a genus that includes destructive fruit pests. In this study, we characterize the timing and evolution of aip56 HGT in the Bactrocera lineage. We use PCR, Sanger sequencing, and analysis of publicly accessible genome assemblies to determine that aip56 was transferred into the nuclear genome of a Bactrocera ancestor ca. 63
myna, and is highly expressed in the fat body. Additionally, we analyze the structure of AIP56 in *Bactrocera* using Phyre2 and find that the C-terminus has a high-confidence match to a rhamnose-binding domain, which suggests an immune function. Our results suggest that HGT of *aip56* served an adaptive function that may have facilitated the success of this agriculturally devastating clade.


The Insulin/Tor signaling pathway is responsible for the uptake of glucose into cells and its metabolism. It has also been linked to cell growth, fat and protein metabolism, and longevity. Its dysregulation in humans plays a major role in Type II Diabetes, Cardiovascular Disease, and cancer. GlyP, a member of the Insulin signaling pathway, encodes for the glycogen phosphorylase enzyme which is responsible for glycogen breakdown in the cell by releasing glucose into the bloodstream. Previous evidence has indicated that a gene’s selective constraint is influenced by its protein connectivity and position in pathway, among other factors, so that genes with less molecular interactions and a closer location to the membrane evolve faster than those with more protein-protein interactions and located closer to the nucleus. We hypothesized that GlyP would exhibit high selective constraint in *D. busckii, D. hydei, D. kikkawai,* and *D. suzukii* when compared to *D. melanogaster,* the reference species, due to its high connectivity and downstream position in the insulin pathway. The GlyP gene was annotated in each species utilizing the Genomics Education Partnership (GEP) tool pipeline, namely, the UCSC Genome Browser, tBlastn, Blastp, the Gene Record Finder, and verified in the Gene Model Checker. Genes models were proposed based on synteny, RNAseq data and other lines of evidence. The protein alignment of GlyP in all species was examined. All species were within 97% similar to the reference species despite *D. busckii* being the furthest diverged species from *D. melanogaster.* This supports our hypothesis that highly connected genes, also known as hubs, are under high selective constraint even in species vastly separated by evolutionary time. The gene’s downstream position may also provide evidence that GlyP is highly conserved and a cornerstone enzyme in the insulin pathway.

**286C  A genome wide model for estimating DNA transposable element excision rates in Drosophila virilis** Stefan Cerbin1, Danny Miller2,3, Justin Blumenstiel1 1) Department of Ecology and Evolutionary Biology, University of Kansas, Lawrence, KS; 2) Department of Genome Sciences, University of Washington School of Medicine, Seattle, WA 98195, USA; 3) Department of Pediatrics, Division of Genetic Medicine, University of Washington and Seattle Children’s Hospital, Seattle, WA

DNA transposons are sequences that are capable of moving in the genome resulting in profligate DNA damage and genome instability. Estimating transposon mobilization rates and the damage they cause is important for understanding genome evolution and genome stability. *Drosophila virilis* has several strains with varying copy number of DNA transposons and differences in germline piRNAs profiles. Transposons become activated when females lacking the requisite piRNAs to unique transposons inherited paternally, resulting in hybrid dysgenesis. With these strains we can study global transposon mobilization to determine the rates of excision and the overall change in copy number. In this study, we use pooled long-read DNA sequencing to identify elements mobilized in dysgenic crosses. These reads will be analyzed to estimate an excision rate using a likelihood model for estimating global excision rates for DNA transposons. This model will incorporate specific DNA transposon insertions, family, and piRNA profiles as parameters. We predict that differences in piRNA profiles, DNA transposon family identity, location, and internal deletion status will jointly determine the excision rate.

**287A  Testing the Effects of Fast-Evolving Heterochromatic Genes on Euchromatic Transposable Elements in Drosophila** Leila Lin, Yuh Chwen G. Lee UC Irvine

Eukaryotic genomes consist of gene-rich, transcriptionally active euchromatin and gene-poor, silenced heterochromatin. The tightly packed heterochromatic regions make up at least 20% of the human genome and about one-third of the fruit fly (*Drosophila*) genome. Heterochromatic genes play a central role in maintaining the stability of the genome. Specifically, their protein products are involved in the epigenetic silencing and compartmentalization of repetitive sequences and transposable elements (TE) that are enriched in heterochromatin. Accordingly, we hypothesize that heterochromatic genes must be fast evolving in order to keep up with the rapidly-changing and oftentimes deleterious repetitive sequences and transposable elements in heterochromatin. Supporting this hypothesis, we have identified fast-evolving heterochromatic genes across sixteen species of Drosophila flies. We achieved this through performing copy number variation analysis, a phylogenetic analysis by maximum likelihood (PAML) to test evolution across long evolutionary time, and a McDonald-Kreitman test (MK) to test evolution across short evolutionary time. To test whether the fast evolution of these heterochromatic genes may also influence euchromatic TEs that are epigenetically silenced with enrichment of heterochromatic marks, we crossed flies with a mutation for the genes of interest to a reporter strain with red fluorescent marker next to with various families of TEs. Changes in the intensity of the fluorescence would provide information about the extent of epigenetic silencing of TEs in the euchromatic genome. Our study will connect how the fast-evolving gene-poor heterochromatin may shape the evolution of gene-rich euchromatin.
Essential genes, which lead to lethality if functionless, are conventionally presumed to be conserved and ancient, whereas young genes are considered dispensable and nongermane to organismal survival. New genes, which are evolutionarily young, have been found to have essential functions in diverse processes such as centromere targeting, gametogenesis, human brain development, reproduction, and protein diversification. As new genes retain characteristics of the evolutionary forces that engendered them, their structures, and their functions; they provide a unique opportunity to gain insight into the evolution of essential functions.

Empirical investigations of essential genes on a whole-genome scale via functional genomic experiments such as RNAi are laborious, expensive, typically available only for model species, and are hardly suitable for complex organisms such as humans or mice. This has resulted in a deficit of functional and phenotypic data for understudied non-model species, limiting our understanding of the evolution of essential gene functions. The intractability of creating knock-out lines for non-model species, or infeasibility in the case of complex organisms, has necessitated the ability to accurately and precisely predict essential genes in silico as candidates for functional interrogation and evolutionary analysis.

In this work, I will develop machine learning methods to utilize features extracted and patterns learned from D. melanogaster 3rd generation sequencing-based assemblies and phenomic data to predict and identify essential new genes in non-model Drosophila species as targets for experimental and evolutionary analyses. Specifically, I will develop deep learning and ensemble learning algorithms to predict essential genes in non-model species followed by gene age dating to identify new genes and their putative origination mechanisms. I will then conduct CRISPR/Cas9-based knockouts of candidate essential new genes to validate their essentiality by measuring lethality and fertility effects in the knock-out lines I create. This work will yield insight into the evolution and development of gene essentiality in both model and non-model species, as well as provide open-source computational software for predicting essential new genes in Drosophila species with the potential for application in other non-model organisms.

289C Extensive genome-wide homozygosity tracts reveal micro-environment population structure in Drosophila populations. Peter Andolfatto1, Clair Han2,3, Patrick Reilly2,4, Andrew Taverner2, Ana Pinharanda1, Sheel Chandra1, Kevin Deitz5, Daniel Matute6 1) Columbia University; 2) Princeton University; 3) HHMI Janelia; 4) Yale University; 5) American Museum of Natural History; 6) University of North Carolina Chapel Hill

Drosophila population samples are usually established as descendent populations of a single wild-caught female (“isofemale lines”). As a result, whole genome sequencing (WGS) data has generally been collected either from isogenized versions of isofemale lines or from pools of wild-caught individuals. Both methods represent a departure from the approach used for most other organisms, which is to sequence outbred diploid individuals. In a population genomic survey of Drosophila santomea, an island endemic sister-species of D. yakuba, we re-sequenced the genomes of 34 wild-caught individuals widely distributed across the island of São Tomé. Plots of population-level genome-wide diversity reveal nothing particularly remarkable. Despite this, closer inspection reveals unusually long runs of homozygosity (ROH) within individuals. Remarkably, a substantial proportion of these individuals (>75%) harbor more ROH than expected for offspring of matings at the level of 1st cousins or closer. The level of diversity between individuals greatly exceeds that of within individuals (Fst = 0.194), strongly suggesting non-random mating. This is particularly surprising given the small geographic scale over which these samples were collected and indicates the existence of a large number of extremely small micro-environments on the island. The general lack of WGS of outbred diploid individuals in Drosophila prevents the easy identification of such ROH patterns. Among the rare exceptions is D. sechellia, for which wild collected individuals exhibit non-random mating patterns similar to that of D. santomea. Simulations reveal that the most likely demographic model for both species is one of many extremely small populations (resembling isofemale lines) with low but non-zero migration between them. Our findings have important implications for the fine-scale structure of Drosophila populations and downstream population genomic inference of demographic and selection parameters.

290A SR drive and the evolutionary history of the Y chromosome in Drosophila simulans Cecile Courret1,2, David Ogerreau3, Amanda Larracuente1, Catherine Montchamp-Moreau3 1) University of Rochester; 2) Laboratoire Evolution Génome Comportement et Ecologie, CNRS UMR9191, Gif sur Yvette and Université Paris-Saclay

Meiotic drive is an infringement of the law of equal allele segregation into the gametes. In heterozygote individuals, the causal genetic elements prevent the production of gametes that do not contain it. Thus, they can spread through populations even if they are deleterious for the carriers. Because they induce sex-ratio bias, sex-linked drivers expressed in the heterogametic sex are an important source of genetic conflict, characterized by the evolution of suppressors which tends to restore a balanced sex ratio.

In Drosophila simulans, X-linked meiotic drivers disturb the segregation of the Y chromosome during male meiosis. The progeny of carrier male is mainly composed of females. The drive is caused by two X-linked elements, acting together. The first one lies within a segmental duplication, the second has been identified as the HP1D2 gene, a young
and fast-evolving member of the Heterochromatin Protein 1 gene family. HP1D2 accumulates on the heterochromatic Y chromosome in spermatogonia, strongly suggesting that it controls the segregation of sister chromatids through heterochromatin modifications.

In natural populations where the drivers have spread, they are neutralized by resistant Y chromosomes. We followed the dynamics of the resistant Y chromosomes in natural populations in relation to the dynamics of the driver. We observed the replacement of sensitive by resistant Ys within a handful of years in populations invaded by the drivers. To go further in the study of Y chromosome variation in this species we sequenced 21 iso-Y lines, their Y chromosomes came from different locations around the world and have different phenotypes. We confirm the very low nucleotide diversity among Y chromosomes in this species, which could be considered as a signature of recurrent genetic conflicts. While we identified a haplotype with fixed differences between sensitive and resistant Ys. The sequence similarity between the sensitive Y chromosomes suggests that they have a recent common ancestor. We also confirm the ancestry of the resistant lineage by examining Y-linked sequences in the sister species of D. simulans, D. sechellia and D. mauritiana. All together the molecular polymorphism allows us to retrace the demographic and evolutionary history of the Y chromosome related to the one of the species. We show that intragenomic conflicts can drive astonishingly rapid evolution of Y chromosomes.

291B Natural Selection Shapes Variation in Genome-wide Recombination Rate in Drosophila pseudobscura Kieran Samuk1,2, Brenda Manzano-Winkler1, Kathryn R Ritz1, Mohamed AF Noor1 1) Duke University; 2) University of California, Riverside

While recombination is widely recognized to be a key modulator of numerous evolutionary phenomena, we have a poor understanding of how recombination rate itself varies and evolves within a species. Here, we performed a comprehensive study of recombination rate (rate of meiotic crossing over) in two natural populations of Drosophila pseudobscura from Utah and Arizona, USA. We used an amplicon sequencing approach to obtain high-quality genotypes in approximately 8,000 individual backcrossed offspring (17 mapping populations with roughly 530 individuals each), for which we then quantified crossovers. Interestingly, variation in recombination rate within and between populations largely manifested as differences in genome-wide recombination rate rather than remodeling of the local recombination landscape. Comparing populations, we discovered individuals from the Utah population displayed on average 8% higher crossover rates than the Arizona population, a statistically significant difference. Using a Q10 analysis, we found that this difference in crossover rate was dramatically higher than expected under neutrality, indicating that this difference may have been driven by natural selection. Finally, using a combination of short- and long-read whole-genome sequencing, we found no significant association between crossover rate and structural variation at the 200–400 kb scale. Our results demonstrate that (1) there is abundant variation in genome-wide crossover rate in natural populations, (2) at the 200–400 kb scale, recombination rate appears to vary largely genome-wide, rather than in specific intervals, and (3) interpopulation differences in recombination rate may be the result of local adaptation.

293A Acetobacter to Lactobacillus Ratios within Drosophila Melanogaster Microbiota, Diet and Environment Across a Latitudinal Gradient Aubrey Johansen, Amanda Morrison, Emma Davis Brigham Young University, Provo, UT

The community of microorganisms associated with an organism (‘microbiota’) can be tightly linked to the evolution and life history variation of that organism in the wild. Therefore, patterns in the composition of these microbial communities are likely to be linked to the adaptive traits and evolution of their hosts. Our lab previously identified a pattern in the abundance of the two dominant bacterial groups in wild Drosophila from two locations in the United States: acetic acid bacteria (AAB) are more abundant at low latitudes and lactic acid bacteria (LAB) are more abundant at high latitudes. While this variation in microbiota composition can be attributed at least in part to host genotype, the likely role of environmental sources of microbes in determining the AAB:LAB ratio of wild flies have not been clearly defined. In this work we define how variation in environmental reservoirs of microorganisms are linked with variation in microbiota composition of the flies from two locations in the United States. We sampled flies, their diets, and soil from thirteen orchards located in the eastern United States and northern Utah, USA. We are currently working to sequence the microbiota of these samples and compare the LAB:AAB ratios between flies and their environmental substrates to determine if there is congruence in overall microbiota composition or abundance of specific microbial groups. Our early analyses on a subset of samples suggests the abundance of LAB in flies and their diets are coupled, although the initiating cause of this coupling has not yet been identified. Our continued work with these and additional samples will test if this pattern is reproducible in multiple locations and if other microorganisms also display covariation between flies and their diets. Together, these approaches will help define the role of environmental sources of microbes in determining microbial compositions in wild Drosophila populations.

294B Chromosomal Rearrangements in two populations of Drosophila yakuba Timothy Ranallo-Benavidez, Rebekah Rogers UNC Charlotte

Understanding how genetic mutations underlie population differentiation is an important question for evolutionary biology. Chromosomal rearrangements can produce new genetic sequences that lead to phenotypic differences
between closely related species. We have identified chromosomal rearrangements in two recently diverged populations of Drosophila yakuba from mainland Africa and the island of Mayotte. To understand the effect that a more contiguous reference sequence has on identifying chromosomal rearrangements, we also use a recent long-read Drosophila yakuba assembly. With this new reference we are able to identify many more mutations, many of which occur in centromeric regions.

295C  **Karyotype evolution - Insights from a D. melanogaster strain with unusual sex chromosome karyotypes** Duoija Li, Dhvey Gandhi, Yukiko Yamashita 1,2,4 1) Whitehead Institute for Biomedical Research; 2) Department of Biology, Massachusetts Institute of Technology; 3) Department of Chemistry, Massachusetts Institute of Technology; 4) Howard Hughes Medical Institute, Massachusetts Institute of Technology

Drosophila melanogaster utilizes the X/Y sex determination system where the X to autosome ratio determines sex. Their sex chromosomes, X and Y, each harbors a ribosomal DNA (rDNA) locus consisting of hundreds of tandemly repeated rRNA genes whose sufficient copy number and transcription is essential for cellular functions. Females carrying Y chromosomes are rare occurrences which usually results from low rates of nondisjunction events. Here we discovered a strain of D. melanogaster from geographically isolated Seychelles archipelago whose X chromosomes barely contains rDNA, necessitating all females to carry at least one Y chromosomes. We found that this strain has two kinds of Y chromosomes, an intact one (Y) and a truncated one (YS) that lacks the majority of the long arm, the latter of which cannot support male fertility. Investigating why such a non-functional Y chromosome may be maintained in this strain, we found that Y chromosome makes female sterile when present in multiple copies. Therefore, Y serves as the essential source of rDNA for female. Our work on the Seychelles strain may hint at how karyotype may evolve, eventually leading to the diversification of species.

296A  **Tandem duplications as targets of selection in local adaptation** Taylor Conway, Rebekah Rogers 1) University of North Carolina at Charlotte; 2) University of North Carolina at Charlotte

Tandem duplications are a source of genetic novelty that create new gene sequences, alter gene expression patterns, and modify existing genes. We have used computational and molecular methods to describe evolutionary impacts from recently diverged populations in Drosophila. We locate and compare tandem duplications in Drosophila yakuba populations from mainland Africa compared with Island populations on Sao Tome and the eastern island Mayotte. These two independent cases of island invasion clarify how tandem duplications respond during habitat shifts. We have identified 1,426 mutations in D. yakuba capturing 1,022 genes. New gene structures include whole gene duplication and chimeric constructs. We are exploring the role these play in population differentiation in island environments.

297B  **Chromosomal rearrangements as a source of local adaptation in island Drosophila** Brandon Turner, Theresa Miorin, Nick Stewart, Cathy Moore, Robert Reid, Rebekah Rogers 1) University of North Carolina at Charlotte; 2) University of Georgia; 3) Fort Hayes State University

Chromosomal rearrangements act as a source of genetic novelty by shuffling DNA throughout the genome. These mutations can produce chimeric genes, induce de novo gene formation, or alter gene expression changes for existing genes. Here, we explore how these mutations may serve as agents of evolutionary change as populations adapt to new environments during habitat shifts. We identify 16,480 rearrangements in mainland D. yakuba and two locally adapted populations of D. santomea and D. yakuba on Sao Tome. Three loci that are associated with signals of strong differentiation in D. santomea lie adjacent to UV resistance or DNA repair genes, suggesting that these rearrangements confer selective advantages in high altitude environments with greater UV stressors. Some 55% of these mutations are facilitated by TE insertions, and 28% are TE facilitated ectopic recombination. In D. santomea 468 mutations are associated with strong signals of differentiation from the mainland while in island D. yakuba we identify 383 candidates of local adaptation. A total of 49.4% of mutations associated with signals of local adaptation also show significant changes in transcript levels, suggesting that the adaptive value of rearrangements is related to effects on gene expression. Together, this survey of structural variation identifies key modes of evolutionary innovation that would be missed in SNP-based screens. This work offers a portrait of how these mutations appear and help organisms to survive during habitat shifts, furthering our understanding of evolutionary processes.

298C  **Genetic variation in recalcitrant repetitive genomic regions in Drosophila melanogaster** Harsh G. Shukla, Mahul Chakraborty, J.J. Emerson University of California, Irvine

About one third of the Drosophila melanogaster genome is heterochromatic and consists of repetitive sequences
like satellites, transposable elements (TE), ribosomal DNA, and occasional single-copy sequence. Despite its role in chromosome segregation, nuclear organization, and gene expression, much of the highly repetitive heterochromatin has been recalcitrant to assembly. This limitation has impeded delineation of genetic variation, evolution, and function of this crucial genomic region. To resolve the sequence of the repetitive genomic regions and map the genetic variation within them, we de novo assembled the genomes of two isogenic strains, the reference strain ISO1 and A4, using Pacific Biosciences highly accurate (HiFi) long reads and compared their assemblies. The euchromatin arms are gapless in our assemblies and provide a complete map of genetic variation in them. We incorporated ~8 Mb of new heterochromatin sequences into the chromosome arm scaffolds, including ~3.5 Mb of X pericentric heterochromatin containing rDNA. We also assembled ~15 Mb of Y Chromosome from the two strains, unveiling the first detailed map of genetic variation for this highly repetitive chromosome. We show that despite being prone to structural mutations, the repetitive regions of the D. melanogaster genome exhibit contrasting patterns of copy number variation across different gene arrays. For example, the size of the Histone cluster is similar (~560 kbp) between A4 and ISO1, whereas the X-linked Stellate (Ste) gene cluster shows striking variation between the two strains. The Histone cluster in both strains consists of ~110 copies, whereas A4 and ISO1 carry 192 and 11 tandem copies of Ste in the X euchromatin, respectively. The varying degrees of structural variation in these two gene clusters are likely because the Histone copy numbers are evolving under stabilizing or purifying selection, whereas the Ste copy numbers are shaped by an evolutionary arms race between X-linked Ste and their Y-linked suppressors Su(Ste). Furthermore, complete resolution of tandem arrays like the Histone cluster at nucleotide level offer an avenue for determining the relative roles of birth-and-death vs concerted processes in the evolution of such clusters. Our results not only show a detailed map of molecular genetic variation within the hitherto unassembled repetitive genomic regions, but also lay the foundation for comparative and functional genomics of complete D. melanogaster genomes.

299A  A tandem duplication in Drosophila melanogaster shows enhanced expression beyond the gene copy number David Loehlin, Caleigh Paster, Jeremiah Kim Williams College

Tandem duplicated genes are common features of genomes, but the phenotypic consequences of their origins are not well understood. It is not known whether a simple doubling of gene expression should be expected, or else some other expression outcome. We describe an experimental framework using engineered deletions to assess any contribution of locally-acting cis- and globally-acting trans-regulatory factors to expression interactions of particular tandem duplicated genes. Acsx1L (CG6300) and Acsx1R (CG11659) are tandem duplicates of a putative acyl-CoA synthetase gene found in D. melanogaster. Experimental deletions of the duplicated segments were used to investigate whether the presence of one tandem duplicated block influences the expression of its neighbor. Acsx1L, the gene in the left block, shows much higher expression than either its duplicate Acsx1R or the single Acsx1 in D. simulans. Acsx1L expression decreases drastically upon deleting the right-hand duplicated block. Crosses among wildtype and deletion strains show that high tandem expression is primarily due to cis-acting interactions between the duplicated blocks. Sequence and phylogenetic analysis suggest that the duplication rose to fixation in D. melanogaster and has been subject to extensive gene conversion. Some strains actually carry three tandem copies, yet strains with three Acsx1 copies do not have higher expression levels than strains with two. Surveys of tandem duplicate expression have typically not found the expected twofold increase in expression. This study suggests that cis-regulatory interactions between duplicated blocks could be responsible for this trend.

300B  Seasonal plasticity and adaptive fluctuations of gene expressions of D. melanogaster Yang Yu, Alan Bergland University of Virginia

Two major mechanisms for populations of short-lived organisms to respond to temporal environmental heterogeneity, such as seasonality, are adaptive tracking and plasticity. Theory predicts that transition from one mechanism to another will have detrimental effects on the populations, thus the genetic architecture between adaptive tracking and plasticity should have limited overlap. Seasonal adaptive tracking can be observed from allele frequency change at individual loci and at genome-wide levels. Although thousands of SNPs have been shown to shift in frequency repeatedly across seasons in Drosophila melanogaster populations, we still have limited understanding of the relative strength of evolutionary mechanisms that underlie seasonal adaptation. In this study, we test the hypothesis that there is distinct genetic architecture between adaptive tracking and plasticity by using genome-wide plastic gene expression profiles across the season. We first identify the genes that plasticly change in expression levels across the season to gain insight into the functional response to seasonal environmental changes. Second, we will combine publicly available eQTL profiles, genome-wide allele frequency data from multiple seasonal populations, with our seasonal gene expression data to test whether the eQTLs associated with plastic genes are de-enriched for seasonal SNPs. We created and reared a genetically controlled F1 fly population from 24h embryos to adulthood in an experimental orchard across 10 seasonal time points (May to Oct in 2019) to examine the differentially expressed (DE) genes across the season. We extracted whole-tissue RNA from 3-5-day-old adult female flies and prepared pooled libraries using bulk RNA barcoding method (BRBSeq). Our next step is to identify the plastic genes and test our hypothesis that the eQTLs associated with these genes are de-enriched for seasonal SNPs. We hope to provide insight into the general understanding of seasonal...
adaptation from an expression perspective, and how seasonal environmental heterogeneity maintains functional genetic variation at eQTLs across time.

301C  Shavenbaby as a model to link phenotypic and gene regulatory changes across *Drosophila* evolution  
*Tatiana Gaitan*¹, Kaelan Brennan¹, Julia Zeitlinger¹,² 1) Stowers Institute for Medical Research, Kansas City, MO; 2) The University of Kansas Medical Center, Kansas City, KS

Phenotypic variability resulting from changes in gene regulation is often mediated by changes at cis-regulatory DNA sequences, i.e., enhancers. However, how changes in DNA sequence at enhancers generate different phenotypic outputs across evolution is not well understood. Understanding how transcription factors (TFs) read out cis-regulatory information at enhancers has been challenging, in part due to technical limitations associated with mapping TF binding at their motifs with sufficiently high-resolution to study crucial facets of transcriptional regulation such as, motif arrangement (syntax) and cooperativity between TFs. In the *Drosophila* embryo, axis patterning signaling defines the expression pattern of Shavenbaby (Svb), a key regulator of epidermal cell shape, which then induces the expression of cellular effectors that specify actin-rich projections called trichomes. How Svb directs the transcriptional activation of its targets presents an excellent opportunity for evolutionary analysis of motif syntax because of the epidermal phenotypes that are strikingly different between species. Thus, to interrogate the syntax of Svb targets, we are collecting genomewide binding information in late-stage *D. melanogaster* embryos and applying our high-resolution ChIP technique, ChIP-nexus, to map TF binding at base-resolution. With these data, we observe high-resolution footprints of Svb at its own motif within known epidermal enhancers. To obtain predictive rules for Svb binding that we can apply in an evolutionary context, we are combining Svb ChIP-nexus data with BPNet, a deep learning model that uses DNA sequences to predict ChIP-nexus binding information. BPNet successfully learns Svb ChIP-nexus data and identifies motifs predictive for Svb binding, which map to known Svb enhancers, and uncharacterized regions that represent new candidate enhancers. Next, we are identifying suitable *Drosophilas*pecies for evolutionary comparisons by performing cuticle preparations, Western blots, and immunostaining. With the strongest candidates, we aim to compare changes in motif syntax, and cooperativity with candidate TFs at target enhancers. Svb ChIP-nexus data together with BPNet provide an unprecedented opportunity to analyze neutral and functional changes in TF binding, and will further our understanding of how gene regulatory logic modulates phenotypic output across evolution.

302A  Identification of Three Novel Paralogs of CG3795  
*Jaquelyn Hester*¹, Amanda Moy¹, Jayda Cavanaugh¹, Kaitlyn Schoonover¹, Kelly Alvarado¹, Stanley Guan¹, Evan Merkhofer³, Gerard McNeil⁶, Howard Granok⁶, Martin Burg³, Michael Foulk⁴, Christopher Ellison¹, Wilson Leung⁶, Cindy Arrigo⁷ 1) Rutgers University - New Brunswick, New Brunswick, NJ; 2) Grand Valley State University, Allendale Charter Township, MI; 3) Mount Saint Mary College, Newburgh, NY; 4) Mercyhurst University, Erie, PA; 5) Washington University in St. Louis, St. Louis County, MO; 6) CUNY York College, Jamaica, NY; 7) New Jersey City University, Jersey City, NJ

The CG3795 gene is located on the Muller A Element (X chromosome) in *Drosophila melanogaster*. The FlyBase Gene Summary indicates that this gene is involved in the breakdown of proteins (proteolysis) and exhibits serine-type endopeptidase activity. As part of an investigation into the expansion of the *Drosophila ananassae* Muller F Element (~19.1 Mb compared to ~1.3 Mb in *D. melanogaster*), we identified four features within the *D. ananassae* Oct. 2018 (AGI/DanaRS2) assembly that show significant sequence similarity to the *D. melanogaster* CG3795-PA protein. Two of the features were located on scaffold QMES02000001 (tentatively assigned to the Muller E Element based on syntenic analysis), and two of the features were located on scaffold QMES0200178 (tentatively assigned to the Muller D Element). Upon further analysis, one of the features on scaffold QMES0200178 was assigned as the putative ortholog based on the following criteria: highest protein alignment coverage (i.e., subject coverage), low E-value, and high percent identity. The other features were determined to be novel paralogs of CG3795 since the RNA-Seq data shows that these regions are being actively transcribed in *D. ananassae*. In addition, the BLASTX alignments of the *D. ananassae* genomic region surrounding each feature against the *D. melanogaster* CG3795-PA protein did not show any in-frame stop codons or frame shifts, thereby supporting the hypothesis that these features are protein-coding genes, not pseudogenes. The RNA-Seq data in *D. melanogaster* suggests that CG3795 is a male-specific gene, with it’s highest expression levels in the testis (modENCODE Tissue Specific RNA-Seq data and Developmental RNA-Seq data). In contrast, the modENCODE RNA-Seq data indicates that CG3795 is not expressed in embryos or in adult females. This CG3795 male-specific RNA-Seq data expression pattern can also be observed in *D. ananassae*. Comparative annotation of species closely-related to *D. ananassae* shows that the CG3795 ortholog and the three CG3795 paralogs are also present in *D. bipectinata*. Given the number of novel paralogs of CG3795 found in both *D. ananassae* and *D. bipectinata*, we expect neofunctionalization of these male-specific genes in these two species. Future investigations will try to identify the CG3795 orthologs and paralogs in other *Drosophila* species in order to better understand the evolution of this male-specific gene.

303B  The evolution of morphology at a single-cell resolution  
*Ella Preger-Ben Noon*, Yifat Yanku, Anna Urum, Stav Naky Technion - Israel Institute of Technology, Haifa, Israel

The evolution of animal body form results from genome divergence. How genome divergence is translated into the...
morphogenetic events that generate new phenotypes is one of the most challenging questions in modern evolutionary biology. We address this question by studying the evolution of male genitalia in Drosophilids. Many anatomical features of the genitalia of D. melanogaster and its close relatives exhibit striking differences in shape and size, providing an excellent system to study the genetic, molecular and developmental basis of phenotypic evolution. Genetic mapping studies revealed that these differences are affected by multiple loci, but the evolved genes and how they coordinately function to generate diverse structures remains unknown. The emergence of single-cell genomics provides an unprecedented opportunity to resolve these problems. Here, we use single-cell RNA-seq to generate gene expression atlases of genital discs of D. melanogaster and its sibling species D. simulans. This approach allows us to unbiasedly identify genes that are differentially expressed in evolved genital substructures between the two species. By combining unsupervised cell-clustering with published gene expression pattern data we obtained transcriptomes for each of the anatomical substructures of the genital disc and identified all the evolved genes within these substructures. Functional analyses using the powerful genetic toolkit of Drosophila will determine the relevance of these genes to male genitalia development and evolution.

304C New Transcript Formation in Hybrid Drosophila Rebekah Rogers1, Nicholas Stewart2, Cathy Moore1 1) University of North Carolina, Charlotte, NC; 2) Fort Hays State University, Hays, KS

The origin of new genes is among the most fundamental processes underlying genetic innovation. The substrate of new genetic material available defines the outcomes of evolutionary processes in nature. Historically, the field of genetic novelty has commonly invoked new mutations at the DNA level to explain the ways that new genes might originate. In this work, we explore a fundamentally different source of epistatic interactions that can create new gene sequences in hybrids. We observe “bursts” of new gene creation in F1 hybrids of D. yakuba and D. santomea, a species complex known to hybridize in nature. The number of new genes is higher in the gonads than soma. We observe asymmetry in new gene creation based on the direction of the cross. Greater numbers of new transcripts form in the testes of F1 male offspring in D. santomea female x D. yakuba male crosses and greater numbers of new transcripts appear in ovaries of F1 female offspring of D. yakuba female x D. santomea male. These loci represent wholly new transcripts expressed in hybrids, but not in either parental reference strain of the cross. We further observe allelic activation, where transcripts silenced in one lineage are activated by the transcriptional machinery of the other genome. These results point to a fundamentally new model of new gene creation that does not rely on the formation of new mutations in the DNA. These results suggest that bursts of genetic novelty can appear in response to hybridization or introgression in a single generation. Ultimately these processes are expected to contribute to the substrate of genetic novelty available in nature, with broad impacts on our understanding new gene formation and on hybrid phenotypes in nature.

305A More than molting: Ecdysone signaling in adult Drosophila Zachary Drum, Joseph Coolon Wesleyan University

Ecdysone signaling is critical for successful life-cycle transitions in developing holometabolous insects, but the role of ecdysone signaling in adult insects is understudied. Previous work in adult Drosophila melanogaster has found roles for ecdysone signaling in many different processes like stress responses, courtship behavior, reproduction, and lifespan, but the ecological relevance of ecdysone in adults is not well understood. Our work examining the transcriptional response of Drosophila sechellia to the toxic volatiles found in its host plant Morinda citrifolia has predicted both the ecdysone receptor (EcR) and the ecdysone induced transcriptional repressor blimp-1 to be involved in the transcriptional response to these compounds. Altering the expression of these genes in adult D. melanogaster alters survival from octanotic acid (OA) toxicity. When D. melanogaster and D. simulans flies are fed food containing the active form of ecdysone (20E), their survival significantly increases and when D. sechellia flies are fed 20E, their survival on OA significantly decreases. The expression of genes involved in ecdysone response and biosynthesis are significantly different in these different fruit fly species, so examining basal ecdysone titers, 20E titers in response to OA exposure, and the genome wide transcriptional response to 20E by RNA-sequencing will help us understand if the ecdysone signaling differs in these species. Understanding how ecdysone signaling relates to OA resistance in D. sechellia may help elucidate how D. sechellia has adapted to specialize on the fruit of M. citrifolia and provide an ecologically relevant role of ecdysone in adult Drosophila.

306B Comparative Analysis of Node Degree on Gene Evolution in the Insulin Signaling Pathway Abigail Myers1, Alyssa Koehler2, Annie Backlund3, Chinmay Rele4, Laura Reed5 1) The University of Alabama; 2) The University of Alabama; 3) The University of Alabama; 4) The University of Alabama; 5) The University of Alabama

A biological pathway is a network of nodes (genes) interacting with each other and other regulating molecules in the cell which determines gene expression and therefore overall gene function of the cell. The insulin signaling pathway is extensively studied and includes a group of genes that regulate glucose metabolism. This, in addition to the pathway’s high conservation across species, makes it a beneficial model to study the function and evolution of biological pathways. In a network, node degree refers to the number of connections a gene has to other genes in the pathway. Previous studies using computational methods found genes with high node degree to be less evolutionarily constrained than genes with low node degree in this pathway. These results contradict the accepted hypothesis that genes with high node
degree are under more selective constraint because connected genes have to adapt to any mutations that occur at the node. In this project we further investigate the impact of node degree on gene evolution using manually curated gene models across the Drosophila genus. We are looking at the ratio of nonsynonymous to synonymous mutations (dN/dS) of three genes with varying node degree. We chose to focus on GlyS, raptor, and Dsor1; which are genes with low, intermediate, and high node degree, respectively. A comparative analysis of the nonsynonymous changes in these genes across Drosophila species related to their node degree will provide a small-scale analysis of the impact of node degree on gene evolution using manually curated gene models. This analysis will grow as more genes in the insulin signaling pathway are annotated and analyzed in the future.

307C  **De novo suppression of a male-harming mitochondrial mutation in Drosophila melanogaster via laboratory passaging** Sarah A. Tomlin1,2, Vada Becker4, David M. Brinkley3, Ching-Ho Chang1,2, Harmit S. Malik1,2 1) Howard Hughes Medical Institute, HHMI, Chevy Chase, MD; 2) Fred Hutchinson Cancer Research Center, Basic Sciences, Seattle, WA; 3) University of Washington, Molecular and Cellular Biology, MCB, Seattle WA; 4) University of Washington, Molecular Medicine and Mechanisms of Disease, M3D Seattle, WA

Mitochondria are specialized double-membrane organelles of bacterial origin present in most eukaryotic cells. Mitochondria contain circular DNA (mtDNA) almost exclusively uniparentally (maternal) inherited through females in most plants and animals. Such uniparental inheritance has been hypothesized to result in the accumulation of ‘male-harming’ mutations that are neutral or beneficial to females due to lack of selection (the ‘Mother’s Curse’ hypothesis). Previously, we found one such exclusively male-harming mtDNA (G177S) mutation in the cytochrome oxidase II gene (COII) of *Drosophila melanogaster*. G177S led to significant impairment of male fertility at 25 and 29 degrees but no impairment in female function. Over five years, the G177S mtDNA mutation was passaged without selection in the w1118 nuclear background. We found that male fertility had recovered substantially but not to wildtype levels in this line, suggesting the possibility that a de novo nuclear suppressor of male infertility arose and swept to fixation in the laboratory. Intriguingly, restoration of male fertility appeared together with lower female fertility in flies carrying the G11S mtDNA mutation. To recreate the original male infertility phenotype, we take advantage of genetic backcrosses to a w1118 line not subject to selection because it carries wildtype mtDNA instead of G1777S mtDNA. We aim to identify the genetic basis of this de novo suppression by whole genome sequencing.

308A  **Maternal mRNAs underlie higher heat tolerance in tropical vs. temperate Drosophila melanogaster embryos** Emily Mikucki, Thomas O’Leary, Brent Lockwood University of Vermont, Burlington, VT

The earliest stages of embryogenesis are particularly sensitive to environmental change, relative to other life stages. This phenomenon has been attributed to the transcriptional silence of early embryos, yet transcriptional responses to environmental change have not been fully characterized in the developing embryo. Previously, we demonstrated that tropical *Drosophila melanogaster* embryos have greater tolerance to acute heat stress than temperate North American embryos, suggesting that there is adaptive genetic variation in embryonic heat tolerance. In order to elucidate the molecular physiological basis of genetic differences in embryonic heat tolerance, we sequenced mRNA from early *D. melanogaster* embryos from tropical isofemale lines that were collected from 5 populations around the globe (Mexico, St. Kitts, Ghana, India, and Guam) and temperate North American lines collected from Vermont, USA. Early embryos (0-1 h old) were exposed to heat stress (32°C, 34°C, or 36°C) or control conditions (25°C) for 45 minutes prior to RNA sequencing. Out of ~29,868 sequenced transcripts, 828 (2.8%) differentiated the transcriptomes of tropical vs. temperate embryos, and all of these transcripts significantly correlated with heat tolerance among the lines. Functional enrichment analysis indicated that tropical embryos had higher abundances of maternally loaded transcripts that encode proteins involved in the oxidative stress response. We also found that tropical and temperate embryos exhibited similar changes in the abundance of 4,534 gene transcripts in response to temperature. 671 of these transcripts were induced by heat shock, indicating that embryos were transcriptionally active in the face of heat stress. Overall, our data indicate that tolerance to environmental perturbation during embryogenesis involves the oxidative stress response, corroborating recent studies on the important role of redox balance in fly development. Further, we demonstrate that transcriptional activity in the early zygote is context dependent.

309B  **Evolutionarily young, gene-silencing piRNA: innovation in gene regulation or control of selfish genetic elements?** Peiwei Chen, Alexei Aravin California Institute of Technology

Every genome is colonized by transposons, a class of selfish genetic elements that can mobilize and replicate themselves within the host genome, causing DNA damage and genome instability. To cope with this, animals employ a small RNA-guided genome defense mechanism to silence transposons. Particularly, the piRNA pathway is responsible for the repression of transposons in animal germline. Though piRNAs have been reported to regulate targets other than transposons, the range of these non-transposon piRNA targets in different species remains poorly explored. Previously, we profiled the piRNA repertoire in the male germline of *Drosophila melanogaster* and found abundant piRNAs from a locus on the Y chromosome that target and silence a protein-coding gene on the X chromosome, *pirate*. Pirate is an evolutionary young gene that encodes a putative deSUMOylase. Intriguingly, we found that in
another Drosophila species, D. mauritiana, pirate is potentially repressed by another small RNA pathway, endogenous siRNA pathway, suggesting that two distinct small RNA-based silencing strategies were independently invented in recent evolution to regulate pirate. I will discuss our ongoing efforts in exploring whether pirate acts as a selfish genetic element and how different classes of small RNAs evolve to suppress pirate in the male germline.

310C Multi-trait genetic characterization of resistance to heavy metal stress Elizabeth Everman, Stuart Macdonald
University of Kansas

Genetic characterization of stress response often relies on the assessment of a single trait; however, single trait assessments do not fully encompass an individual’s response to stressors. We took a multi-trait approach to characterize the genetic factors that contribute to variation in resistance to copper toxicity by examining adult, developmental, and behavioral responses to copper stress. Copper is one of several common heavy metal pollutants that are leached into the environment through mining and agriculture practices. Copper is an especially interesting metal because it is required for normal physiological function and development at low concentrations, but at toxic levels copper exposure can lead to organ damage and failure as well as impairment of neurological function. Furthermore, our previous work has demonstrated that genes and pathways involved in detoxification of non-biologically necessary heavy metals (lead and cadmium) also contribute to variation in the response to copper toxicity. Using the approximately 1500 genetically stable strains that make up the Drosophila Synthetic Population Resource (DSPR), we previously identified 13 regions of the genome that contribute to adult physiological resistance to copper using quantitative trait locus (QTL) mapping. We are currently measuring developmental viability under copper stress in the DSPR, and preliminary QTL analysis indicates that the genetic control of adult and developmental copper resistance may be partially life stage specific despite the lack of a significant phenotypic correlation between the traits. Thus far, only one QTL is shared between the adult and developmental response to copper stress. The shared QTL highlights mekk1 as a potential candidate gene, and estimated DSPR founder haplotype effects on developmental and adult responses to copper at this QTL suggests that allelic variation for mekk1 in the DSPR may have similar effects on copper resistance in both life stages. In addition to multiple physiological traits, we measured variation in copper detection ability in 200 DSPR strains and found that the ability to detect copper at low concentrations in food is genetically variable. We also found that copper-tainted food consumption is negatively correlated with adult physiological resistance to copper stress, underscoring the importance of combining physiological and behavioral assessments of stress response. By measuring multiple traits related to copper stress in the same large panel of genetically stable strains, we will ultimately be able to reveal the genomic independence and interactions underlying multiple traits that shape the organismal response to chemical stress.

311A Discovering Zinc Resistance Loci via Extreme QTL Mapping Katherine Hanson1, Anthony D. Long2, Stuart J. Macdonald1,3 1) Department of Molecular Biosciences, University of Kansas, Lawrence, KS; 2) Department of Ecology and Evolutionary Biology, University of California at Irvine, Irvine, CA; 3) Center for Computational Biology, University of Kansas, Lawrence, KS

Many heavy metals such as zinc, copper and manganese are essential for cellular function and maintaining metal homeostasis is critical. When exposed to toxic levels of heavy metals, including essential metals, organisms can suffer deleterious consequences, including increased risk for cancer and organ failure. Since zinc is involved in many cellular functions, zinc toxicity can have widespread effects, leading to necrosis, inhibition of mitochondria, and impacting the homeostasis of other heavy metals. Zinc toxicity has been primarily studied via functional genetics, employing single-gene mutations and expression knockdowns in a small number of inbred lines. Because the response to zinc toxicity is a complex, polygenic, trait, our goal is to identify those genes segregating for allelic variation for zinc resistance using an unbiased genome wide mapping approach. Drosophila melanogaster is an ideal model to study zinc resistance since it has orthologs of critical genes involved in zinc homeostasis, such as MTF-1 (a transcription factor involved in metal response), zinc transporter (ZnT) proteins and has proved to be a successful model to understand other heavy metal response traits. To identify zinc resistance loci we employed extreme QTL, or XQTL, mapping, a powerful technique that identifies QTL via extreme phenotypic selection of a population. We established a large, outbred population by mixing hundreds of DSPR RILs (Drosophila Synthetic Population Resource Recombinant Inbred Lines), a set of advanced intercross RILs derived from 8 inbred founder lines. We raised animals from this population on media supplemented with toxic levels of zinc, sequencing the pool of surviving, zinc resistant females and a matching control population, replicating the experiment 12 times. We estimate that >4000 animals were tested per replicate, and in each case we selected the top ~7% of animals. At each position in the genome we estimated the founder composition from each pooled sample, identifying QTL as significant frequency shifts between control and selected populations. We succeeded in identifying seven QTL, including one that overlaps with a QTL previously identified for copper developmental viability. Our QTL encompass physical intervals between 320 kb and 880 kb and collectively contain 451 protein coding genes. We implicate MTF-1, Mekk1, a gene involved in cadmium toxicity, present within the shared copper/zinc QTL, and 5 genes found to be associated with zinc resistance in a Drosophila cell line study. For a majority of our QTL only 1 or 2 founder alleles show a substantial frequency change between the control and selected populations, implying that resistant and susceptible alleles are often rare. We identified a series of candidate genes using existing functional data, including
previous metal toxicity studies, and gene ontology, and will further test these via gut-specific RNAi knockdowns.

313C Correlating Regulatory Region and Genetic Evolution Chinmay P. Rele, Laura K. Reed The University of Alabama

Understanding change at an evolutionary scale is primarily done through accession protein and gene evolution; however, understanding how the regulation of those loci occurs would paint a clearer picture of how the proteome and genome interacts with the environment. The regulatory regions of genes are highly variable and important, for the timing of gene expression. Understanding the dynamics between the regulatory regions of an organism with its relevant effects in the genome, transcriptome, and proteome, can inform us of how the evolution of one plays a role on the other. We are studying the evolution of the regulatory regions of Insulin-like peptides (Ilps) and how their evolution correlates with other characteristics of the genes. Ilp is a good family to examine due to high duplication rates, and different mutation rates of those duplicates that occur within the Drosophila phylogeny. In this study, we use the hand-curated gene models of genes within the insulin Signaling Pathway of 28 Drosophila species generated by The Genomics Education Partnership, including the Ilp paralogs. We are using these annotations to anchor the genomic region, extract genomic regions upstream of the start codon, and run Multiple Sequence Alignments. We intend to correlate features within these islands of conservation with genomic features to be able to find an association between regulatory regions and the genes they regulate. The patterns identified within these islands of conservation upstream of genes are likely cis-regulatory elements, or regions where regulatory elements exist that affect the regulation of the gene. Patterns such as indels (insertions/deletions), sequence motifs, as well as overall conservation found in these islands upstream of the gene can be correlated with the metrics of the genes directly downstream, such as ratio of nonsynonymous to synonymous mutations dN/dS, intron size, indel abundance, isoform number, and expression patterns (among other genetic characteristics). We expect to see positive correlation between the evolution of this upstream regulatory region and the evolution of features of the cis-regulated genes. Understanding the evolutionary patterns of these upstream regions is paramount in understanding how regulation of genes occurs.

314A What shall we do with the melanogaster species group? Artyom Kopp1, Bernard Kim2, Amir Yassin2 1) University California, Davis; 2) CNRS Laboratoire Evolution, Génomes, Comportement et Ecologie; 3) Stanford University

The genus Drosophila contains many hundreds of species that can be reared in the laboratory. This feature, along with the prominence of D. melanogaster as an experimental model system, has contributed to the widespread use of Drosophila as a model for molecular evolution, evo-devo, and comparative genomics. Ever since the 1930s, it has been recognized that classifying Drosophila species in a scheme reflecting their relatedness as well as their degree of divergence was fundamental for comparative studies. Therefore, the taxonomic rank of species groups was introduced and gained wide usage in Drosophila research. Of the nearly 60 species groups, the melanogaster group was initially created for 7-14 species since much of the diversity of Australasian and Afrotropical regions was unknown at that time. Over time, this group came to encompass ~200 species, far exceeding the typical size of other groups, and was subdivided into 10 subgroups. Recent phylogenetic studies revealed that those subgroups intermingle with several other species groups; thus, maintaining a monophyletic melanogaster group would require expanding its size and diversity even further. Some authors suggested upgrading the earliest branching ananassae (27 spp.) and montium (94 spp.) subgroups to species-group level, redefining the boundaries of the melanogaster group to a more reasonably sized clade. This solution simplifies the discussion of evolution within each lineage, as was done for example by subdividing the newly created montium species group into 7 new subgroups and 13 species complexes. On the other hand, this approach breaks with a huge body of comparative literature that includes ananassae and montium species as members of the melanogaster group. The aim of this poster is to discuss the taxonomic complexity of the melanogaster species group and to present through a balanced debate the benefits and disadvantages of different solutions. We also show that the ananassae subgroup/group is not monophyletic as currently defined, requiring additional taxonomic revisions.

315B Modelling Satellite DNA organization Sherif Negm, Amanda Larraqueuente Department of Biology, University of Rochester

Repetitive DNAs comprise large portions of eukaryotic genomes. Satellite DNAs (satDNAs) are abundant tandemly repeated DNA sequences found near centromeres, telomeres, and on sex chromosomes. SatDNAs originate through polymerase slippage, recombination between repeat elements, or TE-mediated mechanisms. Arrays of satDNA repeats are highly dynamic over short periods of evolutionary time: they vary in copy number and organization through unequal exchange, and other processes. SatDNA array expansion is thought to decrease organismal fitness but the relative importance of processes shaping satDNA evolution in natural populations is poorly understood. Population genetics studies have primarily focused on studying estimating copy number variation in satDNA arrays, due in part to limits in empirical data, as the repetitive nature of satDNAs make them difficult to study in detail. Recent advances in DNA sequencing now make it possible to infer satDNA organization at the sequence level, providing a richer source of empirical information. Here we provide a novel population genetics approach to study sequence variation in satDNA arrays. We simulate the effects of mutation, unequal exchange, gene conversion, drift, and selection on satDNA array sequence, structure, and organization in populations in a forward simulation framework. We designed a new
probabilistic model for unequal exchange and gene conversion that takes into account sequence divergence between monomers in the repeat array. We have identified summary statistics that capture the variation in repetitive satDNA arrays independent of copy number. We use Bayesian inference and regression approaches to infer recombination rate and gene conversion from simulated data and empirical data from a natural population of Drosophila melanogaster. We show that our approach could be useful for understanding how mutation, recombination and drift shape satDNA arrays under neutral evolution and selection.

316C  Testing long-term evolutionary change and stasis in the pioneer factor Grainyhead Henry Ertl, Patricia Wittkopp
University of Michigan, Ann Arbor, MI

A comprehensive understanding of morphological evolution must entail studying both evolutionary change and stasis in the developmental mechanisms that produce organismal form. Grainyhead (Grh) is a transcription factor critical for epithelial cell formation in the epidermis of flies, mice, and nematodes, suggesting the developmental role of Grh is conserved over 700 million years. To determine whether the underlying molecular functionality of Grh is also conserved, we constructed transgenic flies and tested whether the nematode Grh ortholog could rescue the weak epidermis phenotype of flies lacking functional Grh. We then collected gene expression, chromatin accessibility, and Grh binding data to provide a mechanistic explanation for any divergence between these distantly related Grh orthologs.

317A  Rapid diversification shapes the evolution and function of sperm nuclear basic protein genes in Drosophila species
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Most eukaryotes deploy histones for genome packaging functions. However, many animal species accomplish tighter packaging of genomes in sperm using short, positively charged proteins, called sperm nuclear basic proteins (SNBPs). Although histones are ancient and highly conserved, SNBP repertoires have independent evolutionary origins, differ dramatically across animal lineages, and evolve rapidly in mammalian lineages. Here, we leveraged the sequencing of many Drosophila species’ genomes to perform detailed phylogenomic and genetic analyses of SNBP genes. SNBP genes independently arose in Drosophila species via duplications of HMG (high mobility group) DNA-binding proteins. We found that 11 of 13 SNBP genes have higher protein evolution rates (dN/dS) than 95% of D. melanogaster genes. McDonald–Kreitman tests revealed that five SNBP genes have evolved under positive selection in the D. melanogaster lineage. Surprisingly, we found that the evolutionary conservation of SNBPs is not correlated with functional importance. Several ancestral, strictly retained SNBP genes are dispensable for male fertility in D. melanogaster, whereas two recently evolved SNBPs are essential for fertility. Moreover, SNBP genes necessary for fertility in D. melanogaster have been lost in other Drosophila species. To study the evolution of SNBPs, we ectopically express orthologs of one essential SNBP in D. melanogaster and found they express at different stages. Our result suggests that this SNBP might acquire function in mature sperm post-individualization and become essential. Moreover, we found extensive duplication of protamine genes across Drosophila species, with recurrent expansions of these duplications (>70%) on sex chromosomes. Our population genetic and phylogenetic analyses highlight that one X-linked SNBP amplification in D. mauritiana is under recurrent positive selection. However, the same amplification is independently lost and does not show the signature of positive selection in two sister species of D. mauritiana. These results indicate that sex chromosome-linked SNBP duplicates may be involved in either inducing or suppressing X-versus-Y meiotic drive. Our analyses suggest that the rapid evolution of SNBPs might be a universal phenomenon in animals and partly driven by genetic conflicts during spermatogenesis.

318B  Molecular mechanisms underlying alternating cell polarity establishment in Scaptodrosophila follicle cells
Miriam Osterfield UT Southwestern

Eggshell shape differs substantially among different species of drosophilids, with particularly obvious variation in the number of dorsal appendages. In Drosophila melanogaster for example, two patches of follicle cells are specified to an appendage fate, resulting in two dorsal appendages. In contrast, the follicular epithelium of Scaptodrosophila lebanonesensis (or S. pattersoni) contains only one patch of appendage-fated cells, but this patch gives rise to 4-8 separate dorsal appendages. Previous work has shown that these appendages are formed by the elongation of an alternating set of cell-cell edges within the linear floor cell domain. In other words, there is an alternating left/right pattern, so if a given floor cell lengthens on its left edge, its neighboring floor cells lengthen on their right edges. This study reveals a major molecular mechanism underlying this alternating cell polarity. A series of candidate approaches led to the identification of several molecules that localize specifically to the elongating subset of floor-floor edges; these molecules include Bazooka/Par-3, atypical protein kinase C (aPKC), F-actin, and phosphatidylinositol-4-phosphate (PI4P). In vitro treatment of Scaptodrosophila egg chambers with a panel of phosphatidylinositol kinase or phosphatase inhibitors revealed that PIK93 potently abolishes alternating localization of both PI4P and F-actin and also abolishes dorsal appendage formation. Based on the known selectivity of PIK93, this strongly suggests a role for a type-III PI4 Kinase, namely PI4KI1beta/Fwd or PI4KI1alpha, in this process. Additionally, in vitro treatment of Scaptodrosophila egg
chambers with Latrunculin abolishes the alternating localization of both F-actin and PI4P. The apparent requirement of F-actin and PI4P for each other’s localization suggests that a positive feedback loop involving these components is important for selecting the subset of floor-floor edges that will elongate. Current work is focused on developing genetic tools in *Scaptodrosophila* to further examine this unexpected example of planar polarity.

319C  **Genomic analyses of new genes and their phenotypic effects reveal rapid evolution of essential functions in *Drosophila* development** shengqian xia University of Chicago

It is a conventionally held dogma that the genetic basis underlying development is conserved in a long evolutionary time scale. Ample experiments based on mutational, biochemical, functional, and complementary knockdown/knockout approaches have revealed the unexpectedly important role of recently evolved new genes in the development of *Drosophila*. The recent progress in the genome-wide experimental testing of gene effects and improvements in the computational identification of new genes (< 40 million years ago, Mya) open the door to investigate the evolution of gene essentiality with a phylogenetically high resolution. These advancements also raised interesting issues in techniques and concepts related to phenotypic effect analyses of genes, particularly of those that recently originated. Here we reported our analyses of these issues, including reproducibility and efficiency of knockdown experiment and difference between RNAi libraries in the knockdown efficiency and testing of phenotypic effects. We further analyzed a large data from knockdowns of 11,354 genes (~75% of the *Drosophila melanogaster* total genes), including 702 new genes (~66% of the species total new genes that aged < 40 Mya), revealing a similarly high proportion (~32.2%) of essential genes that originated in various *Sophophora* subgenus lineages and distant ancestors beyond the *Drosophila* genus. The transcriptional compensation effect from CRISPR knockout were detected for highly similar duplicate copies. Knockout of a few young genes detected analogous essentiality in various functions in development. Taken together, our experimental and computational analyses provide valuable data for detection of phenotypic effects of genes in general and further strong evidence for the concept that new genes in *Drosophila* quickly evolved essential functions in viability during development.

320A  **Resolving the evolution and diversification of a Hox-regulated pigmentation trait** Ivan D. Mendez Gonzalez, Mark Rebeiz, Thomas M. Williams 1) University of Pittsburgh, Pittsburgh, PA; 2) University of Dayton, Dayton, OH

A fundamental question in evolutionary biology is how the diversity of forms and colors that we observe in the animal kingdom originated. *Hox* genes are highly conserved genes often implicated in phenotypic evolution, and yet the molecular mechanisms of *Hox* gene evolution have been difficult to pinpoint. In *Drosophila melanogaster*, the *Hox* gene *Abd-B* regulates the production of melanin covering the A5 and A6 segments of the male’s abdomen. This trait evolved from a non-melanic ancestor and in the *melanogaster* species group, within which it was extensively diversified. We tested whether temporal changes in *Abd-B* expression have played a role during the origination and diversification of abdominal melanic pigmentation. We identified two *cis*-regulatory elements (CREs) in *D. melanogaster* that are necessary for *Abd-B* expression in the A5 abdominal segment during late pupal development. Deletion of either CRE disrupts the formation of melanic pigmentation, suggesting that they have partially redundant activities. To test for intraspecific differences in the activity of these CREs, we replaced the *melanogaster* allele with the orthologous region from species with different melanic pigmentation. Our preliminary data suggests that the function of one of these CREs is highly conserved. Our current model is that partially redundant CREs form a relay to maintain *Abd-B* expression over time, and that evolutionary changes in this process that construct and deconstruct these CREs may be involved in the evolution of abdominal melanic pigmentation. Furthermore, the identification of these CREs offers the opportunity to manipulate *Abd-B* expression across different species to causally test how animal morphology evolves in response to changes in the expression of *Hox*-genes. We propose that regulatory evolution of genes with multiple developmental functions, like *Abd-B*, should favor temporal rather than spatial changes, potentially limiting negative pleiotropic effects.

321B  **Germ granule analysis reveals conserved and diverse features among *Drosophila* species** Dominique Doyle, Bianca Ulrich, Bianca Ortega, Matthew Niepielko Kean University

The co-packaging of different mRNA types into macromolecular structures called ribonucleoproteins (RNPs) is a conserved strategy for the regulation of mRNA metabolism. In many animals, the formation of complex RNPs called germ granules is essential for the post-transcriptional regulation of mRNAs that are required for germline development, maintenance, and function. In *Drosophila*, germ granules are assembled at the posterior of the egg and are inherited by the primordial germ cells during embryogenesis. In *D. melanogaster*, mRNAs accumulate in germ granules by forming homotypic clusters, which are distinct aggregates that contain multiple copies of a specific mRNA type. Homotypic clusters in *D. melanogaster* are generated through a stochastic seeding and self-recruitment process, relying on *cis*-regulatory sequences found in the 3'UTR of germ granule mRNAs called “clustering elements.” We hypothesize that clustering elements may be susceptible to evolutionary changes, creating diversity in the abundance of mRNAs found in germ granules from different *Drosophila* species. To test our hypothesis, we first investigated the homotypic clustering of two germ granule mRNAs, *nanos* (*nos*) and *polar granule component* (*pgc*) in three *Drosophila* species. By
combining single molecule in situ hybridization (smFISH), super-resolution microscopy, and quantitative image analysis, we found that seeding and self-recruitment is a conserved process that generates homotypic clusters to enrich germ granules with mRNAs. Interestingly, we found that the mRNA content of nos and pgc homotypic clusters, as measured by absolute transcript number, were strikingly diverse among Drosophila species. Specifically, a 50% difference in the number of transcripts in nos and/or pgc homotypic clusters was discovered between D. melanogaster, D. virilis, and D. pseudoobscura. By employing computational modeling, we recreated the diversity in germ granule mRNA content from all three species. Our simulations suggest that a combination of factors, including differences in mRNA clustering efficacy, generates germ granule diversity. Currently, we are investigating if variability found in clustering elements underlies Drosophila germ granule diversity.

322C Reorganizations in the apical extracellular matrix underlie morphological diversification in Drosophila genital structures

Identifying the genetic changes that cause morphological differences between species is a major goal of evolutionary and developmental biology. While many groups have found success by investigating differences in pigmentation or the regressive loss of entire structures, we know less about the genes and pathways involved in the diversification of three-dimensional body parts. The posterior lobe in the Drosophila melanogaster clade is an ideal system to investigate morphological evolution—this genital structure exhibits staggering diversity among the Drosophila melanogaster subgroup, including the sister species Drosophila mauritiana and Drosophila simulans, and we can track its development by dissecting and staining pupal terminalia. Previous work has shown that the posterior lobe develops in Drosophila melanogaster as a result of cell elongation—individual cells span all the way from the base of the lobe to its tip—and posterior lobe morphology is controlled in part by the apical extracellular matrix (aECM) component Dumpy. We therefore tested whether the aECM also underlies posterior lobe diversification between Drosophila simulans and Drosophila mauritiana. By labeling the aECM with fluorescent lectins, we have found that it forms attachments to the posterior lobe and other genital structures during early development, and these attachments are more extensive in Drosophila simulans, the species with the larger lobe. We also found that the lobe-specific gene expression pattern for dumpy is expanded in Drosophila simulans, which suggests that these morphological changes are controlled at the level of transcriptional regulation. Finally, we have built cellular simulations of posterior lobe development to investigate the role of the aECM in cellular elongation—whether it exerts an active contractile force on these cells or simply functions as a passive scaffold—and whether that role has changed between species. Our results suggest that morphological diversity may be generated by alterations in extracellular matrix organization during development, and that we can find the genes controlling this process within quantitative trait loci associated with genital evolution.

323A Tracking Natural Variation in Tolerance to Transposable Elements Across Time

Transposable Elements (TEs) are fragments of selfish DNA that multiply and propagate throughout the genome. These transposition events can lead to DNA damage, which severely impacts the germ-line integrity of the host, and the fitness of their offspring. While the evolution of resistance via the suppression of TE replication has been well characterised, it has been suggested that tolerance—where germ-line fitness is maintained regardless of transposition—could also potentially evolve in the aftermath of a new TE invasion. The P-element, which first appeared in Drosophila melanogaster mid-way through the 20th century, is one of the best characterised eukaryotic TEs. As D. melanogaster had already been long established as a model organism by the time the P-element incursion occurred, this particular host/TE system provides us with an excellent opportunity to study potential variation in TE tolerance both pre- and post invasion. Previous quantitative trait locus (QTL) analysis utilizing variation in natural strains lead to the identification of 22 SNPs associated with greater P-element tolerance in bruno, a gene thought suppresses P-element induced ovarian atrophy. As founder strains were all established in research labs prior to the P-element invasion, this suggests that the observed variation in tolerance is due to standing variation. In order to develop a historical picture of TE tolerance over time, we performed whole genome sequencing of a panel of D. melanogaster lines collected from different points across the 20th century. We are using these data to determine what alleles were segregating at both the bruno locus and number of genes involved in double-strand break (DSB) repair prior to the invasion of the P-element, and how these allele frequencies changed in the aftermath. In addition to characterising TE tolerance, these historical genomes also give us the opportunity to track and compare other variants that have changed in frequency over the same time period due to other selection pressures, such as climate change and insecticide resistance.

324B Intralocus sexual conflict drives new gene evolution in Drosophila Deanna Arsala1, Shengqian Xia1, Shuaibo Han1,2, Daniel Sanchez1, Manyuan Long1 1) University of Chicago, Chicago, IL; 2) Zhejiang University, Hangzhou, China

Males and females of nearly all sexually reproducing species pursue divergent reproductive strategies to reach their fitness optima despite sharing the same genetic material. These differences can cause intralocus sexual conflict (ISC),
where the presence of a shared genetic trait increases fitness in one sex while decreasing fitness in the other sex. Theoretical and genetic association studies have suggested that ISC can be resolved through the modification of sex-specific gene expression or alternative splicing. However, we still have little empirical evidence of the genes involved in ISC and their direct impact on sex-specific fitness in evolution. A recent study from our laboratory showed for the first time that gene duplication can mitigate sexual conflict in Drosophila melanogaster. The generality of this case is unknown and presents an important problem.

We set out to understand whether intralocus sexual conflict can generally drive the evolution of new genes in Drosophila. We have conducted a CRISPR/Cas9 knockout screen of 36 evolutionarily young, sex-biased gene duplicates in Drosophila. These genes have sex-specific patterns of expression—having high levels of expression in the reproductive tissues of either male or female flies. We have partially assessed the function of half of these duplicates and found that loss-of-function mutations in half of these genes confer a significant reduction in either male or female fertility. Bulk and single-cell RNAseq experiments are being employed to identify the cell types and genetic pathways involved in mitigating sexual conflict in the reproductive tissues. Our preliminary analysis suggests that intralocus sexual conflict drives the evolution of new genes with an unexpectedly large proportion contributing to sex-specific fitness.

325C Identifying the epigenetic determinants of gene-by-environment interactions using Drosophila melanogaster diapause as a model Abigail DiVito Evans, Paul Schmidt, Mia Levine University of Pennsylvania, Philadelphia, PA

Genes and the environment interact to pace developmental transitions. The initiation, cessation, and pausing of such transitions typically depend on genotype-specific responses to environmental cues, i.e. gene-by-environment interactions (GxE). Here we investigate the epigenetic determinants of GxE in the context of developmental pausing. We exploit a seasonally- and geographically-variable reproductive arrest phenotype (“diapause”) in Drosophila melanogaster. Diapausing females suspend reproduction in response to low temperatures and short days that signal the start of winter. When warm temperatures and long days return in the spring, the arrested ovaries re-initiate egg development. This arrest is accomplished by extensive and coordinated changes in gene expression in the ovary. We hypothesize that epigenetic factors control the diapause gene expression program in the ovary and that this epigenetic control depends on genotype. Consistent with epigenetic regulation, inbred lines that lack genetic variation exhibit incomplete penetrance—only a portion of individuals in the same environment with the same genetic background enter diapause. To identify epigenetic factors that mediate gene expression GxE in diapause, we exploit genetic variation between inbred lines with differing diapause penetrance (a “high penetrance” line with 90% diapause and a “low penetrance” line with 5% diapause). Using Western Blots, we screened histone mark abundance of H3K36me1, H3K27me3, H3K9me3, and H3K4me3 between diapause and age- and temperature-matched reproductive ovaries. We discovered that H3K4me3 is depleted in diapause ovaries, but only in the high penetrance line. We predict that experimentally depleting H3K4me3 in the ovary of the high penetrance line will increase diapause penetrance while experimentally elevating H3K4me3 will decrease diapause penetrance. We predict no such effects in the low penetrance line. We will present the consequences of H3K4me3-depletion and -enrichment in both lines and determine whether H3K4me3-dependent penetrance is genotype-specific. Analyzing RNA-seq reads from these same ovaries, we will delineate those H3K4me3-sensitive genes and pathways that are also differentially regulated by the diapause gene expression program. Together, these data may reveal genotype-dependent epigenetic regulation of GxE that mediate an important adaptive phenotype.

326A Redox balance and the oxidative stress response following acute heat stress of the early embryo in temperate and tropical lines of Drosophila melanogaster Thomas O’Leary, Brent Lockwood University of Vermont, Burlington, VT

Early embryos are particularly vulnerable to acute heat stress compared to later life stages. But previous work on Drosophila melanogaster has shown that tropical lines are more heat tolerant than temperate lines, suggesting that selection has made tropical embryos more robust to heat stress. Heat-tolerant genotypes more highly express oxidative stress response genes compared to heat-sensitive genotypes. Beyond the clear benefit of limiting the accumulation of reactive oxygen species and oxidative damage, a robust oxidative stress response may be crucial to maintaining redox balance, which has been shown to be important for progression through early embryogenesis. Here, we characterized redox status and oxidative damage in early embryos (0 – 1 hr after egg laying) following acute heat stress. The response of the embryo was measured through (i) the ratio of redox couples (NADH:NAD+ & GSH:GSSG) as an indication of the general redox state, (ii) superoxide dismutase (SOD) activity to quantify the antioxidant response, and (iii) lipid peroxidation to assess the level of oxidative damage. This work aims to elucidate whether the oxidative stress response is a molecular mechanism underlying acute heat tolerance in early embryos, and therefore, whether maintaining redox homeostasis is critical to heat tolerance in D. melanogaster embryos.

327B Widespread effects of early embryonic thermal stress on morphology, physiology and performance across the lifespan in D. melanogaster Sara Helms Cahan, Andrew Stoloff, Katie Bora, Collin Brown, Caela Flanagan University of Vermont

Environmental temperature has profound effects on every aspect of the development and performance of poikilothermic
organisms, and thermal limits play an important role in setting species geographic distributions. The most obvious effect of temperature extremes is mortality, which is used to determine the extent of the thermal safety margin, the difference between environmental temperatures and the thermal limits of the organism. However, significant negative impacts on performance may occur well below the LT₅₀ value, particularly for thermally sensitive life stages. To investigate the importance of sublethal effects, we compared multiple aspects of survival, development and thermal performance of developing *D. melanogaster* Canton-S eggs or larvae exposed to unstrained or moderately stressful low and high developmental temperatures (18°C, 25°C and 30°C) after one hour, 24 hours, or 60 hours post-laying. One-hour embryos displayed widespread deficits that were largely absent in flies transferred later in development, with somewhat different elements of performance impaired under cold and heat stress. Adult-to-pupal survival of cold-stressed early embryos was 70% of survival in unstressed conditions but was not reduced at 30°C; however, heat-stressed embryos suffered a 45% mortality rate during the pupal stage, and of those that emerged successfully, 35% displayed a crumpled-wing phenotype that rendered them unable to fly, and 30% showed significant locomotion deficits in an adult climbing assay. Adult thermal performance was also impacted, with both cold-stressed and heat-stressed early embryos displaying a ~0.3°C reduction in critical thermal maximum, while cold-stressed embryos showed substantially reduced capacity to improve their critical thermal minimum through developmental acclimation. Altogether, these delayed effects of early thermal stress summed across the lifespan have the potential to reduce mean fitness below levels needed for population replacement, which may have important implications for prediction of population persistence under projected climate change.

328C  **Ultra Violet radiation tolerance between *Drosophila* species from São Tomé and Africa: Adaptation across *Drosophila yakuba* and *Drosophila santomea***  
James Titus-McQuillan, Rebekah Rogers UNC Charlotte

The genetic basis of phenotypic differences between species is among the most longstanding questions in evolutionary biology. How new genes form and the processes selection acts to produce differences across species are fundamental to understand how species persist and evolve in an ever-changing environment. Adaptation and genetic innovation arise in the genome by a variety of sources. Functional genomics requires both intrinsic genetic discoveries, as well as empirical testing to observe adaptation between lineages. Here we explore two species of *Drosophila* on the island of São Tomé and mainland Africa, *D. santomea* and *D. yakuba*. These two species both inhabit the island, but occupy differing species distributions based on elevation, with *D. yakuba* also having populations on mainland Africa. Intrinsic evidence shows genes between species may have a role in adaptation to higher UV tolerance with DNA repair mechanisms (PARP) and resistance to humeral stress lethal effects (Victoria). We conducted empirical assays between island *D. santomea*, *D. yakuba*, and mainland *D. yakuba*. Flies were shocked with UV B radiation (@ 302 nm) at 1650-1990 mW/cm² for 30 minutes on a transilluminator apparatus. Custom 5-wall acrylic enclosures were constructed for viewing and containment of flies. All assays were filmed for reproducibility. Island groups did not show significant differences between fall-time under UV stress and recovery time post-UV stress test. Preliminary evidence shows mainland flies are less resistant to UV radiation than their island counterparts. Further work exploring the genetic basis for UV tolerance will be conducted from empirical assays. Understanding the mechanisms and processes that promote adaptation and testing extrinsic traits within the context of the genome is crucially important to understand evolutionary machinery.

329A  **Genomic Benchmarks: A Collection of Datasets For DNA Sequence Classification**  
Petr Simecek, Katarina Gresova, Vlastimil Martinek, David Cechak, Panagiotis Alexiou Central European Institute of Technology, Masaryk University

Recently, deep neural network models have been successfully applied to identify functional elements in the genomes of *D. melanogaster* and other organisms, e.g., promoters [1], chromatin folding [2], splice sites [3]. Unfortunately, it is not easy to compare the quality of these methods since they use different data preprocessing approaches. In other fields, benchmarks datasets have been established as a gold standard for comparison, e.g., IMDB Sentiment for text classification, SQuAD for question answering, and CASP data for protein folding prediction [4].

We are proposing a collection of datasets that may serve as a benchmark for the classification of genomic sequences. The collection is distributed as a Python package 'genomic-benchmarks' that is distributed through The Python Package Index (PyPI). Each dataset is stored both as a list of genomic interval coordinates and DNA sequences. The package provides utilities for conversion between these two formats, data cleaning procedures and checks. Furthermore, it contains functions that make the training of a neural network classifier easier, like PyTorch and TensorFlow data loaders. We hope other researchers will use our datasets to evaluate the quality of their algorithms.

The package 'genomic-benchmarks' and demo notebooks on how to use it are available on GitHub:

https://github.com/ML-Bioinfo-CEITEC/genomic_benchmarks

[1] Umarov, Ramzan Kh, and Victor V. Solovyev. “Recognition of prokaryotic and eukaryotic promoters using convolutional...
330B Insights into D. melanogaster and D. simulans transcriptome evolution and complexity using transcript distance (TranD) Adalena Nanni$^{1,2}$, James Titus-McQuillan$^3$, Oleksandr Moskalenko$^4$, Francisco Pardo-Palacios$^5$, Sarah Signor$^6$, Srna Vlaho$^7$, Zihao Liu$^{8,9}$, Ana Conesa$^{8,10}$, Rebekah Rogers$^1$, Lauren McIntyre$^{1,2}$ 1) Department of Molecular Genetics and Microbiology, University of Florida, Gainesville, FL; 2) University of Florida Genetics Institute, University of Florida, Gainesville, FL; 3) University of North Carolina Department of Bioinformatics, Charlotte, NC; 4) University of Florida Research Computing, University of Florida, Gainesville, FL; 5) Dept. of Applied Statistics and Operational Research, and Quality, Polytechnical University of Valencia, Spain; 6) Department of Biological Sciences, North Dakota State University, Fargo, ND; 7) Department of Biological Sciences, University of Southern California, Los Angeles, CA; 8) Institute for Integrative Systems Biology, Spanish National Research Council (CSIC), Paterna, Spain

Alternative splicing is an important driver of phenotypic diversity in higher eukaryotes. Understanding how alternative splicing and variation in transcript structure diverge across species can provide insights into phenotypic divergence and speciation. Long-read sequencing of mRNA provides an opportunity to observe transcript structure. We present metrics of complexity and nucleotide-level descriptions of structural phenotypes that can be calculated within an individual transcriptome or compared across transcriptomes. Using this tool, we show how patterns of transcriptome complexity can be compared across species without depending on the identification of orthologs. We further demonstrate that distance metrics can be used to compare transcriptomes of the closely related species, D. melanogaster and D. simulans, and identify novel exons which we validate. We implement out metrics in a PyPi package TranD and in the open source bioinformatics for everyone platform Galaxy (www.Galaxyproject.org) which will empower a wide range of researchers to quickly identify minimum distance transcripts between species, interesting structural variants within species and genome complexity enabling deeper understanding of splicing mechanisms and transcript evolution.

331C Bacterial infection promotes transposable element activation in Drosophila species Sabrina Mostoufi, Nadia Singh University of Oregon, Eugene, OR

Transposable elements (TEs) are short, repetitive sections of ancient viral DNA capable of inserting themselves into new parts of a genome. This ability has allowed TEs to invade the genomes of nearly all organisms, where they make up 5-75% of the genomes of plants, animals, and bacteria. TE activation can also affect gene expression, mutation, and recombination, making them powerful contributors to the evolutionary process. Stressful stimuli can cause TE activation to increase, creating another link between TEs and evolution. Infection can be a particularly stressful event for organisms, leading to a cascade of gene expression changes. Several studies have illustrated that infection can cause increased TE expression, but it is unclear how these processes differ between host species and pathogen species. Here we test the hypothesis that infection causes elevated TE expression in Drosophila species and investigate how pathogen species influences that expression. We analyzed several RNA-seq datasets of varying Drosophila species, infection types (bacterial, fungal, viral), and tissue types. Our results find that infection status is significantly associated with TE expression, but that the magnitude of that effect depends on host species and infection type.

332A Prevalence of galbut virus in wild Drosophila melanogaster populations and to lab colonization Tillie Dunham, Mark Stenglein Center for Vector-Borne and Infectious Disease, Department of Microbiology, Immunology, and Pathology, Colorado State University, Fort Collins, CO, USA

Galbut virus is remarkably common in wild Drosophila melanogaster. Every population that has been tested contains some infected individuals. But on average only 60% of flies are infected and there is evidence that some flies are resistant to infection. To better understand galbut virus-host interactions in wild and lab-reared populations, we pursued three avenues of investigation. First, we quantified galbut virus prevalence in wild populations sampled from different geographic locations over 5 years. We found that galbut virus prevalence varied widely from 0-96%, even in populations sampled from nearby geographic locations. We also found that galbut virus load in individual flies can be binned into 3 phenotypes: high positives, with galbut virus RNA levels exceeding that of an abundant host mRNA, low positives, with galbut virus RNA levels >100,000x lower, and negatives, flies with undetectable infection. Second, we performed an experiment to assess whether galbut virus prevalence would change over time in multiple independent wild-caught populations reared in lab. This will allow us to assess possible fitness costs to infection by tracking changes in infection frequency from generation to generation. Finally, to better understand the low positive phenotype, we bleached eggs then placed individual eggs into vials to be reared to adulthood. We did this in parallel with unbleached eggs that were reared together to have a baseline comparison. This experiment will allow us to determine whether the low positive infection phenotype represents legitimate low-level infection or contamination from highly infected flies. Overall, these
studies contribute to a fuller understanding of the variability associated with infection by this exceptionally common persistent *Drosophila* virus.

333V  **The role of chromatin and DNA sequence changes in de novo gene origin** Logan Blair, Julie Cridland, David Begun, Artyom Kopp UC Davis

Although it’s clear that some genes originate *de novo* from nongenic sequence, the molecular mechanisms through which they first gain expression remain unclear. One hypothesis is that novel transcripts evolve more easily in regions with ancestrally permissive chromatin structure. Alternatively, *de novo* genes may evolve from ancestrally inaccessible regions and the transition to open chromatin is critical to *de novo* gene origin. To distinguish between these models, I used ATAC-seq to examine chromatin accessibility around recently evolved *de novo* transcripts. I compared chromatin states of accessory gland tissue between five *Drosophila melanogaster* strains polymorphic for *de novo* transcript expression, as well as two closely-related species in which orthologous DNA sequences are not expressed. I found that open chromatin in *de novo* transcript promoters was frequently not exclusive to genotypes in which the *de novo* transcript was expressed. However, the magnitude of chromatin accessibility was greater in lines expressing *de novo* transcript, and much greater than that of orthologous regions in related species. Next, to determine the location of cis-regulatory variants associated with *de novo* transcript expression, I generated RNA-seq libraries from accessory gland tissue in 29 *D. melanogaster* strains. I found that SNPs associated with *de novo* transcript expression frequently occur outside the regions of ATAC-seq peaks. Together these results paint a complex picture of *de novo* gene origin in which an increase in chromatin accessibility is required but frequently occurs in ancestrally open regions.

334V  **Experimental Evolution for Longevity Differentiation in *Drosophila melanogaster*** Karen Walsh, Parvin Shahrestani, Molly Burke 1) Cal State University, Fullerton; 2) Oregon State University

The evolutionary theory of aging proposes that the forces of natural selection start to decline after the first age of reproduction and continue to steadily decline until the last age of reproduction, where these forces stabilize at zero or become negligible. Past studies have used *Drosophila melanogaster* populations to show that gradually postponing the first age of reproduction, postpones the age at which the forces of natural selection begin to drop, and results in delayed aging and increased longevity. However, genomic studies involving longevity remain underpowered due to limitations in replication. Long-lived populations were created by progressively postponing the first age of reproduction from 14 days to 70 days. To maximize the statistical power of finding these candidate genes, we used ten replicate populations selected for increased longevity, maintained on 70-day discrete generation cycles, and ten replicate populations that are treated as matched controls, maintained on 14-day discrete generation cycles. We have completed twenty generations of laboratory selection for postponed reproduction. Populations selected for delayed reproduction had increased longevity, increased fecundity, increased desiccation and starvation resistance, and increased wet and dry body weight. Samples from these populations have been frozen/preserved for each generation of selection. We next aim to identify candidate genes involved in longevity using a genome-wide analysis of experimentally evolved short-lived and long-lived populations of *D. melanogaster*. Identifying regions of the genome that are differentiated between short and long-lived populations may provide candidate genes for human longevity.

335V  **Evolution of longevity and immunity differentiation in *Drosophila melanogaster*** Joshua Glowalla, Molly Burke, Parvin Shahrestani 1) California State University, Fullerton, CA; 2) Oregon State University, Corvallis, Oregon

Theoretically a result of declining forces of natural selection, aging is a decline in fitness characters. Previous research has established *Drosophila melanogaster* as a model system to study the evolution of aging. Investigations utilizing experimental evolution have revealed a few well-defined relationships between aging and other traits, but the evolutionary relationship between longevity and immune defense remains unclear. While some studies suggest an evolutionary trade-off between these traits, others suggest positive correlations. Through a 10-fold replicated experimental evolution study, we have created populations that are diverged in age-of-reproduction. Late reproducing populations have evolved increased longevity and also improved immunity against the fungal pathogen *Beauveria bassiana*. We will test these populations for divergence in immunity against other pathogens. Analyses of whole-genome sequences throughout the selection process will provide information both about causative alleles and about their trajectories of change. Moreover, whole-genome comparison of our populations will reveal genetic connections between longevity and immunity. Advances in healthcare and technology have reduced the impact of many diseases on human populations, but consequently the percentage of people living to late age have increased along with the number of people with age-related illnesses. Aging and associated changes to immunity have become global health concerns which are best understood in the context of evolution.

336V  **Trade-offs between cost of ingestion and rate of intake drive defensive toxin use** Tyler Douglas, Sofia Beskid, Callie Gernand, Brianna Nirtaut, Kristen Tamsil, Richard Fitch, Rebecca Tarvin 1) University of California Berkeley, Berkeley, CA; 2) Indiana State University, Terre Haute, IN; 3) University of Texas at Austin, Austin TX

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Animals that ingest toxins can become unpalatable and even toxic to predators and parasites through toxin sequestration. Because most animals rapidly eliminate toxins to survive their ingestion, it is unclear how populations transition from susceptibility and toxin elimination to tolerance and accumulation as chemical defense emerges. Studies of chemical defense have generally focused on species with active toxin sequestration and target-site insensitivity mutations or toxin-binding proteins that permit survival without necessitating toxin elimination. Here, we investigate whether animals that presumably rely on toxin elimination for survival can utilize ingested toxins for defense. We use the A4 and A3 Drosophila melanogaster fly strains from the Drosophila Synthetic Population Resource (DSPR), which respectively possess elevated and reduced metabolic nicotine resistance amongst DSPR fly lines. We find that ingesting nicotine increased A4 but not A3 fly survival against Leptopilina heterotoma wasp parasitism. Further, we find that despite possessing genetic variants that enhance toxin elimination, A4 flies accrued more nicotine than A3 individuals likely by consuming more media. Our results suggest that enhanced toxin metabolism can allow for greater toxin intake by offsetting the cost of toxin ingestion. Passive toxin accumulation that accompanies increased toxin intake may underlie the early origins of chemical defense.

337V Dietary utilization drives the differentiation of gut bacterial communities Chau-Ti Ting¹, Jia-Syuan Chen¹, Shu Fang² 1) National Taiwan University, Taipei, Taiwan; 2) Academia Sinica, Taipei, Taiwan

Gut bacteria have been suggested to play vital roles in the dietary detoxification, digestion, and nutrient supplementation of hosts during dietary specialization. The roles of gut bacteria can be revealed by comparing bacterial communities between specialist and generalist hosts. However, it is often difficult to determine whether bacterial community differentiation is due to host dietary adaptation or divergence. To address this question, we investigated the bacterial communities from two Araceae-feeding Colocasiomyia species and further performed a meta-analysis by incorporating the published data from Drosophila bacterial community studies. By comparing three types of specialists (Araceae-feeding, mycophagous, and cactopholic) with generalist flies, we detected the structural and functional differentiation of the bacterial communities between specialists and generalists. The structure differentiation showed that Bacteroidetes and Firmicutes inhabited specialists, while more Proteobacteria inhabited generalists. The functional differentiation revealed that amino acid and energy metabolic pathways were overrepresented in specialists and generalists, respectively. This differentiation is associated with the higher utilization of structural complex carbohydrates, utilization of proteins, demand for vitamin B12, and demand for detoxification in specialists than in generalists. These results reveal that the interaction of bacterial communities and hosts might consequently facilitate the dietary specialization and ecological adaptation of animals.

338V Identification of a pseudogene derived from Arr1 in D. ananassae Ishtar Olaveja¹, W. Hayden Kee², Evan Merkhofer³, Don Paetkau ⁴, Wilson Leung ⁵, Cindy Arrigo¹ 1) New Jersey City University, Jersey City, NJ; 2) California State University Stanislaus, Turlock, CA; 3) Mount Saint Mary College, Newburgh, NY; 4) Saint Mary’s College, Notre Dame, IN; 5) Washington University in St. Louis, St. Louis County, MO

To facilitate investigations into the expansion of the Drosophila ananassae Muller F Element (~19.1 Mb) compared to Drosophila melanogaster (~1.3 Mb), we are manually annotating the protein-coding genes located on the D. ananassae F Element and a ~1.8 Mb region from the D. ananassae Muller D Element. As part of this analysis, we identified a region within scaffold QMES02000178 of the D. ananassae DanaRS2 genome assembly which shows significant sequence similarity to the Arr1 gene in D. melanogaster. Because past studies have shown that more than 90% of the genes remain on the same Muller Element across different Drosophila species, the D. ananassae scaffold QMES02000178 has been tentatively assigned to the D Element based on the protein-coding genes surrounding this Arr1 feature (i.e., CG14109, CG10725, CG10154, CG10713, CG10222, CG33263, flr). In contrast, the Arr1 gene is located on the Muller B Element (chromosome 2L) in D. melanogaster. Analysis based on the evidence provided by computational gene predictors (e.g., genBlastG, GeMoMa, Genscan, Geneid, and Augustus), RNA-seq data, and FlyBase BLASTP search results placed the putative ortholog of Arr1 on scaffold QMES02000007 in the D. ananassae DanaRS2 assembly. The D. ananassae scaffold QMES02000007 has been tentatively assigned to the B Element, suggesting that synteny for Arr1 is preserved between D. melanogaster and D. ananassae. The BLASTX alignments of the genomic regions surrounding the feature on the D. ananassae scaffold QMES02000178 and the protein sequence for D. melanogaster Arr1-PA include multiple frame shifts and in-frame stop codons, thereby supporting the hypothesis that the feature on QMES02000178 is a pseudogene derived from Arr1. The putative Arr1 pseudogene is also flanked by the DNA2-1_Dan and But2 DNA transposons. Future studies will identify and annotate additional pseudogenes located on the D. ananassae F Element and on the euchromatic reference region from the D Element, in order to assess the impact of the expansion of the D. ananassae F Element on the frequency of pseudogenization.

339V Frequent co-domestication of PIF-like transposable element proteins in insects Fatema ruma¹, Dragomira N. Markova¹, Claudio Casola², Ayda Mirsalehi¹, Esther Betran¹ 1) University of Texas at Arlington; 2) Texas A&M University

Transposable elements (TEs) are genetic units that move and amplify within a host genome. In recent years, an increasing number of studies have shown that one of the most direct contributions of TEs to their host is through the process of
molecular domestication' whereby the genes normally encoded by and serving the replication of a TE are co-opted by the host genome to create new gene(s) with cellular function. Thus, TE proteins are an important contributor to the emergence of novel host proteins. Despite the relative abundance of RNA TEs in eukaryotic genomes, DNA TE proteins and most notably their transposase, responsible for the excision and movement of the elements within the host genome, are generally considered to be more likely to be co-opted by the host than any other TE-derived protein. We have been studying four domesticated transposas from the PIF/Parabinger DNA family of TEs in Drosophila melanogaster, named Drosophila Pif Like Genes (DPLGs). So far, all PIF transposable elements known in plants and animals distinguish themselves from traditional DNA transposons by the presence of two independent transcription units. One encodes a protein representing the catalytic transposase, while the other encodes a protein with a MADF domain. We hypothesize that MADF proteins, a big gene family of regulatory proteins in D. melanogaster, to be derived from the same transposable elements. We also hypothesize that there should often be co-domestications of transposase and MADF proteins because the transposase translocates to the nucleus by the MADF protein. This is true for HARB1 and NAIF1 in humans, DPLG7 and DPM7 in Drosophila and two co-domestications in Arabidopsis (ALP1 and ALP2 and HPD1 and HPD2). To provide further support to this co-domestication model, we investigated numerous insect species genomes for additional evidence of PIF TE domestication events and explore the co-domestication of the MADF protein from the same TE insertion. We present evidence of at least five PIF TE domestication events in insects: two co-domestication of both transposase and MADF proteins in Anopheles (Diptera) and one transposase only domestication event and one co-domestication event Lepidoptera, and one transposase only domestication event in cockroaches (Blattodeae). Thus, our results show that domestication of PIF transposases is frequently accompanied by the co-domestication of a cognate MADF protein potentially for regulatory functions further supporting the common origin of both PIF proteins.

340V  Evolutionary diversification and repeated gene capture by telomeric retrotransposons across the Drosophila genus Jae Hak Son1, Mia Levin2, Christopher Ellison3 1) Department of Genetics, Human Genetics Institute of New Jersey, Rutgers, The State University of New Jersey, Piscataway, New Jersey, United States of America; 2) Department of Biology and Epigenetics Institute, University of Pennsylvania, Philadelphia, United States of America

Transposable elements (TEs) are mobile genetic elements that must replicate faster than their host to avoid extinction. TEs often evolve antagonistically with their host in a classic evolutionary arms race scenario. On the other hand, the co-evolution between TEs and the host genome can be mutualistic, where TEs are co-opted to benefit their host. Telomere-specialized non-LTR retrotransposons in Drosophila have traditionally been known for their mutualistic relationship with their host. These elements have replaced the role of telomerase, which is absent across the Drosophila genus, and replicate specifically to chromosome ends to protect them from erosion. However, more recent work has identified rapid evolution in many telomere binding proteins, which is more consistent with antagonistic evolution rather than a mutualism. Furthermore, we recently found that the D. melanogaster TART-A telomere-specialized transposon has captured a portion of the piRNA pathway gene, nxf2, which allows it to target nxf2 for suppression, again consistent with antagonistic evolution. Here we have examined the evolutionary diversification of telomere-specialized retrotransposons across the Drosophila genus using publicly available long-read genome assemblies from 106 Drosophila species genomes. We identify 9 major telomeric retrotransposon clades from the gage gene and 7 major clades from the pol gene, including the previously described HTT clade from melanogaster group and the TR2 clade found in the annassae and montium groups. We find that host gene capture among these telomere-specialized TEs is relatively common and propose that this phenomenon may serve as an important source of host gene regulation in the Drosophila genome.

341V  Analysis of eIF4E1 Conservation and Synteny across Drosophila Species to Understand the Evolution of the Insulin Pathway Jessica Strand, Jami Feist, Emma Sirjord, Sydney Payeur, Breanna Hoffman, Paula Croonquist Anoka Ramsey Community College

The Insulin/Tor signaling pathway regulates important physiological functions such as glucose and lipid metabolism, cell growth and survival. It is highly conserved from fruit flies to mammals. This pathway is also critical for homeostasis and its dysregulation results in prevalent human diseases such as in Type II Diabetes, Cardiovascular disease and even cancer. The eukaryotic translation initiation factor 4E1 (eIF4E1) encodes for a member of the eIF4F cap-binding complex that is required for cap-dependent translation of mRNA. The purpose of this study was to determine the conservation of the eIF4E1 gene in multiple species of Drosophila relative to D. melanogaster. It was hypothesized that the species highly divergent from the reference species would exhibit less eIF4E1 conservation than those closely related to the reference species. It was predicted that this gene would be most conserved in D. serrata and least conserved in D. arizonae. Gene models were proposed utilizing the UCSC Genome Browser, BLAST, Flybase gene record finder and verified by Gene Model Checker. The synteny analysis and protein alignments observed suggest that eIF4E1 in D. takahashii is the most conserved and D. arizonae is the most divergent. However, all five species had a high eIF4E1 protein identity ranging 81-94%. Previous evidence has demonstrated that position in the insulin signaling pathway may affect selective constraint, with downstream genes evolving at a slower rate than genes closer to the membrane. This may explain the high level of conservation for the eIF4E1 gene. Further studies should aim to investigate the impact of position, connectivity and/or
other factors shown to influence this pathway’s selective constraint.

**342V** Genome-wide relaxation and phylogenetic inertia of codon usage bias in the Neotropical *Drosophila saltans* species group *Carolina Prediger*,*1,2* Erina A. Ferreira*,2* David Ogereau*,2* Amir Yassin*,2* Lilian Madi-Ravazzi*1* 1) São Paulo State University (UNESP), Institute of Biosciences, Humanities and Exact Sciences, São José do Rio Preto, São Paulo, Brazil; 2) Laboratoire Evolution, Génomes, Comportement, Ecologie (EGCE), CNRS, IRD, Université Paris-Saclay, Gif-sur-Yvette, France

It is well established that *Drosophila* flies favor codons ending in C|G but relaxation in codon usage bias (CUB) pattern was reported for the *willistoni* and *saltans* groups, two Neotropical sister clades of the subgenus *Sophophora*. Those reports were, however, carried out with few genes (*saltans* group) or single genome (*D. willistoni*). Here, we generated whole genome sequences for 15 out of 23 species of the *saltans* group and evaluated CUB using a large set of genes in terms of the effective number of codons (ENC) and the relative synonymous codon usage (RSCU) index. We also investigated three forces that may be contributing to CUB, namely, mutational bias, selection pressure and drift. Our results indicated relaxation in CUB (mean ENC=48.33). Average RSCU index showed preference for codons ending in A|T. To evaluate mutational bias strength, we found moderate positive correlation between the RSCU of codons ending in G|C and the average G|C content at third position (GC3) for a given gene and a similarly moderate but negative correlation was seen for codons ending in A|T. Regarding translational selection, we found no correlation between tRNA adaptation index (stAl) and ENC after estimating tRNA diversity and content for four species for which genome assemblies have recently been published. We verified the strength of drift, by estimating the phylogenetic signal of RSCU on a phylogeny inferred from 30 genes, a heatmap with a hierarchical cluster analysis and a correspondence analysis. The first method indicated very strong phylogenetic signal for RSCU in the *saltans* group that was largely confirmed by the phylogenetic clustering of species by the second and third methods. In conclusion, our results demonstrate that relaxation of CUB in the *saltans* group is a general genome-wide pattern that has likely originated by mutational bias in the ancestor of the *willistoni* and *saltans* groups, and has since drifted along the evolutionary history of *saltans* species.

**343V** Resemblances Among Different Romanian Ecotypes of *Drosophila melanogaster* L. *Galia Butnaru*1, Ioan Sarac2 1) Banat University of Agricultural Sciences and Veterinary Medicine Regele Mihai I al României from Timișoara, România; Department of Genetics; 2) Banat University of Agricultural Sciences and Veterinary Medicine Regele Mihai I al României from Timișoara, România

19 *Drosophila* Romanian ecotypes were notice compared with 3 standard genotypes. The ecotypes were collected from geographically differentiated areas and anthropogenic polluted. The aim was to establish the relationship between the ecotypes evolved in the western, central and eastern areas of România. The multivariate analysis of variance revealed the significant differences between individuals [male/female] and among ecotypes. The male body average size represented 82.71% (2.92±0.05mm) compared to the female average size (3.53±0.06 mm). The variability of body size varied from 2.93 to 4.60mm on female and 2.43 to 3.83mm on males. Among female the difference of size averages was larger than among males (dF=1.67mm>dM=1.4mm). Even in this circumstance the male size varied from 80.2% to 84.0% at the same size of females. The results indicated that only 36.4% and 45.6% female and male, respectively, revealed a larger body size than the overall average size. It was found at 81.8% of ecotypes the influence of environment was small and insignificant. The multivariate analysis of variance regarding the participation of females and males at the general body size average of the ecotypes pointed out the population significant involvement of Black Ploșoru and Urdași (A and B) from Gorj County and Ebony, Timișoara-N, Șag (C) from Timiș County (LSD5% = 0.31 mm).

It was concluded that knowledge of quantitative traits, in our case the body size, is directly associated with adaptability to extreme conditions such as anthropogenic pollution and limited motility. Even if the fruit fly populations are not real isolated it has been identified inter-populations and intra-population phenotypic polymorphism. The presence different forms of *Drosophila* in the same niche Ploșoru village [Black Ploșoru (big and normal) and Grey Ploșoru] and in Roșia Montana [[Big and normal]] pointed out their commune origin separated due to disruptive selection. Only at Black Ploșoru the reproductive barrier was prove. This analysis demonstrated that temperature conditions, both within and between generations, influence territorial success of flies [Zamudio et al. 1995].

**344V** Screening for cryptic genetic variation in natural populations of *Drosophila melanogaster* *Gabriella Moreno*, Nicolette Alvdanid, Alexis Long, Emily Jabourian, Lukas Prelooker, David Marcey California Lutheran University

Cryptic genetic variation may be an important component of adaptive, deleterious, or neutral variation that is contingent upon environmental or genetic circumstances to be expressed. We are conducting screens for inducible, phenotypic variation in natural populations of *Drosophila melanogaster*. The screen is based on a model for the production of head defects by the extra eye mutation (ee), which is incompletely penetrant, variably expressed, and conditionally dominant. The proposed epigenetic model predicts that new mutant phenotypes may be uncovered in crosses between ee lines and flies from natural populations that harbor P-elements at various genomic locations. Such novel mutants are predicted to be incompletely penetrant and variably expressed, caused by epigenetic silencing of genes residing at genomic
positions near P transposable elements. In an initial study described here, we conducted a screen for such mutations by scoring flies in lines established from crosses of 40 separate, wild-derived lines from various geographic coordinates to ee. In several such lines, new mutations were recovered that exhibit epigenetic properties. The induction of these mutant phenotypes by crosses to ee was repeatable and not observed in control lines, suggesting that they are not induced de novo, but rather represent cryptic phenotypes induced by genetic elements in the ee line. Cby (crybaby) and Bby (Beady), both affecting ventral eye development, were observed in lines derived from Brazil and Australia, respectively. Tby (tumorous baby), obtained independently in a second line derived from Brazil, yields melanotic tumours in ~3-5% of 3rd instar larvae. Our results to date suggest that variation in natural populations may include cryptic, epigenetic sources, which are revealed under particular genetic contingencies. Screens of multiple, wild strains for additional examples of cryptic phenotypes are ongoing, as are genetic and molecular characterizations of mutant phenotypes so far recovered.

345V  Genotype-dependent effects of human disturbance on organismal fitness  Heidi Johnson, Nicole Riddle  University of Alabama at Birmingham

Human disturbance causes significant changes to home range and movement distance in wild populations. These changes and other behavior modifications often negatively impact body condition and/or fitness. While disturbance in wild populations is well-documented, there is a lack of systematic investigations defining the impact of the disturbance on energetic tradeoff and body composition. To address this need, we will utilize tools developed to study exercise in Drosophila to mimic disturbance. The Treadwheel induces movement in Drosophila through rotation of the enclosures. To sample genetic variation, we are using strains from the Drosophila Genetic Reference Panel (DGRP). The DGRP is a set of inbred strains derived from wildtype genotypes that models naturally occurring population variation. The DGRP strains differ in baseline activity approximately 525-fold for males and 110-fold for females. They also differ in their weight (1.7-fold), and quantitative magnetic resonance analysis revealed significant variation in body composition among the lines. Differences in weight, activity, and body composition is sex- and genotype-dependent. We use this diverse collection of Drosophila strains to probe the consequences of disturbance on movement patterns, body condition, and organismal fitness. We compare the effects of short-term (5 days of daily treatment on the Treadwheel) and long-term (15 days of daily treatment on the Treadwheel) disturbance regimes. Specifically, we investigate how disturbance alters longevity, offspring count, and physical condition and quantify the impact on organismal fitness, in experiments that are in progress. We expect that the impact of long-term disturbance to be more severe than the impact of short-term disturbance. We also expect to find variation in the responses to the disturbance for all outcome measures and that the outcomes are sex- and genotype-dependent.

346V  Evolutionary conservation and divergence of 3D genome organization in Drosophila  Nicole Torosin, Weihuan Cao, Christopher Ellison  Rutgers University

Topologically associating domains (TADs) are 3D organizational units of chromatin that are believed to regulate gene expression by constraining enhancer/promoter interactions. Several early studies found that TADs are highly conserved in both vertebrates and in Drosophila. However, more recent research suggests that TADs may diverge rapidly and that their reorganization is not associated with widespread divergence in gene expression, in contradiction to their supposed role in constraining enhancer/promoter contacts. In this study, we use a comparative genomics approach to estimate the rate of TAD evolution and determine whether the evolutionary conservation of TAD structures is associated with conservation of gene expression. We generated Hi-C chromosome conformation capture data for eleven Drosophila species from the melanogaster group, diverging between 4-32 million years ago. We used a phylogenetic approach to estimate the rate of TAD evolution and found that TADs evolve roughly 10-20-fold faster than other genomic features in Drosophila, such as gene duplicates and chromosomal rearrangements. Next, we found that highly conserved TADs are enriched for the BLACK and BLUE chromatin states which contain Polycomb-repressed and developmentally-regulated genes. These TADs evolve at a significantly slower rate compared to other TADs and show significantly higher constraint in gene expression levels. On the other hand, TADs that are enriched for the YELLOW chromatin state, which contains broadly-expressed, transcriptionally-active genes, evolve faster compared to all TAD domains and show less gene expression constraint. After controlling for chromatin state, we do not find a significant relationship between TAD conservation and interspecies variation in gene expression levels. These results suggest that, in general, most TADs evolve rapidly and their divergence has little effect on gene expression. However, the higher levels of evolutionary conservation and gene expression constraints in TADs enriched for developmentally-regulated chromatin suggests that these TAD subtypes may be more important for regulating gene expression, likely due to the larger number of long-distance enhancer promoter contacts associated with developmental genes.

347V  The interaction between male courtship plasticity and female mate choice in Drosophila melanogaster  Samuel Marston1, Dean Castillo1,2  1) University of Utah, SLC, UT; 2) University of Nebraska at Omaha, Omaha, NE

Sexual selection drives rapid speciation by creating divergence in sexual signals and preferences. Geographically isolated populations of D. melanogaster exhibit strong asymmetric behavioral isolation and divergence in male courtship
These lineages are often classified into Z-type lineages from Southern Africa and M-type lineages that occur outside of Africa. Z-type females almost exclusively choose Z-type males over M-type males in mate choice assays. Both Z-type and M-type males adjust their courtship behavior depending on the genotype of the female they are courting. This behavioral plasticity could potentially optimize male mating success when encountering diverse females. However, the mechanism responsible for triggering a plastic behavioral response is unknown. Males might respond to female behavioral feedback or female chemical cues. To determine which female trait facilitates male courtship plasticity we quantified courtship behaviors, focusing on the time spent singing, for males interacting with females wherein we have genetically manipulated their cuticular hydrocarbon (CHC) profile. Using a Z-type male strain that exhibits strong plasticity, we quantified the time spent singing when interacting with an M-type female and a Z-type female to establish baseline plasticity. We then tested these males with the Z-type females in which the coding sequence of the gene desat2 was replaced using CRISPR, creating a null mutant. The gene desat2 contributes to the production of the CHC 5,9-heptacosadiene (5,9-HD) a compound that is characteristic of Z-type females. This CHC is an isomer of the CHC 7,11 HD which is the most abundant CHC in M-type (non-African) females. Our Z-desat2 null strain has a CHC profile resembling an M-type female but in choice tests behaves like a Z-type female. Thus we believe this genotype would provide an M-type chemical cue to a male but Z-type like behavioral feedback. We hypothesized that if males use these heptacosadiene isomers to adjust their courtship we would expect to see male courtship of Z-desat2 null females resemble courtship of M-type females. This experiment will determine which cue triggers male courtship plasticity and how this might contribute to the evolution of reproductive isolation, furthering our understanding of the co-evolution of male behavior and female preference of closely related populations.

**348V** A locus affecting pigmentation evolution and male mating success between two sibling species in *Drosophila* Jean David, Erina Ferreira, Laure Jabaud, David Ogereau, Héloïse Bastide, Amir Yassin Laboratoire Évolution, Génomes, Comportement et Ecologie, CNRS, IRD, Université Paris-Saclay, Gif-sur-Yvette, France

New species originate when diverging populations become reproductively isolated and morphologically distinct. In theory, genetic correlation between the two phenotypes through pleiotropy or partial linkage significantly accelerates speciation. Here, we investigate the genetic correlation between mate choice and pigmentation difference in a pair of sibling species, *Drosophila yakuba* and *D. santomea*. We introgressed the “light” phenotype of *D. santomea* into the “dark” background of *D. yakuba* through a series of backcrosses. Two introgressed “light”*yakuba* strains that slightly differed in the degree of pigmentation were obtained. Genome sequencing revealed that the darker of the two strains had two introgressed X-linked loci, each centered on a melanin-synthesis gene, namely *yellow* and *tan*. The lighter introgressed strain had, in addition to the two X-linked loci, a third introgressed locus on autosomal arm 3L. The third locus centered on the transcription factor *Grunge*, which regulates the expression of the melanin-synthesis genes *ebony* and *tan* in *D. melanogaster*. We then conducted reproductive isolation assays between the two introgressed strains and strains from the parental species, measuring male and female mating latency and choice. We found no evidence for reproductive isolation in the darker introgressed strain, in contrast to the lighter strain, which exhibited decreased male mating success most likely due to increased latency. Those findings indicate that reproductive isolation and pigmentation difference between *D. yakuba* and *D. santomea* are partly linked through a single autosomal locus. Future dissection of this locus will elucidate the molecular underpinnings of such partial correlation in this primary model of speciation research.

**349V** Evidence of horizontal transmission of *Wolbachia* in *Drosophila sturtevanti* and *Drosophila lehrmanae* (saltans group) Bruna Roman1, Carolina Prediger1,2, Amir Yassin2, Lilian Madi-Ravazzi1 1) São Paulo State University (UNESP), Institute of Biosciences, Humanities and Exact Sciences, São José do Rio Preto, São Paulo, Brazil; 2) Laboratoire Evolution, Génomes, Comportement, Ecologie (EGCE), CNRS, IRD, Université Paris-Saclay, Gif-sur-Yvette, France

*Wolbachia* is an endosymbiotic genus of Alphaproteobacteria that infects a wide range of arthropods and nematodes. This endosymbiont is typically transmitted vertically, but in some cases, horizontal transmission has been reported. The horizontal transmission can occur in different ways in nature, such as parasitism, cannibalism and predation of infected individuals, hybrid introgression and shared ecological niches. Some species of *Drosophila saltans* group are infected by this endosymbiont, such as *Drosophila sturtevanti* (wStv), *D. prosaltans* (wPro) and *D. septentriosaltans* (wStv). Here, we analyze the horizontal acquisition of *Wolbachia* in the host species *D. sturtevanti* and *D. lehrmanae*. We inferred a phylogenetic tree, in MrBayes, with 129 *Wolbachia* genomes (7 sequenced by us and 122 from database) and compared it with the phylogeny of the hosts. The single copy orthologs genes (17) were identified in OrthoFinder and used in the analysis. In general, the phylogeny generated showed that most strains were grouped into their defined supergroups according to the literature. However, six strains (namely, wAlb, wAgra, wNik, wAdent, wStv and wStv-like), belonging to supergroup A, grouped together and distant from the other members of this supergroup. The formation of this last clade is interesting because the phylogeny of these strains demonstrates disagreement with the phylogeny of their hosts. Thus, this result suggests wStv (host *D. sturtevanti*) and wStv-like (host *D. lehrmanae*) were acquired by horizontal transmission, since wPro and wSpt strains also infective of saltans group species were positioned phylogenetically
we find that there are increases in embryo length associated with maternal changes in metabolism and ovariole expression, and normalized cuticles after ten generations. Using a multi-omics approach across the evolved populations, followed by experimental evolution. We find that the evolved populations have increased fitness, canalization of gene evolution across populations with two extra copies of Bicoid, we performed unbiased genome-wide mutagenesis, causes a posterior shift of the networks' regulatory outputs and a decrease in fitness. To directly monitor network patterning in early embryos. This network can be synthetically perturbed by increasing the dosage of regulatory network evolution at the system level, and such knowledge learned from experimental evolution can help degree laboratory evolution can be used to explore these processes. To address these questions, we have used Lautaro Gandara, Kerstin Richter, Justin Crocker EMBL

350V  Sexual Selection is not a Driver of Female Sperm Storage Organ Length in Drosophila Cameron Himes, Tiffini Smith, Mollie Manier The George Washington University

In sexual selection, male trait evolution is often driven by female preference in a way that can generate exaggerated male phenotypes via a Fisherian runaway process. In this way, character transitions in the male trait are expected to follow female preference transitions. However, factors driving the evolution of female preference are less understood. Sperm in Drosophila are extremely long and are stored within the long, coiled female sperm storage organ, the seminal receptacle, or SR. When females mate with multiple males, long SRs favor long sperm, and short SRs favor short sperm. Thus, sperm length and SR length are a post-copulatory male trait-female preference system. Consistent with this, sperm length and SR length are coevolving across the Drosophila lineage, driven by a genetic correlation between the two traits. However, it is unknown what drives SR evolution. To test the hypothesis that SR length evolution is correlated with post-copulatory sexual selection, we obtained remating rates, SR length, sperm length, body size, and fecundity in 81 Drosophila species through direct experimentation and conducted a comprehensive literature search. We performed a comparative phylogenetic trait evolution test to determine if there is correlated evolution between remating rate and SR length in a phylogenetic context. There was no significant correlation between remating rate and SR length, suggesting that sexual selection is not driving the evolution of SR length on a macroevolutionary scale. The genetic correlation between SR and sperm lengths and their respective competitive phenotypes may be enough to drive sperm-SR coevolution without selection specifically acting on SR length.

351V  Intermolecular interactions drive protein adaptive and co-adaptive evolution at both species and population levels Junhui Peng, Nicolas Svetec, Li Zhao Rockefeller University

Proteins are the building blocks for almost all the functions in cells. Understanding the molecular evolution of proteins and the forces that shape protein evolution is essential in understanding the basis of function and evolution. Previous studies have shown that adaptation frequently occurs at the protein surface, such as in genes involved in host-pathogen interactions. However, it remains unclear whether adaptive sites are distributed randomly or at regions associated with particular structural or functional characteristics across the genome, since many proteins lack structural or functional annotations. Here, we seek to tackle this question by combining large-scale bioinformatic prediction, structural analysis, phylogenetic inference, and population genomic analysis of Drosophila protein-coding genes. We found that protein sequence adaptation is more relevant to function-related rather than structure-related properties. Interestingly, intermolecular interactions contribute significantly to protein adaptation. We further showed that intermolecular interactions, such as physical interactions may play a role in the co-adaptation of fast-adaptive proteins. We found that strongly differentiated amino acids across geographic regions in protein-coding genes are mostly adaptive, which may contribute to the long-term adaptive evolution. This strongly indicates that a number of adaptive sites tend to be repeatedly mutated and selected in evolution, in the past, present, and maybe future. Our results highlight the important roles of intermolecular interactions and co-adaptation in the adaptive evolution of proteins both at the species and population levels.

352V  Synthetic evolution of a Drosophila developmental network predicts trends in wild populations Xueying Li, Lautaro Gandara, Kerstin Richter, Justin Crocker EMBL

It remains unknown how developmental regulatory networks evolve, the predictability of their evolution, or to what degree laboratory evolution can be used to explore these processes. To address these questions, we have used experimental evolution to examine the bicoid (bcd) network in Drosophila, which is essential for anterior-posterior patterning in early embryos. This network can be synthetically perturbed by increasing the dosage of bicoi, which causes a posterior shift of the networks’ regulatory outputs and a decrease in fitness. To directly monitor network evolution across populations with two extra copies of Bicoid, we performed unbiased genome-wide mutagenesis, followed by experimental evolution. We find that the evolved populations have increased fitness, canalization of gene expression, and normalized cuticles after ten generations. Using a multi-omics approach across the evolved populations, we find that there are increases in embryo length associated with maternal changes in metabolism and ovariole development. Consistent with our observation in laboratory evolution, we find that a wild population with larger embryos similarly rescues progeny with increased bcd expression. Together, our results necessitate a broader view of regulatory network evolution at the system level, and such knowledge learned from experimental evolution can help predict evolutionary trends in nature.
353V  *fushi tarazu* and *fushi tarazu factor 1*, novel re-wiring in the *Tribolium castaneum* pair-rule gene network  
Ximena Gutierrez Ramos, Patricia L. Graham, Leslie Pick University of Maryland, College Park

Segmentation is a fixed feature of the insect body plan, yet surprisingly, the genetic network controlling the formation of segments has changed over the course of insect evolution. In *Drosophila*, pair-rule genes are responsible for allocating groups of cells to specific body regions that develop into separate, morphologically distinct body segments. Transcription factor (TF)-encoding genes regulate sets of downstream genes that include those encoding other TFs, signaling proteins, and products directly involved in morphogenesis. The evolutionary variation seen for pair-rule genes contrast with the conservation of expression of downstream target genes. How does the regulatory network re-wire to maintain downstream gene expression while upstream regulators are lost, gained, or changed in function? We are exploring this question by examining the expression, function, and interactions of the pair-rule gene *fushi tarazu* (ftz) in the flour beetle, *Tribolium castaneum* (Tc). Tc-FTZ contains an LXXLL motif that is necessary for the functional interaction of FTZ and its obligatory partner FTZ-F1 in *Drosophila*. Despite this, a large genomic deletion (Stuart et al., 1991) and RNAi experiments (Choe et al., 2006 and data not shown) indicate that Tc-ftz plays no role in segment formation in *Tribolium*. In contrast, FTZ-F1 is expressed in pair-rule stripes in *Tribolium* and RNAi knockdown resulted in pair-rule defects (Heffer et al., 2013). To definitively determine whether Tc-ftz plays a role in segmentation, we are using CRISPR/Cas9 to generate a genomic deletion of this gene. Further, to understand how FTZ-F1 regulates pair-rule patterning in *Tribolium* without the input of FTZ, we are examining cis-regulatory elements of the conserved target gene *enlarged* (en). We are testing approaches to using CRISPR-based and piggybac-based enhancer reporter genes to identify regulatory elements and assess their dependency on Tc-FTZ-F1, using dsRNA to knockdown Tc-ftz-f1 expression. Other TFs binding to identified cis-regulatory elements will be identified and characterized. Simultaneously, we have generated a FLAG-tagged version of Tc-FTZ-F1 to identify other genomic targets, determine whether or not Tc-FTZ plays a role in their regulation and to identify the DNA binding partners using co-immunoprecipitation. This will allow us to understand how a rearrangement of upstream regulators occurs to maintain a conserved downstream gene expression pattern, and provide information on the transcriptional regulation of the downstream genes within the *Tribolium* pair-rule gene network.

354V  Probing evolution by *Hox* locus replacement  
ANKUSH AURADKAR1,2, Emily A. Bulger3, Sushil Devkota3, William McGinnis1, Ethan Bier1,2 1) Section of Cell and Developmental Biology, University of California San Diego, La Jolla, CA 92093, USA; 2) Tata Institute for Genetics and Society-UCSD, La Jolla, CA 92093-0335, USA; 3) Developmental and Stem Cell Biology Graduate Program, University of California San Francisco, and Gladstone Institutes, San Francisco, CA, USA

*Hox* genes determine positional codes along the head-to-tail axis. Here, we replaced the entire *Drosophila melanogaster* proboscipedia (*pb*) *Hox* locus, which controls the development of the proboscis and maxillary palps, with that from *Drosophila mimica*, a related species with highly modified mouthparts. The *D. mimica* replacement rescues most aspects of adult proboscis morphology; however, the shape and orientation of maxillary palps were modified, resembling *D. mimica* and closely related species. Expressing the *D. mimica* *Pb* protein in the *D. melanogaster* pattern fully rescued *D. melanogaster* morphology. In contrast, expression of the *pb* locus directed by *D. mimica* *pb* cis-regulatory sequences was reduced from that of *D. melanogaster* *pb* in cells that produce altered maxillary structures, potentially altering the balance between *pb* and a competing *Hox* gene *Deformed* (*Dfd*). This hypothesis is consistent with the observation that over-expression of *Dfd* in *D. melanogaster* results in altered maxillary palp orientation similar to that in *D. mimica* replacement. These findings suggest that *pb* regulatory sequences evolved in related species to alter mouthpart morphology.


355V  Genetic architecture of male-female coevolution in *Drosophila melanogaster*  
Cameron Himes, Galvin Jake, Mollie Manier George Washington University

In sexual selection, male trait evolution is often driven by female preference in a way that can generate exaggerated male phenotypes via a Fisherian runaway process. This male-female co-evolution requires a genetic correlation between the male trait and female preference, but the genetic architecture behind this correlation is typically unknown. In Drosophila, sperm are extremely long and are stored within the long, coiled female sperm storage organ, the seminal receptacle, or SR. SR length is a mechanism of cryptic female choice for sperm length, such that when females mate with multiple males, long SRs favor long sperm, and short SRs favor short sperm. Thus, sperm length and SR length are a post-copulatory male trait-female preference system. Consistent with this, sperm length and SR length are coevolving across the Drosophila lineage, driven by a genetic correlation between the two traits. Here, we have used the Drosophila Synthetic Population Resource (DSPR) to characterize the genetic architecture underlying sperm-SR coevolution and to identify candidate genes underlying each trait. We found that sperm and SR length are each associated with a single QTL peak that are closely linked on chromosome 3R, leading us to conclude that the genetic mechanism of male-female
co-evolution in this case is linkage rather than pleiotropy. Within the sperm QTL is the gene scotti (soti), which encodes a testis-specific protein that localizes to late-stage spermatids and regulates the onset of individualization. We examined soti mutants and found that spermatid cysts are 350 microns shorter than wild type controls. Within the SR QTL are two developmental regulators of epithelial cell migration, heartless (htl) and stripe (sr). Both of these genes are expressed in the female reproductive tract and enriched in SR tissue, which is epithelial in origin. Ongoing experiments are further characterizing the roles of htl and sr in SR development.

356V Genetic basis of variation in high sugar-induced diabetes-associated traits and development delay in Drosophila Xuan Zhuang1, Fabio Morgante2, Abaranjitha Muniyasamy1, Michael Ludwig3, Soo Young Park3, Yang Li3, Matthew Stephens3, Graeme Bell3, Martin Kreitman1 1) Department of Biological Sciences, University of Arkansas, Fayetteville, AR; 2) Clemson Center for Human Genetics, Clemson University, Clemson, SC; 3) Department of Ecology and Evolution, University of Chicago, Chicago, IL

Drosophila is a well-established model for investigating complex traits and many human diseases. It provides powerful tools for dissecting the contributions of both genes and environment on development and metabolism. We developed a Drosophila model to study high sugar diet (HSD) induced Type 2 diabetes (T2D) associated traits in large populations with different genetic backgrounds. We examined HSD induced phenotypes in larvae and in adults across a subset of Drosophila Genetic Reference Panel (DGRP). Flies on HSD display an increase in whole body glucose and glycogen level, a decrease in developmental rate, survivorship, body weight, and longevity, compared with flies under low sugar diet (LSD). The examined DGRP lines display a continuous and wide range of these phenotypes and large broad-sense heritability, suggesting great potential for quantitative trait loci (QTL) mapping. In the meanwhile, we developed a unique experimental system for genetic mapping named Drosophila Recombinant Populations (DRPs), which are consisted of 16 outbred advanced intercross populations (AIPs), each founded with 8 inbred DGRP lines. The DRPs provide about 70,000 genotypically distinct flies that allow us to apply the HSD induced T2D model to each individual to investigate the genetic architecture of these complex traits. We used one of the HSD induced traits, namely developmental delay, to perform a bulk segregant mapping analysis of extreme phenotypes. We developed a computational pipeline with a Hidden Markov model (HMM) to impute the whole genome of each fly based on their low-coverage sequenced genomes and the available high-coverage founder genomes. Genome-wide association studies (GWAS) identified 76, 2009, and 373 polymorphisms at p<10^{-5} for the analysis of HSD, LSD, and G x E (gene by diet) respectively. Gene ontology (GO) analysis indicates predominant enrichment for genes involved in some hormones and ketones cellular metabolic processes among HSD candidates, and reproduction and mediator complex among LSD candidates. RNAi validation studies of the top candidates (Cyp9b2 and CG15088 from GxE analysis, wun and Ten-m from HSD analysis) indicated their moderate but still significant effects in increasing the development rate on HSD. This study provides a broad survey of diabetes associated traits in HSD induced T2D flies, and further dissects the genetic architecture of development time on diets of different sugar concentrations, and interaction of gene and diet.

357V Effective label of XL/XR and Neo-X chromosomes of Drosophila miranda using oligopaints probes Henry Bonilla1, Isabela Pimentel de Almeida1, Mara Lisboa Santana Pinheiros1, Maria Vibranovski1,2 1) Bioscience Institute, University of São Paulo; 2) New college for Interdisciplinary Arts and Sciences, Arizona State University

Experiments of fluorescent in-situ hybridization (FISH) are a valuable resource for investigating chromosome evolution, chromosome behavior in individual cells and, ultimately, for assessing the quality of a sequenced genome. Drosophila miranda is a model species to study the evolution of sex chromosomes due to their X chromosomes of different ages. The outcome of independent autosome-sex chromosome fusions produced the so-called XR and Neo-X that emerged about 15 MYA and 1.5 MYA ago, respectively. Oligopaint represented a new generation of probes (Beliveau et al 2012) that are custom-synthesized oligonucleotides, short (26–46 bases), versatile and very specific for targeting single-copy chromosome regions that can be extended to any organism whose genome has been sequenced. Here, we developed oligopaints to label the entire set of X (XL/XR and Neo-X) and the fourth autosomal chromosomes. We used OligoMiner pipeline (Beliveau et al., 2018) and the latest D. miranda genome sequence (Manhajan et al., 2018). The specificity of oligopaints was assessed by FISH experiments on mitotic chromosomes. Shortly, we designed oligopaints under the stringent condition defined in OligoMiner in addition to Kmer and secondary structure filters to guarantee specificity. A Python script was developed to diminish the number of oligos but maintain an appropriate distribution along entire target regions. Finally, flanking primer regions were added to produce oligos from a custom complex oligo-pool library.

We obtained 25553 oligos covering the entire Neo-X chromosome with a density of 1.01 oligos/Kb. The XL/XR chromosome has been divided in four regions according to their different evolutionary history and the genome assembly. By separating the XL arm in pericentromeric and distal regions (XL-1 and XL-2), we obtained 11180 oligos (0.79 oligos/kb) and 27282 (1.08 oligos/kb), respectively. The XR arm was split in XR-A and XR-D. XR-A comes from the ancient X chromosome and it’s located in the pericentromeric region whereas XR-D in the distal portion of the XR arm. We obtained 12717 oligos (1.02) and 25382 oligos (0.99) in the XR-A and XR-D respectively. Only 10336 oligos (0.32 oligos/kb) were designed for the fourth chromosome.
The follicle stem cell (FSC) niche in the adult *Drosophila* ovary balances the actions of stem cell self-renewal and output through the guidance of graded extracellular signals. Anterior sources (cap cells, escort cells [ECs]) produce Hh and Wnt pathway ligands, and posterior sources (polar follicle cells [FCs]) produce JAK-STAT pathway ligands. These opposing gradients intersect over the FSC domain, and regulate FSC division and differentiation into quiescent ECs (anterior) or proliferative FCs (posterior) across FSC positions. We have previously shown through combined lineage and morphological studies that adult ECs, FSCs, and FCs derive from intermingled cell (IC) precursors present at the start of pupation, and acquire their adult function based on their final anteroposterior (AP) location, rather than a series of cell fate decisions. Our studies also revealed how the first FCs and egg chamber are formed. Precursors accumulate posterior to germline and ICs of the developing gerarium. These Extra-Germarial Crown (EGC) and Basal Stalk (BS) cells are then invaded by the most posterior germline cyst without any attached ICs, and surround this cyst in an epithelial monolayer. Here we investigate how signaling pathways influence somatic cell AP locations and adult fates during pupal development. Wnt signaling initiates early from the anterior. MARCM lineage studies showed that cells with elevated Wnt pathway activity are biased towards more anterior locations, while reduced Wnt signaling favored more posterior locations. Detectable JAK-STAT signal was seen only after budding of the first pupal egg chamber, when polar FCs are specified. Loss of JAK-STAT signaling favored more anterior locations. Elevated JAK-STAT signaling increased precursor output, suggesting faster division. Hh signaling has previously been demonstrated to begin before pupal development and to be important for generating adult somatic cells of the ovary. Preliminary results suggest that Hh signaling during pupal stages is required for ovarian somatic cell survival and proliferation, with effects on cell location still to be determined. We are particularly interested in exploring which signals affect formation of the first FCs because that process occurs in a different morphological setting to the conversion of adult FSCs to FCs.

**359A  Diapause extends female germline stem cell longevity in *Drosophila***

*Sreesankar Easwaran, Denise Montell  
University of California - Santa Barbara*

A consequence of aging in many species including humans is reduced female fertility. Intriguingly, some species can preserve fertility longer under specific environmental conditions. For example, at low temperature and short day-length, *Drosophila melanogaster* can enter a dormant state called adult reproductive diapause. During this cold-induced quiescence, as in other stressful conditions, ovarian development arrests at the yolk uptake checkpoint. However, mechanisms underlying fertility preservation and post-diapause recovery are largely unknown. Here, we report the effects of cold-induced arrest and recovery on *Drosophila* ovarian development. We found that diapause affects all stages including germline stem cells (GSCs) and is distinct from other stress responses in causing more complete arrest yet preserving the potential for recovery. Further, GSCs incur DNA damage, activate p53 and the Chk2 checkpoint, and divide less during dormancy. Niche signaling is reduced yet germline precursor cells do not differentiate. Thus, GSCs adopt an atypical, suspended state connected to cystoblast daughters. Post-diapause recovery of niche signaling, resumption of GSC division, and dedifferentiation contribute to restoring GSCs. Mimicking one feature of the quiescent state, reduced juvenile hormone production, enhanced GSC longevity in non-diapausing flies. Thus, studies of adult reproductive diapause in *Drosophila* can provide approaches to GSC longevity enhancement.

**360B  The impact of cell cycle and DNA damage response on germline stem cell survival in the *Drosophila* testis**

*Jasmine Grey, Salman Hasan, Erika Matunis  
Johns Hopkins University School of Medicine, Baltimore, MD*

Stem cells are integral to the development and maintenance of a tissue. Adult stem cells have the unique ability to self-renew and produce differentiated daughter cells and have been shown to survive radiation-induced DNA damage in many tissues. However, the exact pathways these cells use to repair DNA damage in an intact tissue are not very well understood. Here we use the *Drosophila* testis stem cell niche to understand the processes of DNA damage response in an intact niche. Germline stem cells (GSCs) give rise to spermatogonia in the *Drosophila* testis and have been shown to resist high (75 Gy) irradiation (IR), while their differentiating progeny, spermatogonia die. The radio-resistance of the GSCs suggests that they efficiently and successfully use the DNA double-strand break (DSB) repair pathways, non-homologous end joining (NHEJ) and homologous recombination (HR). Our goal is to understand the specific mechanisms GSCs use to resist IR induced damage and maintain tissue homeostasis. Preliminary data suggest that the response of GSCs to IR in flies null for NHEJ is phenotypically indistinguishable from wild type, but flies null for HR are more sensitive to IR. We hypothesize
HR is required for damage-induced regeneration and that the ability of the GSCs to survive irradiation depends on its cell cycle phase at the time of irradiation, which dictates the DSB repair pathway used. To test this, we are using *Drosophila*-specific fluorescent ubiquitination-based cell cycle indicator (fly-FUCCI) system to visibly mark the stages of the cell cycle in live and fixed imaging, providing an in-depth analysis into the survival mechanisms carried out by IR-damaged GSCs. A deeper understanding of stem cell resistance to IR will add greatly to the *Drosophila* testis field and may have implications in other radio-resistant stem cells, such as cancer stem cells.

### 361C  Tnpo-SR maintains ovarian cyst connectivity and is required for GSC fusome dynamics morphogenesis in *Drosophila* ovarian germline stem cells

Anna Williams, Elizabeth Ables East Carolina University

The maintenance of species is dependent on parental genetic information being passed onto the next generation. Germ cells are crucial for this transfer and therefore drive evolution. Evolutionarily conserved across phyla, germ cells in many systems are supported by a stem cell population (GSCs). In *Drosophila*, these stem cells divide via incomplete cytokinesis to form interconnected daughter cells that will mitotically divide and differentiate to form oocytes and nurse cells, which supply the oocyte with maternal factors necessary for embryogenesis. Connection of daughter cells is facilitated by the fusome, an endoplasmic reticulum-like structure with cytoskeletal and membranous components. The fusome is remodeled upon division and will become increasingly branched as the cysts continue to divide. Tnpo-SR, an ecdysone-responsive gene, was identified in a genetic screen for its putative role in stem cell self-renewal. Tnpo-SR shares amino acid similarity to mammalian β-importin proteins; however, its specific roles in gamete production has not been characterized. Using genetic mosaic mutants alongside germline-specific RNAi, we found that Tnpo-SR is necessary in GSCs for their self-renewal. Depletion of Tnpo-SR in dividing germ cells generates egg chambers with cells varying from the typical 16 suggesting abnormal cyst development and cyst fragmentation. Here, we investigated the cytoskeletal dynamics of Tnpo-SR mutant GSCs and cytoeblasts in order to further characterize the role of Tnpo-SR in GSC self-renewal. We find that GSC fusome density is decreased when Tnpo-SR is depleted, suggesting its role in GSC fusome morphogenesis. The alteration of cyst formation also impacts oocyte selection. This research identifies a novel role of the karyopherin Tnpo-SR on not only GSC formation and maintenance but also the formation of a healthy egg needed to drive the maintenance of species.

### 362A  Effects of nuclear lamina aging on oogenesis

William Zaremba, Pamela Geyer University of Iowa

Tissue homeostasis depends on replacement of injured and defective cells by proliferation of resident adult stem cells. Homeostasis declines with age, largely due to decreased stem cell function and/or number. The *Drosophila* ovary provides an outstanding model for investigating age effects on tissue homeostasis, as oogenesis depends upon a well-defined germline stem cell (GSC) population that sustains oocyte production for many weeks. Age-dependent declines of oocyte production are associated with decreased mitosis, not GSC loss. Strikingly, we discovered that the asymmetric mitoses of GSCs are specialized, wherein the nuclear envelope and nuclear lamina remain throughout mitosis. During these specialized mitoses, spindles are nucleated from centrosomes embedded in the retained NL. The mode of GSC mitosis raises questions about how the mitotic NL contributes to homeostasis. To this end, we have studied age effects on accumulation of mitotic NL components, finding that lamin-B and emerin levels decline with age. These observations support predictions that deterioration of mitotic NL integrity reduces mitotic division and oocyte production. Further support for this prediction comes from our studies of the role of Checkpoint kinase 2, a transducer kinase that becomes activated upon NL defects and elevated DNA damage in germ cells. Notably, loss of Chk2 prolongs oocyte production, delaying loss of egg laying by 15 days, suggesting that activation of Chk2 contributes to reduced fecundity of older females. Taken together, our studies provide new insights into age-dependent contributions of the NL to homeostasis of the ovary.

### 363B  Function of Bazooka in dedifferentiation of the male germline stem cells

Muhammed Burak Bener, Mayu Inaba University of Connecticut Health, Farmington, CT

*Drosophila* male germline stem cells (GSCs) divide asymmetrically to generate a self-renewing stem cell and a differentiating cell. The outcome of asymmetric division is compromised when cells that once initiated differentiation program revert back to stem cell state (i.e., dedifferentiation). Dedifferentiation has been proven to be an essential mechanism for the maintenance of the GSC pool. However, compromised asymmetric outcome may result in the clonal expansion/depletion of stem cells, leading to various pathological conditions. Therefore, the process of dedifferentiation must be carefully regulated. In this study, we characterized the process and outcome of dedifferentiation to understand the underlying mechanisms. By experimentally inducing a GSC-loss and recovery cycle in the testis, we found that the dedifferentiated GSCs reestablish the patch-like structure of a polarity protein, Bazooka (Baz, *Drosophila* homolog of mammalian Par-3), at the hub-GSC interface quickly within 1 to 3 days after induced GSC loss. Moreover, we found that differentiating spermatogonia start expressing Baz prior to migration to the niche, which suggests a role of Baz in the regulation of the dedifferentiation process. Baz has been shown to be required for stem cell-specific polarity checkpoint, centrosome orientation checkpoint (COC). Consistent with the quick recovery of Baz-patch, we observed the presence of intact centrosome orientation checkpoint, correct spindle orientation, and intact mitotic rates in dedifferentiated
GSCs. Moreover, our live observation demonstrates that dedifferentiated GSC immediately divides after positioned back to the niche. Taken together, our results demonstrate that dedifferentiated GSCs quickly regain the ability to undergo asymmetric cell division and Baz plays an essential role in this process.

**364C Investigating the Regulation of Germline Stem Cell Cytokinesis by Somatic Stem Cells** Carlos Billini, Kari Lenhart Drexel University

Most adult tissues have multiple specialized stem cell populations residing within a single microenvironment, or niche, that repopulate and regenerate the tissue over time. While much is known about how the niche controls a single stem cell line, not much is known regarding the interaction and coordination between the different stem cell types contained that repopulate and regenerate the tissue over time. While much is known about how the niche controls a single stem cell, not much is known regarding the interaction and coordination between the different stem cell types contained by a single niche. The *Drosophila* testis niche contains two populations of stem cells: Cyst Stem Cells (CySC) and Germline Stem Cells (GSC) that must be precisely coordinated for proper tissue function. It is known that two CySC daughter cells envelope a single GSC daughter cell, insulating the GSC daughter from niche signals. Our lab has identified that somatic control of GSC cytokinesis helps coordinate daughter cell release from the niche, resulting in this 2:1 ratio. Further, we determined that the 2:1 ratio is required for complete and proper GSC daughter cytokinesis. Inhibition of CySC cytokinesis leads to failure of abscission, or membrane severing, between GSC-daughter pairs and results in failure of release of a differentiating germ cell from the niche. Therefore, we hypothesize that a signal from CySCs is necessary to trigger proper abscission of the GSCs. We have additionally found that Wnt pathway activation within CySCs is required for successful completion of GSC abscission. Surprisingly, we find that somatic Wnt pathway activation does not directly regulate GSC abscission. Instead, we have discovered that Wnt activity in CySCs is necessary to dampen GSC cycling rate. Inhibition of the Wnt pathway in somatic stem cells results in a significant increase in GSC cycling rate and a consequent failure of these cells to successfully complete cytokinesis prior to mitotic entry. Further understanding how somatic stem cells regulate the cycling rate of adjacent GSCs will elucidate the collaborative mechanisms in place within the niche to ensure tissue homeostasis.

**365A Investigating re-initiation of stem cell cytokinesis during tumor proliferation** Beth Kern, Kari Lenhart Drexel University

Adult stem cells become depleted throughout the lifetime of an organism, necessitating regenerative mechanisms to maintain the stem cell niche. Interestingly, signals for plastic regeneration mechanisms are often shared with those initiating tumorigenesis. A common feature of both plasticity and tumorigenesis is a reversion of differentiating cells to a stem-like state. The *Drosophila* male testis provides an invaluable model to study the process whereby differentiating cells transform into overproliferative tumor cells due to its distinct location and genetic tractability. Within the testis, the stem cell niche maintains both germ line stem cells (GSCs) and somatic cyst stem cells (CySCs). GSCs divide asymmetrically to produce a differentiating daughter, which must become encapsulated, or encysted, by two CySC daughters. Our lab has developed a model whereby the coordinated release of this 2:1 soma-germline grouping is achieved via a modified cytokinesis program that pauses completion of GSC cytokinesis until somatic encystment of the germline is achieved. Whether this modified cytokinesis program becomes re-engaged during tumorigenesis and to what degree tumorous germ cells reinstate stem cell specific processes has yet to be explored. By genetically inducing tumorigenesis through ectopic expression of niche signals, we have visualized actin misregulation at the intercellular bridge and subsequent tumor fractionation by single cell abscission. We now aim to directly examine if tumor progression requires re-initiation of the stem cell-specific cytokinesis program. Examination of *in vivo* tumor proliferation will provide critical insights into the cell biology underlying tumorigenesis and may be useful in elucidating novel therapeutics for treatment of carcinoma.

**366B Programmed changes of interaction of Stat92E homologous loci regulate transcription during the stem cell differentiation** Mayu Inaba, Matthew Antel UConn Health, Farmington, CT

Correlation between strength of pairing of homologous chromosomes and gene expression status has been demonstrated in genome-wide, implicating the functional impact of homolog pairing on local transcriptional activity. *Drosophila* male germline stem cells (GSCs) constantly divide asymmetrically to produce one GSC and the other daughter as a differentiating gonialblast (GB). GB then enters differentiation program in which stem cell specific genes are quickly downregulated. Here we demonstrate that a change of local pairing status of homologous Stat92E loci is required for downregulation of Stat92E gene during the differentiation. Using Oligopaint fluorescent in situ hybridization (FISH) technique, we found that the interaction between homologous loci of Stat92E is always tight in GSCs and immediately loosened in GBs. When one of the STAT92E loci was absent or relocated on other chromosome, Stat92E did not pair and failed to downregulate. Same defect was observed upon knocking down of pairing factors, suggesting that the pairing is likely required for switching transcriptional status. Moreover, Stat92E enhancer element but not cis-transcription is required for the change of pairing status, indicating that pairing change is not a consequence of transcriptional changes. GSCs are known to inherit pre-existing histones (H3 and H4), while newly synthesized histones are distributed in GBs. When this histone inheritance was compromised, the change of Stat92E pairing did not occur, suggesting that the pairing change is an intrinsically programmed process during the asymmetric stem cell division. Taken
together, we propose a possibility that the change of local pairing status may be a common process to promote rewriting gene activity status during cell-differentiation.

367C  Examining the Role of Adipokines in Regulating Oogenesis Chad Simmons, Alissa Armstrong University of South Carolina

Obesity and malnutrition can influence nutrient signaling pathways that relay nutrient status throughout the body, yet the molecular components that allow communication about dietary input between different tissues is not completely understood. *Drosophila melanogaster* ovarian function is sensitive to changes in diet and serves as an ideal model to decipher how nutrient sensing by peripheral tissues influence the ovary. Previous studies demonstrate that *Drosophila* adipose tissue uses nutrient sensing pathways, like insulin/insulin-like growth factor and Target of Rapamycin signaling to modulate ovarian germline stem cells and their progeny (Armstrong et al., 2014; Armstrong and Drummond-Barbosa, 2018). Multiple adipokines, factors secreted from the adipose tissue, are required for tissue growth during larval development. The range of adipocyte-derived factors that contribute to adipocyte-to-ovary communication is not well understood. We propose that the adipokines with roles in larval growth, eiger (egr), unpaired2 (upd2), stunted (sun), growth blocking peptides (Gbp1, Gbp2, Gbp3) and CCH-amides (ccha2) also function to promote adipose tissue nutrient sensing control of oogenesis. We find that all are expressed in the adult female fat body. Using RT-PCR, we are determining if their expression is modulated by protein-poor and obesogenic diets. To determine their functional role, we use the Gal4/UAS gene expression system for RNAi-mediated knockdown of each adipokine specifically in adult adipocytes and measure effects on oogenesis using whole mount immunofluorescence and confocal microscopy. Thus far, we have observed that adipocyte knockdown of *upd2* and *egr* leads to increased adipocyte size but does not affect germline stem cell number in the ovary. We are currently investigating the effects of adipokine knockdown on other aspects of oogenesis, including germline stem cell maintenance, germline cyst survival, progression through vitellogenesis and ovulation. Understanding how these adipokines communicate to the ovary will allow us to begin uncovering the cellular and molecular mechanisms that underlie signaling between adipose tissue and the germline stem cells.

368A  Assessing the interactions between *W. pipientis* genotype and titer on the bag of marbles partial loss of function mutant (hypomorph) in *Drosophila melanogaster* Catherine Kagemann, Charles Aquadro Cornell University

*Wolbachia pipientis* are maternally transmitted endosymbiotic bacteria commonly found in arthropods and nematodes. *W. pipientis* have complex interactions with their hosts, and many of these interactions serve to increase transmission. *W. pipientis* commonly manipulate reproduction of the host via cytoplasmic incompatibility, resulting in embryonic mortality. However, there are other known interactions between *W. pipientis* and its host. For example, *W. pipientis* rescues the bag of marbles (*bam*) partial loss of function (hypomorph) fertility phenotype in female *Drosophila melanogaster*. *Bam* is an important germline stem cell (*GSC*) gene involved in GSC renewal and cystoblast differentiation. GSCs are required for the production of egg and sperm, making the genetic interaction between *W. pipientis* and GSC genes such as *bam* of great evolutionary interest to us. While we understand that *W. pipientis* contributes to the rescue of the *bam* hypomorph phenotype, we have yet to determine the functional mechanisms that are behind this interaction. Therefore, we aim to elucidate 1) whether variation in *W. pipientis* variant genotype and titer influence the rescue of the mutant *bam* phenotype at different ages in adult females and 2) whether *W. pipientis* variants cause differential rescue of the *bam* hypomorph phenotype at the transcriptional level as the host fly ages. Results show that rescue of the mutant *bam* phenotype does in fact depend on the genotype of *W. pipientis* as the flies age and the magnitude of rescue is dependent on the age of the female fly. Relative quantification of *W. pipientis* titer via qPCR shows that titer increases in all *W. pipientis* genotypes at the peak rescue of the *bam* hypomorph phenotype. Our RNA-seq analysis revealed that *W. pipientis* infected *Drosophila* differentially express many of *bam*’s genetic and physical interactors in the *bam* hypomorph genotype. Additionally, pairwise comparisons between *W. pipientis* genotypes showed differential expression of one of *bam*’s genetic interactors, *zpg*, and several other GSC genes. RNAI will be used to determine whether any of these candidate genes identified from our RNA-seq analysis are responsible for aiding in the rescue of the *bam* hypomorph phenotype at the transcriptional level.

369B  Lineage decisions and competency in early *Drosophila melanogaster* neurogenesis Fiona Kerlin, Robert Zinzen Max Delbrück Center for Molecular Medicine in the Helmholtz Association Berlin-Mitte (BIMSB)

During embryogenesis in *Drosophila* melanogaster, a tightly controlled gene regulatory network generates a specific number of neurogenic stem cells, also called neuroblasts, in distinct positions over the course of early development. This process of neurogenesis generates a total of ~30 neuroblasts per segment in a spatially and temporally controlled manner, where neuroblasts are successively selected and delaminate from an epithelial sheet of cells along 3 dorsoventral columns and several anteroposterior rows. Once born, neuroblasts divide asymmetrically several times to self-renew and produce ganglion mother cells, neurons, and glia. This series of events is characterized by a temporally and spatially distinct gene expression, so that each neuroblast has a unique pattern of gene expression that determines its developmental trajectory. Importantly, the spatial and temporal origin of the neuroblast is thought to determine its...
ensuing lineage and cell fate.

After decades of research the neurogenic system in the early Drosophila embryo is quite well understood phenomenologically: Not only are we aware of many gene expression combinations that are distinct for individual neuroblasts, but fluorescent labeling experiments of individual neuroblasts have yielded maps of their developmental trajectories. However, our understanding of the molecular mechanisms that restrict, drive and distinguish neuroblast fate decisions is extremely limited.

New studies in my lab using single-cell sequencing are for the first time able to assess the genome-wide gene expression complements of individual neuroblast identities with the aim to determine the expression of factors that delineate these identities.

Mapping cell fate decisions over the course of embryonic neurogenesis of Drosophila will allow for a more comprehensive understanding of lineage decisions that give rise to the emerging neurogenic cell identities in the early nervous system. My aim is to reveal the exact lineages that give rise to neurons and glia with spatial resolution in vivo. Specifically, the building of a lineage tree for neuroblasts will allow to infer a relatedness of the cells to each other. So far only the common ancestry of cells could be shown, but not when and in which succession those cells separated from each other.

I aim to map lineage decisions and couple the description of lineages with knowledge of when and where molecular markers are expressed.

The lineage of neuroblasts will be recorded with the intMEMOIR method, that relies on a genomic editing process via an integrase and makes cells express the resulting barcode. Image acquisition of the barcode transcript with RNA in situ hybridization will be followed by lineage tree modelling. This should lead to the identification of new marker genes and foster a more detailed understanding about the underlying molecular mechanisms that drive early neurogenic trajectories.

### 370C  Neural Circuits Involved in Nutrient-Dependent Neuroblast Reactivation

**Susan Doyle**, Cami Kellinui, Xin Yuan, Sarah Siegrist University of Virginia, Charlottesville, VA

The precise control of cell growth and division is critical for proper tissue development, integrity, and function. Growth control involves the integration of intrinsic cellular programs with extrinsic cues, including growth factors, hormones, and the availability of nutrients. At the end of Drosophila embryonic development, proliferating neural stem cells, known as neuroblasts, enter into a period of mitotic dormancy called quiescence. Neuroblast exit from quiescence and cell cycle resumption is tightly coupled to nutrient availability; as larvae hatch and begin feeding, amino acids are sensed, resulting in release of Drosophila insulin-like proteins (Dilps) and subsequent activation of the PI3K growth pathway in neuroblasts. Both Dilp2 produced by insulin producing cells (IPCs) in the brain and Dilp6 from glial cells have been shown to mediate the PI3K activation in neuroblasts that results in exit from quiescence. The fat body, a key nutrient sensing tissue, has likewise been shown to be important for both sensing amino acids and producing systemic factors that result in Dilp secretion and neuroblast reactivation in the larval CNS. Here, using animals in which the fat body has been ablated and cultured brain explants from freshly hatched larvae, we demonstrate that the fat body is not required per se for nutrient-dependent reactivation of neuroblasts from quiescence. This suggests that other cell types and tissues capable of amino acid sensing, local brain production of Dilps, and/or uptake of Dilps from IPCs may be capable of driving nutrient-dependent neuroblast reactivation. We further explore this hypothesis using the trans-Tango and DenMark systems to trace pre- and post-synaptic contacts of Dilp2 and Dilp6 producing neurosecretory neurons as well as neurons in the brain expressing the insulin binding protein ImpL2. Using the GAL4/UAS system to express the proapoptotic gene grim and the potassium channel kir2.1 for synaptic silencing, we test the role of specific sets of neuropeptide and neurotransmitter-producing neurons in reactivation of neuroblasts in response to nutrients. Our results shed light on the programs that govern neural stem cell proliferation decisions in response to dietary nutrient availability.

### 371A  A Screen for Amino Acid Transporters Involved in Nutrient-Dependent Reactivation of Quiescent Neuroblasts

**Erik Miao**, Jonathan Day, Susan Doyle, Bharath Sunchu, Sarah Siegrist University of Virginia

In the developing Drosophila nervous system, neural stem cells, called neuroblasts, undergo a period of quiescence at the end of embryogenesis. Neuroblasts exit from quiescence after larvae hatch and begin feeding. Dietary amino acids are required for neuroblast reactivation, however little is known regarding the exact nature of amino acids required and the cell and tissue types involved in sensing them. Insulin signaling is a key regulator of nutrient-dependent growth and metabolism in neuroblasts, and Drosophila insulin-like peptides (Dilps), produced by the insulin-producing cells (IPCs) of the brain, activate PI3-kinase signaling in neuroblasts. Recently, two L-type amino acid transporters expressed in the IPC’s have been shown to be required for release of Dilp2 in response to the amino acid leucine. Therefore, we hypothesized that amino acids directly sensed by the IPCs may play an important role in nutrient-dependent reactivation of neuroblasts from developmental quiescence.

To test this hypothesis, we used the Gal4/UAS binary expression system to disrupt transmembrane amino acid transporters in the IPCs. UAS-RNAi lines for the 26 genes in the solute carrier family that encode AA transporters were...
Sphingolipids are an essential component of cellular architecture. Sphingolipid metabolism regulates cellular processes such as proliferation, growth, apoptosis, inflammation, and senescence. Genetic defects in the metabolism of sphingolipids are associated with several metabolic disorders, including colorectal cancer. The molecular mechanisms expressed under the control of an IPC-specific GAL4 driver (dilp2-GAL4), in combination with a pcnaGFP transgene to assess cell proliferation. Freshly hatched larvae were raised on a standard Bloomington diet, and after 24 hours of feeding, live larvae were removed from food, rinsed and mounted on a glass slide with a coverslip to immobilize them. Brains were imaged in whole mount through the cuticle using epifluorescence, and reactivation from quiescence scored based upon GFP expression.

After screening through 21 of the proposed RNAi lines, no significant reduction of reactivation was found with comparison to the control line (OregonR). This suggests that redundancy exists in nutrient uptake pathways, such that knocking out one AA transporter is not enough to show a significant reduction in reactivation. In the future, we plan to expand our amino acid transporter screen by using Gal4 drivers to express AA transporter RNAi lines in the fat body, glia, and neuroblasts themselves. The results of these experiments may give us valuable information on the tissues and nutrients required to reactivate neural stem cells from quiescence, and in addition may lend insight into improving human stem cell regeneration.

372B Activin signaling controls ISC proliferation and cell fate to maintain adult gut homeostasis Christian Christensen, Julien Colombani, Ditte Andersen University of Copenhagen, Department of Biology

The intestinal tract is the most proliferative organ in the body, and coordination of intestinal stem cell (ISC) activity and cell fate is dynamically tuned to ensure epithelial integrity at homeostasis and in response to injury. To identify niche-derived signals controlling adult gut homeostasis, we recently carried out a large-scale functional genetic screen using RNAis targeting secreted peptides expressed in the ISC niche. Our screen identified several of the evolutionarily conserved TGF-beta/activin ligands suggesting an important function of activin signaling in controlling gut homeostasis. Consistent with this, we found that niche-derived activin-beta signals through the type I activin receptor, baboon, in the stem- and progenitor cells to promote regenerative growth in response to an oral Ecc15 infection. Additionally, we find that activin signaling in intestinal stem- and progenitor cells participate in coordinating proliferation and cell fate decisions during homeostasis to restrict gut epithelial turnover rates and ensure gut integrity. Activin signaling was recently reported to regulate salivary gland stem/progenitor cell function in mice, suggesting a conserved role of activins in controlling adult stem cell activity and tissue homeostasis.

373C Functional analysis of Escargot and STAT targets in intestinal stem cells of the Drosophila melanogaster posterior midgut Armen Khanabaei, Aaron Lemus, Cynthia Petrossian, Donnie Ca, Courtney Frazier, Ithan Cano, Marziah Hossine, Mariano Loza-Cell California State University, Northridge, Los Angeles, CA

Many organs contain adult stem cells (ASCs) that replace cells lost to damage, disease or normal turnover. Like their embryonic counterparts, ASCs can divide asymmetrically, giving rise to a new copy of themselves (i.e. self-renewal) and a sister cell that commits to differentiation into a specific cell type. Previous work has led to the identification of so-called master regulators (MR) genes in ASCs, i.e. pleiotropic genes that directly or indirectly affect the expression and activity of hundreds of downstream effectors, and whose functional alteration simultaneously affects diverse aspects of normal stem cell homeostasis. Genome-wide screens around MR genes have identified hundreds of their putative targets. However, the experimental validation of these bioinformatic predictions and the further characterization of putative downstream effectors as true stem cell regulators is notoriously lacking.

To begin addressing this gap between bioinformatic predictions and experimental validation, we use intestinal stem cells (ISCs) of the Drosophila melanogaster posterior midgut. We first integrated genome-wide DNA mapping data for two known MR genes in these cells: the transcription factor Escargot (Esg) and the signal transducer STAT. We then used qPCR analysis to test the response of over 80 predicted targets to the genetic manipulation of either or both MR genes in vivo, which allowed us to validate several of them as true targets. This analysis revealed a noticeable false discovery rate (FDR) with regards to mode of regulation by the MR genes among the bioinformatic predictions, as reflected by a higher than predicted rate of co-regulation by both Esg and STAT. Furthermore, the genetic manipulation of nearly all confirmed targets that we have tested thus far has resulted in noticeable phenotypes in ISC homeostasis, including changes to their number and/or morphology. Therefore, while our data would indicate that bioinformatic prediction of downstream target regulation by these MR genes is rather noisy, it is nonetheless highly predictive of a true ISC regulatory role for those targets that have been validated experimentally by independent means.

374A Sphingolipid metabolism regulates intestinal stem cell homeostasis M. Mahidur Rahman, Chenge Zhang, Marco Marchetti, Chloe Kraft, Collin Clark, William Holland, Scott Summers, Bruce Edgar 1) Huntsman Cancer Institute, University of Utah; 2) Department of Nutrition and Integrative Physiology, Diabetes and Metabolic Research Center, University of Utah, Salt Lake City, Utah, USA

Sphingolipids are an essential component of cellular architecture. Sphingolipid metabolism regulates cellular processes such as proliferation, growth, apoptosis, inflammation, and senescence. Genetic defects in the metabolism of sphingolipids are associated with several metabolic disorders, including colorectal cancer. The molecular mechanisms
involved in sphingolipid metabolism-mediated intestinal stem cell (ISC) homeostasis and tumorigenesis are poorly understood. We do not know if and how the intermediate metabolites in the anabolic or catabolic processes of the ceramide metabolism regulate ISC fates in the adult midgut. To check if any of the sphingolipid metabolic enzymes are involved in the ISC homeostasis, we have genetically perturbed these enzymes in the specific cell types of the ISC lineages and the progenitor cells together. We found that the perturbation of different enzymes in the ceramide metabolism affects each cell type differently. Overall, the effects on the progenitors are manifested differently than those on the terminally differentiated enterocytes and enteroendocrine cells. For example, reducing ceramide synthases such as lace strongly affects enteroendocrine cell morphology compared to the ISCs or enteroblasts. Over-expression of the ceramidase bwo in the enteroblasts leads to robust proliferation of the ISCs, besides the increased size of the enteroblasts. On the other hand, the enterocytes are highly sensitive to any perturbation of the enzymes involved in the sphingolipid metabolism. Both increased or decreased levels of the enzymes kill the ECs, which in turn promote ISC proliferation. Since the EGFR signaling pathway is an important driver for ISC proliferation, we probed if the activated EGFR pathway would regulate the sphingolipid metabolism. Transcriptomic and lipidomic analysis of the Spitz activated EGFR pathway in the gut shows that the EGFR pathway can upregulate genes encoding ceramide biosynthesis as well as other sphingolipids. These results suggest that different enzymes and products of the sphingolipid metabolic pathway can act as signaling molecules and interact with major stem cell signaling pathways such as EGFR signaling to affect ISC homeostasis.

375B  Identifying factors that maintain the adult testis niche Gabriela Vida, Elizabeth Botto, Stephen DiNardo University of Pennsylvania

Stem cells are important for repairing and regenerating our tissues, and often reside in a niche that controls their behavior. The testis niche has been a paradigm for niche-stem cell interactions. Recently, our lab has focused on the construction of this niche. Examining initial assembly of the niche in embryonic gonads has revealed both how extra-gonadal signals drive niche assembly (Anllo and DiNardo, 2021), and begun illuminating the cell mechanics involved in first forming a compact, functional niche (B. Warder poster). Here we use the adult testis niche to address the cell biological features that maintain niche structure and function during its steady-state operation. The niche resides at the testis tip and is comprised of a group of quiescent cells that send renewal signals to the neighboring stem cells. These quiescent cells are organized spherically, and are radially surrounded by two stem cell populations, the germline stem cells and the cyst stem cells. Our preliminary evidence suggests that acto-myosin contractility (AMC) is important for the maintenance of the spherical nature of the stem cell niche. The recently released single nuclear, RNA-SEQ Fly Cell Atlas (FCA) for the testis is a resource to suggest genes that might be candidates involved in niche structure. Center-divider (cdi) is suggested as enriched in the niche based on the FCA data and encodes a serine/threonine kinase known to induce actin remodeling through phosphorylation of cofilins. Our preliminary evidence shows that knockout of cdi indeed affects maintenance of testis niche structure. We therefore hypothesize that the AMC and actin remodeling are both important for preserving niche structure and potentially critical for the maintenance of stem cells.

376C  The role of ESCRTs in signaling within the testis stem cell niche Mara Grace, Erika Matunis Johns Hopkins University, Department of Cell Biology

Adult stem cells are crucial for regeneration, tissue repair after injury, and developmental processes such as spermatogenesis. Stem cells exist in a dynamic microenvironment termed the niche that provides signals to ensure the maintenance and self-renewal of the adult stem cell population. An appreciation of the dynamic communication between stem cells and their niche is vital to understand processes such as reproduction, oncogenesis, aging, development, and regeneration. Here I use the testis of Drosophila melanogaster as a model to investigate the role of endocytosis, and the ESCRT complexes specifically, in signaling within the stem cell niche. Endocytosis regulates a myriad of signaling pathways as well as cellular communication. The ESCRT complexes are involved in a variety of cellular processes, such as multivesicular body formation and particle budding, and are considered endocytic tumor suppressor genes due to their role in signal attenuation. Knockdown of ESCRTs in somatic stem cells causes niche cells to become significantly enlarged and exhibit abnormal morphology. Furthermore, Unpaired, a ligand for the JAK-STAT pathway normally secreted by niche cells, accumulates within the cytoplasm of niche cells upon ESCRT knockdown in somatic stem cells. As knockdown of ESCRTs in somatic stem cells affects the morphology of niche cells, this suggests that ESCRTs mediate signaling from somatic stem cells back to their niche to prevent niche hypertrophy. I am further exploring the role of additional pathways in modulating this signaling. A deeper understanding of the signaling dynamics within the Drosophila testis stem cell niche will have further implications for stem cell niches in other tissues and organisms as well as processes such as regeneration and renewal.

377A  Drosophila Holes in muscles is required for ongoing adult muscle function and muscle stem cell maintenance. Robert Hoff, Richard Cripps San Diego State University

The Drosophila gene Holes in muscles (Him) is a mesodermal transcriptional corepressor that functions in myogenesis. Previously, we and others have characterized Him expression in the early embryonic mesoderm where it functions...
downstream of known muscle specification genes as a repressor of the myogenic transcription factor MEF2 to delay muscle differentiation until the appropriate stage. Here, we extend our analysis of Him to other stages of Drosophila muscle development and function by analyzing the phenotype of the first-generated Him mutant allele. Him mutants show a reduction in the imaginal disc associated myoblast pool that form the adult thoracic muscles, including those required for jump and flight. Consistent with this observation, we find that Him mutants have reduced myonuclei in adult differentiated flight muscles. To further characterize the role of Him in adult muscle function, we assayed flight performance in young and aged flies. Him mutants perform comparably to wild type controls at young (3-5 days) of age, however have significantly reduced flight ability at two weeks of age, suggesting a role for Him in continued muscle maintenance. Additionally, we find that the flightless phenotype correlates with the number of somatic muscle stem cells associated with the flight muscles. Young Him mutants and control flies have a similar number of stem cells, but this number is significantly reduced in the mutants at two weeks of age. Altogether, these data suggest an ongoing role for Him in muscle beyond early development potentially mediated through regulating myonuclear number and stem cell maintenance.

### 378B  Assessment of cellular and functional heterogeneity within the Drosophila testis stem cell niche
**Jennifer Viveiros,** Erika Matunis Department of Cell Biology, Johns Hopkins University School of Medicine

Adult stem cells reside in dynamic, supportive microenvironments termed niches, which are generated by specialized niche cells. Niches are often complex and composed of functionally cooperative subpopulations of cells rather than uniform populations, and gaining insight into niche composition underpins our understanding of tissue homeostasis. Therefore, our ultimate goal is to further our understanding of stem cell niches by investigating their cellular composition and gene expression programs, using the *Drosophila* testis stem cell niche as a model. This niche contains three cell populations: post-mitotic somatic hub cells (or niche cells), which are surrounded by two types of stem cells, germline stem cells (GSCs) that differentiate into sperm and cyst stem cells (CySCs) that give rise to somatic support cells. Previous *in situ* hybridization (ISH) and immunostaining experiments have suggested that hub cells may not uniformly express the same transcriptional program, raising the possibility that they are a heterogeneous population of cells. Intriguingly, hub cells descend from somatic gonadal precursors (SGPs), which arise from three different parasegments (PS) in embryogenesis, suggesting that developmental origin could underly previously observed heterogeneity. Here we use the HACK and Split-Gal4 ternary systems to genetically control individual populations of SGPs. Preliminary results suggest that combinatorial patterns of homeobox gene expression allow genetic dissection of SGPs arising from differing PS. These tools will allow lineage tracing and manipulation of the distinct populations of SGPs according to their origin to begin to determine if hub cell origin correlates with function, localization within the niche, and transcriptional program.

### 379C  Investigating somatic stem cell cytokinesis and coordination of daughter cell release from the testis niche
**Tiffany Roach,** Kari Lenhart Drexel University

Stem cells rely on instructional cues from their specialized microenvironment, or niche, to balance the production of self-renewing and differentiating daughter cells. Many adult tissues must coordinate the divisions and behaviors of multiple stem cell lineages to generate and maintain tissue. The *Drosophila* testis niche exhibits coordination between germline stem cells (GSCs) and cyst stem cells (CySCs) such that each differentiating GSC daughter must become completely encapsulated, or encysted, by two daughters of the CySC lineage. Our lab has found a modified cytokinesis program in GSCs that is critical for coordinated release of differentiating daughters. However, almost nothing is known about the cytokinesis program and cellular dynamics of the CySC lineage that may be acting in conjunction to facilitate this interaction. By establishing a combination of fluorescent markers to live image the CySC lineage, I will investigate CySC cytokinesis to determine the presence of a modified cytokinesis program that may be operating in tandem with GSC cytokinesis to enable coordinated release of stem cell daughters. Ultimately, this powerful new imaging system will permit us to identify the cell biology and temporal dynamics by which the 2:1 soma to germline ratio is established for tissue homeostasis. This work will extend our knowledge of the interactions and complex dynamics of two distinct stem cell populations within the testis niche.

### 380A  Characterizing the Novel Protein Asperous Involved in Tissue Regeneration
**Si Cave,** Robin Harris Arizona State University

The mechanisms of tissue regeneration are not fully understood. Using the model of the wing imaginal disc, we have identified factors upregulated during regeneration through RNA-sequencing. Asperous (CG9572) is a novel and mostly uncharacterized protein that is strongly transcribed specifically during regeneration. The goal of this work is to characterize Asperous in the context of regeneration, using a three-pronged approach to understand 1) its protein structure, 2) its localization, and 3) its cellular function in both damaged and undamaged imaginal disc tissues.

To do this, we have utilized computational tools to predict the structure and found the presence of a WD40 domain, suggesting a role in coordinating multi-protein complex assemblies. We have also used genetic manipulations via our unique genetic ablation tool, DUAL Control, to knockdown Asperous and have found it is essential for regeneration.
Finally, we are using fluorescent reporters of tagged Asperous proteins to understand its localization, which suggests it may be extracellular and contribute to differential growth in a compartment-specific manner. As Asperous has a uniquely regeneration-specific expression pattern and is associated with promoting regeneration in the wing disc, investigating the features of this novel protein will further our understanding of uncharacterized elements of a regeneration program.

381B Genetic determinants of cell fate plasticity during regeneration after radiation damage in Drosophila Caitlin Clark, Michelle Ledru, Jeremy Brown, Shay Segal, Shelpi Verghese, Sarah Ferrara, Andrew Goodspeed, Tin Tin Su. 1) Department of Molecular, Cellular and Developmental Biology, University of Colorado, Boulder, CO; 2) University of Colorado Cancer Center, Anschutz Medical Campus, Aurora, CO; 3) College of Veterinary Medicine and Biomedical Sciences, Colorado State University, Fort Collins, CO; 4) Department of Cell Biology, Emory University School of Medicine, Atlanta, GA

Ionizing radiation (IR) is used to treat approximately half of all cancer patients due to its ability to induce cell death. However, IR can also induce cancer stem cell-like properties in non-stem cancer cells, potentially promoting tumor regeneration and diminishing therapeutic success. Our published studies have shown that we can model IR-inducible stem cell-like properties in Drosophila melanogaster larval imaginal wing discs. After irradiation, IR-resistant cells from the hinge-region of the disc translocate to the pouch-region of the disc and change fate to help regenerate the pouch (Verghese and Su, 2016, PMID: 27584613; Verghese and Su, 2018, PMID: 30462636). Genome-wide RNAseq analysis of dissociated wing disc cells identified IR-induced gene expression changes specific to the hinge such as down-regulation of hinge determinants and up-regulation of Myc targets and ribosome biogenesis genes. Functional testing shows that Myc and genes that function in ribosome biogenesis are required specifically in the hinge for IR-induced fate change (Ledru et al., in revision for PLoS Genetics). Current efforts are directed at using hinge-specific ribosome profiling to identify mRNAs whose increased translation promotes cell fate change after irradiation.

382C Necrosis-induced apoptosis promotes regeneration in Drosophila wing imaginal discs Jacob Klemm, Robin Harris Arizona State University

Cell death is essential for the proper regeneration of tissues following injury. Dying cells serve as crucial signaling centers that promote cell proliferation and tissue remodeling following damage. Much of our understanding of these events comes from studies in Drosophila imaginal discs. For example, apoptotic cells release Wg and Mmp1, among other factors, to adjacent cells to promote tissue repair. This phenomenon been extensively characterized with respect to apoptosis-induced regeneration, however, much less is understood about how tissues respond to unregulated forms of death, like necrosis. Necrosis is a rapid, disordered cell death in which cell membrane integrity is lost and intracellular contents are released to the external environment. To study the regenerative response to necrotic tissue death, we developed a genetic ablation system that drives necrosis in the Drosophila wing imaginal disc.

With this new model, we have found that necrosis leads to a unique regenerative response. Immunofluorescent staining reveals significant apoptosis is induced at a distance from the wound, which we have termed necrosis-induced apoptosis (NIA). Unlike other damage-associated apoptosis in the wing disc, NIA cells are not regulated by the JNK pathway. Moreover, NIA cells are required for regeneration following necrosis; inhibition of NIA cell activity leads to lower levels of proliferation at the wound edge. Consequently, NIA cell inhibition results in a reduced capacity to regenerate, as assayed by adult wing size. This research shows that upon necrotic tissue death, wing discs rely on apoptotic signaling to facilitate regeneration.

We are currently focused on identifying the signals that lead to NIA cell activity and understanding the mechanism by which these cells promote proliferation. Preliminary data highlights cytokine and calcium signaling as potential regulators of NIA cells. Additionally, lineage tracing shows NIA cell progeny may contribute to the reconstruction of the salm domain. To comprehensively identify the mechanisms that define NIA cell activity, transcriptome data will be generated for NIA cells and wound edge regenerating cells. These data will characterize the tissue response to necrosis and provide insight to designing therapeutics for humans afflicted by necrotic injuries and diseases.

383A Elucidating The Roles of Zelda and Taranis During Late Regeneration in Drosophila Wing Imaginal Discs Anish Bose, Keaton Schuster, Rachel Smith-Bolton. 1) University of Illinois at Urbana Champaign; 2) New York University

Regeneration is a complex process that enables damaged tissues to be replenished and restored back to their correct morphology and function. As organs and tissues are prone to damage either by external forces or chronic illness, the identification of factors and mechanisms important for regeneration may have clinical significance. Our laboratory has described a regeneration-specific mechanism that stabilizes cell fate after damage, where the gene taranis maintains proper cell fate during late regeneration in Drosophila 3rd instar larval wing imaginal discs. Reduced Taranis levels allow damage-induced JNK signaling to overexpress engrailed. Overexpressed engrailed initiates a negative feedback loop that silences the engrailed locus, resulting in impaired patterning during regeneration, including posterior-to-anterior cell-fate transformations. Taranis prevents these posterior-to-anterior cell-fate changes by preventing engrailed overexpression. However, it is unclear which signals activate taranis expression during late regeneration, or how engrailed is regulated by JNK signaling or Taranis. Preliminary data showed that the pioneer transcription factor zelda was upregulated.
during late regeneration, and in a reduced zelda background, taranis reporter expression was also reduced, suggesting that Zelda may be an activator of taranis. Furthermore, zelda may have additional roles beyond activating taranis in the regenerating wing imaginal disc. Finally, we are also examining how engrailed expression is misregulated by JNK signaling, and how that misregulation is prevented by Taranis.

**384B The epithelial apical-basal polarity regulator Lgl constrains imaginal disc regeneration** Faith Karanja, Rajan Bhandari, Thu Tran, Adrian Halme University of Virginia  

Mutation of the neoplastic tumor suppressor lethal (2) giant larvae (lgl) in Drosophila melanogaster imaginal disc epithelia produces neoplastic tumors that elicit systemic responses, developmental delay, and growth inhibition, similar to those produced by regenerating imaginal discs. We observe that critical systemic and local regenerative signaling pathways are activated in lgl-mutant tumors, including expression of the relaxin-family hormone Dilp8, which is responsible for developmental delay and growth control during regeneration, and Wingless (Wg), which regulates growth and patterning in the wing during regeneration and normal development. Wg activation during regeneration is mediated by a downstream regulatory region, the damage responsive element (DRE), which we demonstrate is also responsible for Wg activation in lgl tumors. We observed that removal of the DRE has little effect on wingless expression during normal disc development but severely reduces the regenerative capacity of imaginal tissues and prevents neoplastic transformation as well as neoplasia-mediated developmental delay. These observations led us to hypothesize that Lgl might function to limit regenerative activity that promotes the neoplastic transformation of discs. Supporting this hypothesis, we observed that: 1) inducing damage and regeneration in imaginal discs promotes the neoplastic potential of lgl clones, which is blocked by removing the DRE, and 2) expression of a phosphorylation-resistant allele of Lgl strongly inhibits disc regeneration. Finally, during our analysis of lgl clones in the wing imaginal disc, we found unexpected heterogeneity in the neoplastic potential and the DRE dependence between clones. This heterogeneity may reflect differences in regenerative pathways and potential found in different regions of the developing tissue. These results demonstrate an unexpected role for the epithelial apical-basal polarity regulator Lgl in regulating regenerative activity in imaginal discs.

**385C Inducing limb regeneration in Drosophila melanogaster** Michael Abrams1 3, Fayth Tan1, Yutian Li3, Ty Basinger1 5, Martin Heithe1, Anish Sarma1, Iris Lee1, Zevin Condiotte1, Misha Raffiee1 4, John Dabiri1 2, David Gold1 6, Lea Goentoro1 1) Division of biology and Biological Engineering, California Institute of Technology, Pasadena, CA ; 2) Graduate Aerospace Laboratories and Mechanical Engineering, California Institute of Technology, Pasadena, CA; 3) Department of Molecular and Cell Biology, University of California at Berkeley; 4) Department of Bioengineering, Stanford University; 5) Department of Biology and Allied Health Sciences, Bloomsburg University; 6) Department of Earth and Planetary Sciences, University of California at Davis  

Can limb regeneration be induced? Drosophila belongs to the holometabolon insects, which do not regenerate limbs or other appendages as adults. However, developing imaginal discs in larvae can regenerate, possibly suggesting an inherent ability to regenerate. In this study, we found that limb regeneration in adult flies can be induced. Strikingly, limb regeneration can be induced using, simply, nutrient supplementation with the amino acid L-leucine and the growth hormone insulin. Almost 50% the treated flies partially regrew their amputated limbs. The new leg part redeveloped cuticles and sensory bristles. Some flies completely reformed the amputated leg segment, followed by a joint and the beginning of next segment. This is the first demonstration that patterned regenerative response can be induced in adult Drosophila limb. Finally, the same strategy of leucine and insulin/sugar administration induce appendage regeneration in the moon jellyfish and digit regeneration in mice. These results suggest that latent regeneration is prevalent in animals and can be activated with relatively simple environmental stimuli. Our results present Drosophila as a novel system for studying how to induce regeneration.


**386A Adapting the Nitroreductase Cell Ablation System to Drosophila** Gary Teeters, Sarah Siegrist University of Virginia  

Targeted cell ablation is a common method used to assay cell function during development and to determine cellular limits during regeneration. Many techniques are available to achieve cell ablation, yet improvements are still needed to increase spatial and temporal control. One method commonly used in the Zebrafish, Danio rerio is a Nitroreductase (NTR) mediated ablation system. In this technique cells expressing NTR can be induced to undergo cell death following drug treatment. For example, when metronidazole (MTZ) is exposed to cells expressing Nitroreductase, NTR converts MTZ into a DNA cross linker inducing cells to die. We set out to determine whether this commonly used cell ablation system could be used in Drosophila. We cloned the Nitroreductase gene from Escherichia Coli used in Curado et al. 2007 with a GFP reporter to examine cell death in vivo. This technique would allow for greater temporal control of cell ablation
without the need of temperature sensitive Gal80 or mutant alleles instead by regulating exposure to nitroimidazole antibiotics. This project means to adapt the protocols from Danio rerio to Drosophila melanogaster which will modify the exposure method, concentration, and time. We aim to show that exposure to nitroimidazole antibiotics through ingestion at sub toxic levels will be able to ablate cells in Gal4 specific manner with temporal control and no off-target toxicity.

387B The Role of dMyc in Drosophila wing imaginal disc regeneration Felicity (Ting-Yu) Hsu, Rachel Smith-Bolton University of Illinois at Urbana-Champaign

Regeneration is a complicated process through which some animals restore missing tissue upon damage. While many signaling pathways crucial for regeneration are critical in normal development, regeneration is not as simple as recapitulating normal development. We study regeneration using our genetic ablation system in Drosophila wing imaginal discs. One of the genes that are important in regeneration is Myc, whose expression levels and transcription levels are upregulated in the regenerating region of wing imaginal discs. Overexpression of Myc improves wing imaginal disc regeneration, while reduction of Myc expression worsens wing disc regeneration. During normal development, one primary role of Myc is ribosomal biogenesis. However, the exact function of Myc in regeneration is unknown. Our recent findings suggest that the nucleoli sizes, indicating ribosomal biogenesis activity, in the regenerating wing pouch are similar to those in the undamaged pouch. Interestingly, smaller nucleoli were found in the undamaged hinge tissue of the damaged wing discs. This finding suggests that one possible role of Myc might be maintaining ribosomal biogenesis activity in the regeneration blastema while the activity is reduced in other parts of the damaged disc. Thus, we aim to identify the exact role of upregulated Myc in wing disc regeneration. In addition to ribosomal biogenesis, Myc is also involved in cell competition. After tissue damage, the difference in Myc expression levels between the regeneration blastema and the rest of the disc is increased, as Myc is only upregulated in the regenerating tissue. During normal development, Myc expression level differences in the wing imaginal disc lead to cell competition, where cells with lower Myc are eliminated. However, we observed no extreme cell death at the boundary between high and low Myc expression levels in regenerating wing discs. Therefore, we aim to study whether cell competition takes place in regenerating wing discs.

388C Wear and Tear of the Intestinal Visceral Musculature by Intrinsic and Extrinsic Factors Ho Kim¹, Eric So¹, Jiae Lee¹, Yi Wang², Vikram Gill³, Anna Gorbacheva³, Hee Jin Han¹, Katelyn Ng¹, Ken Ning¹, Inez Pranoto¹, Alejandra Cabrera¹, Dae Seok Eom³, Young Kwon¹ 1) University of Washington; 2) University of California Irvine

The gut visceral musculature plays essential roles in not only moving substances through the lumen but also maintaining the function and physiology of the gut. Although the development of the visceral musculature has been studied in multiple model organisms, how it degenerates is poorly understood. Here, we employ the Drosophila midgut as a model to demonstrate that the visceral musculature is disrupted by intrinsic and extrinsic factors, such as aging, feeding, chemical-induced tissue damage, and oncogenic transformation in the epithelium. Notably, we define four prominent visceral musculature disruption phenotypes, which we refer as ‘sprout’, ‘discontinuity’, ‘furcation’, and ‘crossover’ of the longitudinal muscle. Given that the occurrence of these phenotypes is increased during aging and under various stresses, we propose that these phenotypes can be used as quantitative readouts of deterioration of the visceral musculature. Intriguingly, administration of a tissue-damaging chemical dextran sulfate sodium (DSS) induced similar visceral musculature disruption phenotypes in zebrafish larvae, indicating that ingestion of a tissue-damaging chemical can disrupt the visceral musculature in a vertebrate as well. Our study provides insights into the deterioration of the gut visceral musculature and lays a groundwork for investigating the underlying mechanisms in Drosophila as well as other animals.

389A Transition from acute nerve injury to central sensitization requires metabotropic driven astrocyte store-operated Ca²⁺ entry Mariya Prokhorenko, Jeremy Smyth Uniformed Services University of the Health Sciences

Chronic pain following injury is a debilitating condition associated with serious comorbidities and opioid dependence. Understanding the neurological mechanisms that underlie the transition from initial injury to chronic pain is essential for prevention and treatment. This transition is mediated by central sensitization, which involves reversible changes to nociceptive neurocircuitry pathways in the central nervous system. Central sensitization research and therapeutics have focused on neuronal circuitry, whereas the roles of glial cells are poorly defined. Notably, astrocytes that normally maintain and modulate neuronal synapses become reactive and exhibit aberrant Ca²⁺ signals during central sensitization. However, how these Ca²⁺ signals are generated and how they modulate astrocyte function in central sensitization are unknown. Importantly, central sensitization involves significant upregulation of glutamate release at nociceptive synapses, and glutamate acts on astrocytes via metabotropic glutamate receptors to stimulate Ca²⁺ release from endoplasmic reticulum (ER) Ca²⁺ stores via inositol 1,4,5-triphosphate receptors (IP₃R). We are testing the hypothesis that central sensitization requires astrocyte metabotropic signaling involving IP₃R-mediated Ca²⁺ release from ER Ca²⁺ stores and activation of store-operated Ca²⁺ entry (SOCE). We are testing this by combining powerful genetic tools with a robust model of central pain sensitization in Drosophila melanogaster. We assay central sensitization in flies by
amputating a single leg and monitoring the response to subnoxious temperatures. Injured, sensitized animals exhibit increased jumping behavior at the subnoxious temperature, whereas animals without injury do not respond to this temperature. Using a combination of fluorescence imaging techniques, we show enhanced astrocyte Ca\(^{2+}\) signaling in the ventral nerve cord precedes behaviorally measured central sensitization after injury, including changes in live signaling using GCaMP and cumulative signaling using Transcriptional Reporter of Intracellular Calcium (TRIC). We also found that astrocyte-specific suppression of IP\(_R\) as well as the SOCE components Stim and Orai attenuates nociceptive jumping behavior at one week following nerve injury, suggesting a key role for IP\(_R\) and SOCE-mediated Ca\(^{2+}\) signaling in astrocytes during pain sensitization. Our results will bring new understanding to the role of astrocyte signaling in pain sensitization and may suggest novel therapeutic targets for the prevention or treatment of chronic pain.

390B  Wound-induced changes in epithelial tension  Ivy Han, James White, James O’Connor, M. Shane Hutson, Andrea Page-McCaw Vanderbilt University, Nashville, TN, USA

Understanding the mechanisms of epithelial wound healing is a major challenge in biology and medicine. *Drosophila* is a great model to study wound healing and has contributed significantly to identifying how epithelial tissue repairs a wound through coordinated changes in cellular behavior (e.g., cell migration). Wound-induced epithelial cell migration is preceded by dynamic changes in cytoskeletal structures and in tissue tension. However, it is unclear how epithelial wounding immediately affects tension in nearby epithelial cells or whether changes in tension play a role in the wound response. This study aims to characterize how wounding alters the cortical tension of apical cell borders in epithelial wound through coordinated changes in cellular behavior (e.g., cell migration). Wound-induced epithelial cell migration is preceded by dynamic changes in cytoskeletal structures and in tissue tension. However, it is unclear how epithelial wounding immediately affects tension in nearby epithelial cells or whether changes in tension play a role in the wound response. This study aims to characterize how wounding alters the cortical tension of apical cell borders in epithelial cells and determine factors that modulate these wound-induced tension changes. We use a laser-recoil assay to measure response. This study aims to characterize how wounding alters the cortical tension of apical cell borders in epithelial cells and determine factors that modulate these wound-induced tension changes. We use a laser-recoil assay to measure response. This study aims to characterize how wounding alters the cortical tension of apical cell borders in epithelial cells and determine factors that modulate these wound-induced tension changes. We use a laser-recoil assay to measure response. This study aims to characterize how wounding alters the cortical tension of apical cell borders in epithelial cells and determine factors that modulate these wound-induced tension changes. We use a laser-recoil assay to measure response. This study aims to characterize how wounding alters the cortical tension of apical cell borders in epithelial cells and determine factors that modulate these wound-induced tension changes. We use a laser-recoil assay to measure response. This study aims to characterize how wounding alters the cortical tension of apical cell borders in epithelial cells and determine factors that modulate these wound-induced tension changes. We use a laser-recoil assay to measure response. This study aims to characterize how wounding alters the cortical tension of apical cell borders in epithelial cells and determine factors that modulate these wound-induced tension changes. We use a laser-recoil assay to measure response. This study aims to characterize how wounding alters the cortical tension of apical cell borders in epithelial cells and determine factors that modulate these wound-induced tension changes. We use a laser-recoil assay to measure response.

Further investigation on connections between epithelial tension and other established wound responses such as calcium signaling, cell fusion, and cell migration could pave the way for deeper understanding of the mechanobiology of epithelial wound healing.

391C  The Role of Polyploidy During *Drosophila* Epithelial Wound Repair  James White, Kimi LaFever Hodge, Ivy Han, Jasmine Su, M. Shane Hutson, Andrea Page-McCaw Vanderbilt University

In the past decade, there has been increased awareness of cellular polyploidy in development, homeostasis, and cancer. In addition to being a conserved developmentally programmed behavior, polyploidy is also induced in response to injury. Work by Vicki Losick has shown adult *Drosophila* epithelia repair by becoming polyploid through endoreplication and cell-cell fusion. In Zebrafish epicardial explants, regeneration is led by a wavefront of polyploid cells. We have found that both endocycling and cell fusions occur after laser wounding in the *Drosophila* pupal notum epithelium, and controls expressing Rok-RNAs confirmed that the technique could identify cells with reduced cortical tension. We discovered that within minutes after wounding, tension was reduced in epithelial cells closer to the wound in a distance-dependent manner. Wounds were administered by a single-shot pulsed-laser ablation, which causes a cavitation bubble that damages cells around the wound. Using a different method of laser-wounding, we found that plasma membrane microtears from the laser-induced cavitation bubble were necessary for reducing cortical tension near the wound. Previously, our group identified that the G-Protein Coupled Receptor (GPCR) Methuselah-like 10 (Mthl10) is required for calcium signaling within about a minute after wounding. Our preliminary results show that knockdown of *mthl10* alters tension around the wound, abolishing the distance-dependent tension reduction after single-shot pulsed-laser wounding. This suggests that Mthl10 may be a key factor that modulates wound-induced tension changes, either by preventing tension loss or restoring tension at locations distal to the wound. Further investigation on connections between epithelial tension and other established wound responses such as calcium signaling, cell fusion, and cell migration could pave the way for deeper understanding of the mechanobiology of epithelial wound healing.

392A  Insulin receptor/Akt/TOR signaling regulates muscle stem cell pool in *Drosophila*  Kumar Vishal, Richard Cripps San Diego State University, San Diego, CA

Indirect flight muscles (IFMs) are the largest muscles in *Drosophila* and are made up of hundreds of myonuclei. The generation of these giant muscles requires a large pool of wing disc associated muscle stem cells. To achieve this, there is an activation of quiescent muscle stem cells followed by a rapid scaling up of the pool in the postembryonic stages, where the number of adult muscle stem cell increases from 10-20 to 2500 within 4 days. However the factors that control activation and proliferation to form this muscle stem pool are incompletely known. The goal of this study is to examine the role of Insulin receptor/Akt/TOR signaling in the regulation of muscle stem cell. We find that blocking the
components of Insulin receptor/Akt/TOR signaling results in a diminished myoblast pool. This reduction in the pool size is due to decreased muscle stem cell activation and proliferation. Disrupting the Insulin receptor/Akt/TOR signaling at initial stages of larval development results in an absence of myoblast pool activation. On the other hand, abrogating signaling in the late larval stage reduces muscle stem pool. Furthermore, our results demonstrate that the attenuation of the Insulin receptor signaling leads to complete absence of IFM formation. By contrast, activating the pathway increases the pool size and the proliferative capacity of the muscle stem cells. Finally our genetic interaction data suggests that Akt interacts with a pseudokinase Tribbles to regulate muscle stem cell population. Collectively, our studies identify a novel role for Insulin receptor/Akt/Tor signaling in the activation and proliferation of the muscle stem cell pool.

393B  Cell cycle exit and stem cell differentiation are coupled through regulation of mitochondrial activity in the Drosophila testis Dieg0 Sainz de la Mazo1, Silvana Hof-Michel2, Lee Phillimore3, Christian Bökel1, 2, 3, Marc Amoyel1 1) University College London, UK; 2) Philippus University Marburg, Germany; 3) Universität Ulm, Germany

Stem cells maintain tissue homeostasis by proliferating to replace cells lost to damage or natural turnover. To do so, they need to balance self-renewal and differentiation. Whereas stem cells proliferate, terminal differentiation is accompanied by exit from the cell cycle, so we asked how stem cell identity was coordinated with cell cycle exit. We use the Drosophila testis as a model. The testis niche supports two stem cell populations, germ line stem cells and somatic cyst stem cells (CySCs) by providing signals necessary for their self-renewal. CySCs proliferate while their differentiating daughter cells do not, providing an ideal model to ask how stem cell identity is linked to proliferation. We show that Cyclin E is required for CySC self-renewal and blocking the G1/S transition causes premature differentiation. Conversely, knocking down Rbf, the homologue of Retinoblastoma that inhibits the G1/S transition, expanded the CySC population and blocked its differentiation. Rbf functions by inhibiting the complex E2f1/Dp. Surprisingly, E2f1/Dp activity was not required for self-renewal, suggesting that the endogenous role E2f1/Dp activity is not in promoting cell cycle progression but to prevent differentiation in cycling cells. To determine how Rbf inhibited differentiation, we analysed gene expression upon Rbf knockdown. Genes regulating mitochondrial biology and energy production were downregulated in Rbf-deficient testes. Promoting mitochondrial biogenesis and activity in Rbf knockdowns by expressing the PGC1α homologue, Spargel (Srl) and Ets97D/NRF-2 rescued differentiation of CySCs. However, we observed ectopic cycling in rescued, differentiated cells suggesting that oxidative metabolism is essential for CySCs differentiation but is not sufficient to exit the cell cycle. Thus, our results indicate that Rbf coordinates cell cycle exit and differentiation by inhibiting E2f1/Dp and promoting a metabolic state that is permissive for differentiation.

394V  Molecular mechanisms behind adult muscle stem cells specification and activation. Hadi Boukhhatmi1, 2, 5, 6, Nourhene Ammar1, 2, Sarah Bray5, 6, Jennifer Zannet3, 4, François Payré5, 6 1) Institut de Génétique et Développement de Rennes, CNRS, Rennes ; 2) Université de Rennes 1 ; 3) Molecular, Cellular and Developmental Biology Department (MCD), Centre de Biologie Intégrative (CBI), CNRS, Toulouse ; 4) Université Paul Sabatier ; 5) University of Cambridge ; 6) Department of Physiology, Development and Neuroscience.

To compensate for damages and cell death, adult tissue homeostasis requires regeneration operated by long-lived stem cells. Skeletal muscles are regenerated by muscular stem cells called satellite cells (MuSC), which are quiescent and, upon injury, become activated and differentiate into myoblasts to repair muscles. A key question is how such cells are specified and then protected from differentiation for a prolonged developmental period. We discovered that MuSC maintenance involves persistent expression of the Zfh1/ZEB Transcription Factor (TF). We showed that sustained Zfh1 expression, in MuSC requires production of an alternate zfh1 RNA isoform, which is insensitive to microRNAs (miR8/miR200) degradation. Zfh1 protein is thus maintained in these cells, enabling them to escape differentiation and persist in the adult and contribute to muscle homeostasis. The key role of ZEB1 in the control of mammalian MuSCs maintenance has been recently reported, showing the broad relevance of Drosophila toward a better understanding of MuSC regulation. While MuSC specification and differentiation starts to be well understood, little is known about the genetic control that underlies the early steps of their activation. Our results revealed that another conserved TF, OvoL/Shavenbaby (Svb), is expressed in adult MuSCs. Svb is the unique Drosophila member of the OvoL family of TFs, which are emerging as key regulators of epithelial tissues and stem cells. We found that the Svb TF binds in vivo to zfh1 cis-regulatory regions suggesting a direct regulation of zfh1 transcription by Svb. Interestingly, svb is rapidly turned off in new-born differentiating progenies that replenish the muscle fibre. Upon svb loss of function in adult MuSCs, we observed dramatic increase in the rate of myogenic progenitor production indicating that Svb controls the frequency of MuSC division rate. Building on these data, we concluded that Svb have a dual role in MuSCs: a) Maintaining stemness through zfh1, b) Acting as an early activating gene by regulating the rate of MuSC division.

395V  Regulation of Damage-Responsive Maturity-Silenced enhancers in Drosophila John Quinn, Robin Harris Arizona State University

Regeneration is a complex process that occurs in a variety of different organisms. In early larval stages, the imaginal discs of Drosophila — the precursors to the adult appendages — have a significant capacity to regenerate that is lost as the organism matures. Using a genetic ablation method established by our group we have found that several genes involved
in disc regeneration are regulated by damage-responsive and maturity-silenced (DRMS) enhancers. These regulatory elements are activated upon damage to induce regenerative gene expression, but epigenetically silenced as the organism approaches pupariation. The goal of my research is to investigate what specific signals activate DRMS enhancers and how they become progressively silenced as discs mature.

The genes wg and Wnt6 are activated during regeneration and controlled by a single DRMS enhancer (DRMS\textsuperscript{Wnt}). We found that JNK signaling is necessary for the activation of DRMS\textsuperscript{Wnt} but not sufficient, since developmental JNK signaling does not activate the enhancer. The cells contributing to regeneration co-express both JNK and JAK/STAT consistent with previous data that shows these pathways are important for regeneration. Using a GFP reporter for DRMS\textsuperscript{Wnt} we found that reducing JAK/STAT pathway activity via STAT92E RNAi decreases damage-induced activation of the reporter, suggesting a role for JAK/STAT in the activation of the enhancer. Although, this decrease indicates that JAK/STAT is involved in regeneration, it does not show whether STAT is directly interacting with the enhancer DNA. To test this interaction, we generated a DRMS\textsuperscript{Wnt} GFP lacking the consensus transcription factor binding sites to STAT92E, which shows a similar reduction in GFP to that seen with STAT RNAi. To understand the dynamics of how JAK/STAT activates DRMS we examined how a reduction in STAT might affect regeneration over a 48hr period. Our results show that STAT is required for full enhancer activity and does not preferentially work at a certain point of regeneration. Going forward, we are overexpressing the JAK/STAT pathway elements to determine the necessity and sufficiency in DRMS enhancer activation and whether it can improve regeneration. Furthermore, we are also testing whether this genetic relationship is relevant to other DRMS enhancers that we have identified.

396V Switching On/Off the Hh signalling Pathway Determines Niche Cell Fates of Ovarian Germline Stem Cells Yu-Ting Wang\textsuperscript{1,2}, Chun-Ming Lai\textsuperscript{1}, Hwei-Jan Hsu\textsuperscript{1,2} 1) Academia Sinica, Taipei City, Taiwan; 2) National Defense Medical Center, Taipei City, Taiwan; 3) Louisiana Cancer Research Center, Tulane University, LA, USA

The balance between stem cell renewal and differentiation is critical for tissue homeostasis, and that is largely regulated by stem cell microenvironments, or niches, formed by specific groups of cells. The stem cell maintenance niche recruits and maintains stem cell pool, while the stem cell differentiation niche promotes stem cell differentiation. However, little is known about how these niches are formed. Here, we report that off and on states of Hh signaling determine the fate of maintenance (cap cells) and differentiation niche (escort cells), respectively, of the Drosophila ovarian germ line stem cell (GSC). During development, Hh signaling is activated in intermingled cells (ICs, niche precursors), while it is absent in cap cells but remained in escort cells. Suppressing Hh signaling turnover in ICs, results in cap cell reduction, and induces cap-escort transitional cells that eventually shift toward the escort cell lineage. We also report that Hh signaling in cap cells is silenced via Notch signaling and Cullin3 (Cul3)-HIB-mediated proteasome in a parallel or an upstream-downstream manner. From a small-scale RNAi screening for Cul3 regulators combined with the in silico promoter analysis, we find that the promoter of Ubc10, an E2 ubiquitin-conjugating enzyme, carries two putative Notch responsive elements, and Ubc10 knockdown in ICs phenocopies the above-described phenotypes. Interestingly, Ubc10 with its E3 partner, Ari-1 (Ari-1-Ubc10) is proposed to coordinate with Cul3-HIB complex for protein degradation, an E3-E3 tagging model. Although more studies are needed to pinpoint the involved mechanisms, our studies have added knowledge on the establishment of stem cell niches, and that may be applied to other stem cell systems.

398V The role of Diaphanous in the reactivation of quiescent neural stem cells Kun-Yang Lin\textsuperscript{1}, Wei Yung Ding\textsuperscript{2}, Hongyan Wang\textsuperscript{1,2} 1) Neuroscience & Behavioral Disorders Programme, Duke-NUS Medical School, Singapore, ; 2) Department of Physiology, Yong Loo Lin School of Medicine, National University of Singapore, Singapore

The switch between quiescence and proliferation of stem cells is critical for tissue development and homeostasis. The failure of neural stem cell reactivation is associated with neurodevelopmental disorders, such as microcephaly. Drosophila neural stem cells (NSCs) have emerged as an ideal \textit{in vivo} model for studying quiescence and reactivation. Previously, we discovered that the primary protrusion, a hallmark of quiescent NSCs, is enriched for the actin cytoskeleton. However, the role of the actin cytoskeleton and its regulators have not been established in quiescent NSCs. Here, we report that Diaphanous (Dia), an actin regulator promoting actin polymerization, regulates F-actin polymerization and reactivation of quiescent NSCs. \textit{dia} mutants show delayed NSC reactivation and small brain phenotype. Depletion of \textit{dia} via RNA interference in the quiescent NSCs causes a reactivation delay and the reduction of Filamentous actin. Moreover, overexpression of constitutively active form of Dia leads to the delay of NSC reactivation and larval growth, while overexpressing wild-type form of Dia does not affect NSC reactivation. Together, our results suggest that Dia regulates actin polymerization and is a novel regulator for the reactivation in quiescence NSC.

399V Regulation of nutrient-independent proliferation of the mushroom body neuroblasts (MB NBs) in Drosophila melanogaster Md Ausrafuggaman Nahid\textsuperscript{1}, Conor Sipe\textsuperscript{2}, Sarah Siegrist\textsuperscript{1} 1) University of Virginia; 2) Shepherd University

\textit{Drosophila} neural stem cells (NSC), known as neuroblasts (NB), undergo asymmetric cell division throughout development in order to make the adult fly brain. Dietary nutrients provide essential building blocks necessary for NB growth and proliferation. However, there is a subset of NBs, known as mushroom body neuroblasts (MB NBs), which are able to continue proliferation regardless of extrinsic dietary nutrient availability. My research aims to understand the
molecular mechanism regulating nutrient-independent MB NBs proliferation. Transcriptional co-activator Yorkie (Yki) is well known for its role in maintaining tissue growth and size. However, it remains unclear whether Yki controls the nutrient-independent proliferation of the MB NBs. Upon NB specific Yki knockdown, MB NBs stopped proliferation in response to dietary nutrient withdrawal and resumed proliferation upon re-feeding. Upon expression of a constitutively active form of Yki, all NBs in the fly brain continued cell proliferation independent of dietary nutrient availability. This suggest that Yki regulates NB proliferation decision in response to nutrient availability. Yki requires a DNA binding partner for its function and Sd is the most well-known in Drosophila. I knocked down Sd in a NB specific manner and it did not stop MB NBs from proliferating in a nutrient-independent manner which suggests that Sd might not be the Yki binding partner in the brain. NB specific knockdown and over-expression of Myc resulted in a similar phenotype as Yki. I will conduct two experiments to determine whether there is a genetic interaction between Myc and Yki in the MB NBs. I will over-express Myc in Yki knockdown NBs, and also over-express Yki in Myc knockdown NBs.

400V Consequences of monosomy: How stem cells can lose their female identity and start tumors. Annabelle Suisse, Lara Alzouabi, Allison Bardin Institut Curie, PSL Research University, CNRS UMR 3215, INSERM U934, Stem Cells and Tissue Homeostasis Group, Paris, France.

Aneuploidy, the loss or gain of chromosomes, is the most prevalent genetic hallmark of cancer. However, in a normal context, it is difficult to assess its frequency and its impact as aneuploid cells are often eliminated. It is not known whether chromosome gain or loss happens in healthy stem cells, whether this changes during aging, or how stem cells cope with it. Here, we use the Drosophila intestinal stem cell model to explore these questions.

Our previous data from whole genome sequencing in aged female Drosophila midguts have revealed a loss of an entire X chromosome (monosomy), suggesting that stem cells in healthy tissue can undergo aneuploidy. We have developed a genetic system to study this further, based on the loss-of-heterozygosity (LOH) of an X-linked tumor suppressor gene, whereby loss of the wild-type X chromosome results in neoplastic growth. Using a specific marker, we confirmed that loss of an X chromosome occurs frequently (1 in 800 stem cells).

How do these cells survive and compensate for the sudden loss of an entire chromosome?

We believe that part of the answer lies with buffering by the dosage compensation (DC) pathway, which in male cells deposits H4K16ac on the single X and increases its transcription. We show that DC becomes activated in female monosomic cells, indicating for the first time that a X chromosome counting mechanism still exists in adult cells. We are now testing whether DC is required in monosomic cells for neoplasia formation and effectively buffers the loss of one X.

Our data suggest that another factor promoting the survival and fitness of aneuploid cells could be loss tumor suppressor activity. Indeed, X chromosome monosomy is more frequent when a tumor suppressor is inactivated than in a wild-type context. This suggests that the inactivation of tumor suppressor genes could foster an environment where aneuploid cells can survive and thrive.

Together, our work shows that monosomy happens frequently in healthy stem cells, is an important mechanism of LOH and can be viable, making it a driving force of tumorigenesis.

401V Role of the PIWI protein Aubergine in the regulation of intestinal regeneration Karen Bellec1, Lynsey R Carroll1, Yu Yachuan2, Rippei Hayashi3, Julia Cordero1,2 1) University of Glasgow; 2) CRUK Beatson Institute; 3) Australian National University

Adult stem cells are essential actors in the maintenance of basal homeostasis and damage-induced regeneration in self-renewing tissues. This relies on an accurate regulation of the proliferation and differentiation of stem cells by highly conserved and coordinated signaling pathways. Consistently, disorders in signaling activity may lead to homeostasis disruption inducing age-associated tissues dysfunctions as well as a wide range of cancers.

One of the interests of our laboratory is to identify and understand the molecular mechanisms controlling and adjusting intestinal stem cell (ISCs) behavior to fit the proliferative demands of the gastrointestinal epithelium. For this, we work with the adult Drosophila intestine, an excellent model system that shares functional and structural homologies with its mammalian counterpart. Previous pioneer works shed light on multiple evolutionarily conserved signaling pathways involved in the regulation of ISCs proliferation such as Wnt signaling, JAK/STAT or EGFR signaling. Unexpectedly, the Piwi-interacting RNA pathway (PIWI pathway), known for its role in the repression of transposable elements in the germline, has also been shown to be essential for ISCs maintenance and function through the repression of transposable elements (Sousa-Victor et al., 2017).

Interestingly, we discovered that Aubergine (Aub), another component of the PIWI pathway, is specifically expressed in ISCs and required for their proliferation upon intestinal damage. The abolition of stem cell proliferation upon knockdown of other PIWI pathway components point to a canonical role of Aub in intestinal regeneration. This is consistent with our small RNA sequencing data revealing an increase of piRNA in the intestine upon damage which is abolished upon the knockdown of Aub. In the Drosophila germline, Aub interacts with the translation initiation factor eIF3 to promote target gene translation. In the intestine, this interaction is conserved: eIF3 is expressed in ISCs upon intestinal damage and the loss of Aub is associated with a decrease of eIF3C protein. In addition, the knockdown of eIF3 recapitulates the
phenotypes observed upon the loss of Aub. Altogether, our experiments reveal a new mechanism involved in the regulation of stem cell behavior. Our work bears implications for regenerative medicine, ageing and cancer research as it may lead to the identification of new therapeutic targets to prevent stem cell dysfunction.

402V  WD40 Wuho regulates intestinal stem cell homeostasis for Gut integrity and Longevity Kreeti Kaja1,2,3, Hwei-Jan Hsu1,2,3 1) Molecular and Biological Agricultural Sciences Program, Taiwan International Graduate Program, National Chung Hsing University and Academia Sinica, Taipei; 2) Graduate Institute of Biotechnology, National Chung Hsing University, Taichung; 3) Institute of Cellular and Organismic Biology, Academia Sinica, Taipei, Taiwan

Animal health and longevity require proper maintenance of gut homeostasis, which is largely supported by intestinal stem cells. However, the genetic regulators of stem cell activation and differentiation remain poorly understood. Here, we report that Wuho (Wh, a conserved WD40 protein) maintains the balance between intestinal stem cell (ISC) proliferation and differentiation, which affects the intestinal epithelial barrier and longevity in Drosophila females. Wh is enriched in ISCs; mutation of wh or knockdown in ISCs leads to ISC over-proliferation and aberrant differentiation of ISCs toward absorptive enterocytes. Furthermore, wh mutants exhibit leaky gut and upregulation of antimicrobial peptide transcripts, partially accounting for the shortened lifespan of wh mutants. In addition, ROS levels and JNK signaling are increased in the ISC lineage of the wh mutant gut, possibly due to bacterial infection. Elevated ROS and JNK signaling are known to enhance ISC proliferation and cause gut dysplasia. Thus, Wh appears to intrinsically and extrinsically regulate ISCs in order to promote gut homeostasis. Surprisingly, wh mutant ISCs, generated by FLP/ FRT-induced mitotic recombination, show limited proliferation and differentiation capacities, suggesting a role of Wh in keeping ISCs in a quiescent state within the young gut. We speculate that when Wh is dysfunctional in all ISCs, a compensatory mechanism may be triggered to promote ISC proliferation, resulting in gut dysplasia. Furthermore, Wh expression is decreased in aged ISCs, and wh mutant guts mimic the aged gut phenotype, suggesting that Wh maintains gut physiological aging. We are currently working to identify Wh-interacting partners in the ISC lineage. Nevertheless, our results thus far suggest that Wh may prevent age-associated intestinal dysplasia and inflammation.

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403V  Kinetics of blood cell differentiation during hematopoiesis revealed by quantitative long-term live imaging Kevin Ho1, Rosalyn Carr2, Alexandra Dvoskin1, Guy Tanentzapf1 1) Department of Cellular and Physiological Sciences, University of British Columbia, Vancouver, Canada, V6T 1Z3; 2) School of Biomedical Engineering, University of British Columbia, Vancouver, Canada, V6T 1Z3

Stem cells typically reside in a specialized physical and biochemical environment that facilitates regulation of their behavior. For this reason, stem cells are ideally studied in contexts that maintain this precisely constructed microenvironment while still allowing for live imaging. Here, we describe a long-term organ culture and imaging strategy for hematopoiesis in flies that takes advantage of powerful genetic and transgenic tools available in this system. We find that fly blood progenitors undergo self-renewal, suggesting that they are true stem cells. Using quantitative imaging to simultaneously track markers for stemness and differentiation in progenitors, we identify two types of differentiation that exhibit distinct kinetics. Moreover, we find that infection-induced activation of hematopoiesis occurs through modulation of the kinetics of cell differentiation. Overall, our results show that even subtle shifts in proliferation and differentiation kinetics can have large, aggregate effects to transform stem cells from a quiescent to an activated state.

404V  Enteroendocrine control of intestinal health and disease in Drosophila Andre Medina1,2, Julia Cordero1,2 1) Cancer Research UK Beatson Institute; 2) University of Glasgow

The last few years have witnessed an increased interest in the gastrointestinal tract beyond stem cell research. This is especially true for enteroendocrine (EE) cells, a secretory lineage from intestinal stem cells (ISCs) that plays a key role in sensing local and external stimuli, which are translated into major physiological outputs. As in vertebrates, Drosophila has a diverse EE system scattered along the midgut epithelium and organized into subpopulations of cells expressing different neuroendocrine (NE) peptides. EE cells and NE peptides alterations are recognized features of intestinal damage and gastrointestinal pathologies, such as inflammatory bowel diseases (IBD). However, the functional significance of EE cell disruptions in intestinal physiopathology remains poorly understood. Here, we used adult Drosophila melanogaster as an in vivo model organism to study EE cells function and the functional implications of EE cell disruptions in intestinal damage and inflammation. Previous studies have shown the importance of EE cells to maintain intestinal homeostasis. Depletion of EE cells has a strong impact on intestinal regeneration. On the other hand, we have observed that long-term damage or inflammation favors the increase of EE cell proportion and ectopic expression of NE peptides. To unravel the functional significance of this phenomenon, we performed a genetic screen to identify gut-derived NE peptides that impact ISC proliferation. Our results suggest a novel role for the EE peptide DH31 in this context. DH31 expression in the adult midgut is upregulated upon intestinal damage by feeding with the bacteria Pseudomonas entomophila or chemical treatment with ROS, resulting in previously unrecognized signaling
Epithelia are frequently exposed to extracellular insults and damage due to their exposed position. The restoration of homeostasis upon damage relies on regulated activation of complex stress-responsive signaling pathways to coordinate cellular responses such as apoptosis, G2 stalling, proliferation and survival. Consequently, deviations from these coordinated responses, due to prolonged or potent damage, lie at the center of chronic wound pathologies. Despite their importance, the understanding of how spatial patterns of signaling pathways guide repair behaviors, how the extent of damage is matched by a response of appropriate duration and strength, and how precisely these processes become deviated within damaged tissues remains elusive. Using TNF-α/Eiger-mediated damage to the wing imaginal disc of Drosophila, we observe a stratification of cellular responses coinciding with the spatial patterns of the JNK and JAK/STAT pathways. We find a distinct cell-autonomous repression of JAK/STAT by JNK, and demonstrate through mathematical modeling the importance of such repression to generate our observed spatial patterns. Importantly, JNK-dependent G2 stalling is overridden by ectopic expression of JAK/STAT, leading to increased apoptosis in the tissue. This suggests that the repression of JAK/STAT by JNK is essential to stratify distinct cellular repair behaviors at the wound site. Understanding how perturbations to these patterns impair regenerative responses would be a step towards creating new perspectives into previously uncharacterized links between important signaling pathways and their underlying repair behaviors - which fundamentally influence wound healing pathologies.

Regeneration is the ability to rebuild a body part that has been damaged or amputated. Drosophila imaginal discs are able to undergo wound healing and regenerative growth after injury or genetic ablation. Reactive oxygen species (ROS) act as early signals that are sensed by the MAP3 kinase Ask1, which in turn activates downstream by phosphorylation the MAP kinases p38 and JNK. Apoptosis can occur as the result of sustained or high activation of these kinases, whereas short or low activation can promote regeneration. In search for the conditions that cells require to activate Ask1-dependent regeneration program we find that PI3K/Akt signaling is necessary for Ask1 to activate p38, but not JNK. In addition, nutrient restriction or mutations that target Ser83 of the Drosophila Ask1 protein, a PI3K/Akt-sensitive residue, impairs regeneration. However, these effects can be reversed by the ectopic activation of p38, but not of JNK. Our results demonstrate that the phosphorylation of p38 during regeneration is nutrient sensitive, and that Ask1 controls the activation of p38 through Ser83. This mechanism is important for discriminating between p38 and JNK signaling pathways in the cells involved in tissue repair and regenerative growth.

Regeneration requires surviving cells to mount a response that promotes localized proliferation and repatterning to replace lost and damaged tissue. In Drosophila, the larval imaginal discs regenerate through the formation of a blastema, a zone of localized cell proliferation and increased cellular plasticity. Many important processes during regeneration occur in small subpopulations of cells, the study of which has been revolutionized by single-cell technologies. By profiling the transcriptomes of thousands of individual cells from developing and regenerating imaginal discs, we have identified regeneration-specific transcriptional programs and unique cellular states, including two distinct cell populations within the blastema. These regeneration-specific cell states are characterized by the upregulation of a myriad of genes encoding secreted proteins that establish the pro-regenerative microenvironment. The transcription factor Ets21C is specifically expressed during regeneration in this regenerative secretory zone, and we have demonstrated that Ets21C controls the expression of multiple regeneration-promoting genes, including Mmp1, Ilp8, upd3, and asperous. While eliminating Ets21C function has no discernible effect on development, it severely compromises regeneration. Regenerating tissues in Ets21C−/− mutants fail to maintain a less differentiated blastema and to pause tissue-wide transcriptional changes. As a result, regenerative growth terminates prematurely. Thus, the Ets21C-controlled transcriptional program within the blastema cells is required to effectively coordinate a regenerative response. We also find that this Ets21C-dependent gene regulatory network is activated in small populations of blastema-like cells in tumorous discs, suggesting that pro-regenerative mechanisms can be co-opted by tumors to promote aberrant growth. Our findings highlight unappreciated heterogeneity within the imaginal disc blastema, reveal a critical regenerative gene regulatory network orchestrated by Ets21C, and suggest that this gene regulatory network might function in subpopulations of cells to organize both regenerative and tumorous growth.
During spermatogenesis, many animals replace histones with small positively charged proteins that can tightly package DNA, known as sperm nuclear binding proteins (SNBPs). Unlike highly conserved histones, Drosophila SNBPs recently acquired essential functions for fertility and expanded lineage-specific functional paralogs via duplications. To study how SNBPs acquired sperm-specific function in Drosophila, we focus on two young SNBPs, tHMG1 and tHMG2. tHMG1/2 has originated in the Sophophora subgenus from the duplication of a ubiquitous transcription factor (HmgD or HmgZ) with a high mobility group box (HMGB), and is further tandemly duplicated in D. melanogaster. The roles of tHMG1 and tHMG2 during spermatogenesis are unknown, but they appear to have non-redundant functions. Here we show that tHMG1 is under positive selection using population genetic analyses and its knockdowns show a reduction in progeny. However, we did not detect the same signature of positive selection and phenotypes in tHMG2. Consistent with our data, previous studies also show tHMG1 and tHMG2 have different expression and cytological patterns in sperm. We hypothesized that the highly diverged C-terminus, instead of the conserved HMGB domain, between tHMG1 and tHMG2 determine the functional differences between these two recently diverged duplicates. We are generating CRISPR knockouts and transgenic flies of tHMG-1 and tHMG2 to understand their biological function. Our study will highlight how new SNBPs acquired function during sperm development and further shed light on the evolutionary forces shaping SNBPs evolution.

Phosducin-like protein 3 (PhLP3) has been shown to possess redox-activity, and it is believed to function as a co-chaperone in the folding of certain cytoskeletal proteins. PhLP3 is very well conserved across the animal kingdom from humans to Drosophila melanogaster. PhLP3’s homolog in Drosophila melanogaster is the uncharacterized CG4511. We find that CG4511 plays a role in the regulation of spermatogenesis in Drosophila melanogaster. A P-element inserted in the 5’ UTR of CG4511 leads to a decrease in its expression, infertility in males homozygous for the P-element and testes that fail to produce mature sperm. Further examination of these CG4511 mutant testes reveals that actin-based individualization cones are absent, as are the needle-like nuclei indicative of mature sperm. Differential interference contrast microscopy and phase contrast microscopy reveal defects in spermatid maturation. Additionally, seminal vesicle size in these CG4511 mutants is severely reduced. Excision of the P-element restores male fertility, spermatogenesis, and seminal vesicle size. Our results suggest the importance of CG4511 in the regulation of spermatogenesis. Given its hypothesized role as a co-chaperone for the folding of cytoskeletal proteins, CG4511 may be required to promote the folding of cytoskeletal proteins needed for spermatogenesis. The absence of actin individualization cones in males homozygous for the CG4511 P-element insert supports this hypothesis. We are currently exploring the hypothesis that CG4511 also regulates microtubule dynamics during spermatogenesis.

Nucleoside transporters are essential mediators for nutrient transmission and energy metabolism. Although there have been intensive studies on nucleoside transporters and their biochemical function, no study has investigated the physiological role of concentrative nucleoside transporters (cnts). In this study, we use Drosophila melanogaster to explore the role of cnt1. We generated a mutation in the cnt1 gene. Our results show partial male sterility. Electron microscopy revealed abnormalities in the sperm tail and mitochondrial defects. In addition, we observed high ATP levels in the male testes. We therefore hypothesise that these phenotypes result in altered sperm motility. These results support the hypothesis that energy balance is required for full maturation of spermatids.

Drosophila gonads are highly sexually dimorphic with somatic and germ cell components. The karyotype (XX and XY) of somatic and germ cells must match in addition to the germ cell requiring the non-autonomous signals from soma for the production of functional gametes. Y chromosome ~40mb, is mostly heterochromatin contains ~16 known protein coding genes. The Y linked fertility factors are known to transcribe in primary spermatocytes in a specific spatiotemporal manner for ~90 hours during the spermatogenesis that is important for male fertility. Nevertheless, the Y expression if modulated by non-autonomous cues from soma has never been addressed, which is the purpose of this study. We adapted transcriptomics and generated RNA-seq libraries from gonads of XY larvae (L3) as control and from XY larvae with ectopically expressing transformer (tra) that will allow us to investigate the Y expression when the somatic sex cue is reversed. The transcriptome analysis revealed that none of the Y linked genes that are expressed in XY gonads (12 protein coding genes, 4 pseudogenes with Transcript Per Million value >1) are expressed when the somatic sex
is reversed with the exception of flagrante delicto Y (FDY) that has an autosomal paralog. Our cytological capture of Y transcription (male fertility factor kl3 (kl-3), first exon transcripts detection by in situ) supported our transcriptome analysis as kl-3 is transcribed in XY spermatocytes evidently whereas it was not in XY germ cells when the somatic sex cue is reversed. In case of XXY karyotype, kl-3 is inactive in germ cells possibly because of female cues coming from the soma. In that case it will be interesting to see if reversing the somatic sex in XXY soma leads to Y transcription that we are currently reviewing.

Our results infer that the non-autonomous male cues from soma (tro absence) are necessary for the most Y expression in XY germ cells. The male somatic cues might initiate a special transcriptional program in the germ cells that regulates Y activation which is inhibited when the somatic sex is reversed. Although, it is possible that the male cue from soma is necessary for XY germ cells to progress into the primary spermatocyte stage and proper spermatocyte specific transcription and/or parallel transcriptional program that in turn initiate Y transcription, that we are excited to investigate in near future.

412C Exploring The Sperm Head-Tail Connection Apparatus Kathleen Mulhern, Brian Galletta, Nasser Rusan Nation Heart, Lung and Blood Institute, NIH, Bethesda, MD

The Head-Tail Connection Apparatus (HTCA) establishes and maintains the connection between the sperm’s nucleus and elongating axoneme. Failure of the HTCA results in decapitation which can lead to male infertility. The structure of the HTCA has been investigated through electron microscopy but only a few proteins have been implicated in establishing and/or maintaining this connection. It is highly likely that there are unknown factors contributing to the integrity of this connection as neither direct nor indirect interactions between the known factors have been elucidated. In order to identify unknown factors, we are conducting a GFP localization screen of proteins in the Drosophila Sperm Proteome (Wasbrough et al 2010). Of the 1108 proteins in the sperm proteome, there are 167 proteins with endogenously-tagged GFP stocks available through Bloomington Drosophila Stock Center. We started this screen by looking at 30 of these 167 available stocks. In this initial selection, we prioritized unnamed proteins and proteins with known cytoskeletal binding activity. So far, we have identified 2 proteins with localizations at or near the HTCA and are following up on both by looking at knockdown phenotypes. We have also identified several proteins which localize to other structures during sperm development. This initial success has encouraged us examine other GFP-tagged lines. While the primary goal of this project is to identify factors involved in the HTCA, our initial data also indicates that this screen will uncover factors involved in other aspects of Drosophila sperm development.

413A The N-end rule Pericentrin degradation is required for centrosome assembly and function in Drosophila spermatogenesis Ramya Varadarajan, Brian Galletta, Carey Fagerstrom, Karen Plevock, Nasser Rusan National Heart Lung and Blood Institutes, NIH

Centrosomes are the major microtubule organizing centers (MTOC) that are composed of centrioles and the pericentriolar material (PCM). Centrosomes are essential part of diverse cellular processes that require precise regulation of their protein levels. One protein whose levels must be regulated is Pericentrin – PCNT in humans and PLP in Drosophila. Increased PCNT expression and its protein accumulation are linked to many clinical conditions especially in patients diagnosed with cancer, mental disorders, and ciliopathies reflecting the requirement for regulating PCNT level. However, the mechanisms by which the PCNT is regulated remain less explored. In this study, we took advantage of Drosophila spermatogenesis to characterize the mechanistic of PCNT/PLP degradation. Our previous study by Galletta, 2020 demonstrated that PLP levels are sharply downregulated during early spermatogenesis and this regulation is essential to spatially position PLP in the meiotic centrioles. We therefore first performed a structure-function analysis to identify the functional domain required for regulating PLP level and localization. We found that the N-terminal region of PLP is essential to regulate its protein level and depletion of these regions led to PLP stabilization, which in turn mislocalized PLP on the centrioles. As a consequent, the PCM was also mispositioned leading to defects in spermatids, which ultimately compromised the sperm function. Our biochemical and proteomic analysis revealed that the N terminal region of PLP harbors degradation signals and engages interactions with multiple proteosomal components that hinted a precise degradation process to act via N terminal region. To identify the mechanisms, we performed a candidate RNAi screen composed of proteasome components. We found Rad6, UbcD1 and its related UBR box family E3 ligases, Poe (UBR4) and Hyd (UBR5) to promote PLP degradation in spermatocytes and therefore regulate its spatial localization on the centrioles. The identified candidates are known regulators of the N-end rule degradation pathway, in which their substrates are degraded based on the characteristics of the N-termini amino acids that require single or sequential enzymatic mortification to form N-degron. By characterizing the N terminal amino acids, we found that PLP is a potential substrate of the N-end rule pathway. Collectively, our study identified N-end rule pathway to regulate PLP levels for proper centrosome assembly and may also be relevant for PCNT homeostasis in human as well.

414B A mutation in the gene for kinetochore protein Spc25 disrupts both homolog and sister chromatid connections in male meiosis and causes very high levels of meiosis I nondisjunction Elsie Adams, Bruce McKee University of Tennessee
Genetic and molecular analyses of ethyl-methanesulfonate (EMS) generated mutants has provided a solid forward genetics approach for discovering novel gene products and their roles in various mechanisms over many decades. Cytological analysis of the 3rd chromosome EMS mutant 122-044 strongly suggests a gene mutated plays role in mediating accurate connections, alignment and segregation patterns of chromosomes in both Drosophila male meiosis I and meiosis II. Fluorescence In Situ Hybridization (FISH) analysis with 3rd chromosome specific DNA probe Dodeca reveals 35% of mutant prometaphase I spermatocytes show loss of proper homolog conjunction (n=37), and 83% of mutant anaphase I divisions exhibit homolog nondisjunction of 3rd chromosome autosomes (n=30). Additionally, 20% of metaphase II mutant spermatocytes display premature sister chromatid separation (n=39). Efforts to map this mutated gene led gene candidates within identified a proximal euchromatic region on chromosome arm 3R. Sequencing of genomic DNA from one candidate, the kinetochore gene spc25, from 122-044 mutants revealed a potential 5’ splice site mutation predicted to generate a truncated Spc25 protein. The combination of our observed meiotic phenotypes appears to be novel in that proper chromosome segregation during meiosis I and II of male Drosophila has been understood to rely on two distinct mechanisms - homolog conjunction and sister chromatid cohesion, respectively. However, our understanding of the distinct roles of kinetochore protein complexes during Drosophila male meiosis is incomplete. Our study aims to better understand how specifically spc25 is involved in Drosophila male meiotic chromosome segregation as we complete gene specific complementation analyses, but also determine if some phenotypes may be attributed to mutations of additional genes.

415C Robustness of the canonical mitochondrial fusion machinery promotes Nebenkern formation in Drosophila spermatids Alina Kolpakova, Shmuel Pietrokovski, Eli Arama Weizmann Institute of Science, Rehovot, 76100

Mitochondria, the bioenergetics powerhouses and biosynthetic centers of the cell, are also implicated in many important cellular processes, such as cell death, autophagy, aging, and regulation of immune response and inflammation. Although stereotypically drawn as static organelles, mitochondria are in fact constantly changing shape and subcellular distribution according to function, energy and metabolic demands of the cell. Mitochondrial shapes usually range from small spheres and short tubules to elongated tubules and reticular networks, and these changes are mainly controlled by the balance between two opposing mechanisms of membrane dynamics, fusion and fission (fragmentation). Proper membrane dynamics is essential for maintenance and function of the mitochondria, whereas abrogation of this balance can lead to common diseases, including several neurodegenerative diseases and cancer. The first gene involved in mitochondrial dynamics, fuzzy onions (fzo), was discovered by Hales and Fuller in 1997 as a mediator of the fusion of the entire Drosophila spermatid mitochondria into a giant sphere called Nebenkern. Consequently, orthologs of Fzo, termed mitofusins, have been discovered in organisms from yeast to human, belonging to the dynamin-related protein superfamily of large GTPases. Mitofusins are expressed on the outer mitochondrial membrane (OMM), tethering together adjacent mitochondria by promoting mitochondrial docking through their auto-oligomerization in trans.

In Drosophila spermatids, individual mitochondria aggregate near the newly formed haploid nucleus and subsequently coalesce and fuse into a Nebenkern. The Nebenkern is composed of two giant mitochondria wrapped around each other and arranged in an onion-like structure of layers upon layers of mitochondrial membranes. Although detailed ultrastructural description of Nebenkern formation was already reported five decades ago, the molecular mechanisms underlying the formation of this extraordinary organelle remains largely obscure. Furthermore, it remained unknown whether Fzo has been evolved to uniquely promote fusion of the mitochondria into a giant sphere rather than to reticular network, as well as to what extent the canonical fusion machinery might be involved in Nebenkern formation. Here, I will present our recent studies aiming to address these and related questions, presenting our ongoing unpublished work on the molecular and anatomical mechanisms underlying Nebenkern formation.

416A Regulation of cycB translation by a four-protein complex in Drosophila spermatocytes Catherine Baker, Margaret Fuller Stanford Univ Sch Medicine

The Drosophila male germline contains both mitotic cells (spermatogonia) and meiotic cells (spermatocytes), and the regulation of cell division in these two cell types is dramatically different. Spermatogonia divide regularly and efficiently; spermatocytes, in contrast, undergo a meiotic G2 prophase that lasts 3.5 days, and the concurrent delay of the meiotic divisions is mediated by fine-tuned control of the temporal expression of core cell cycle components. One such cell cycle factor is Cyclin B (CycB). CycB protein expression is high in mitotic spermatogonia, and then low in immature spermatocytes. CycB protein levels spike again just before spermatocytes enter the meiotic divisions. Published work from our lab has shown that the RNA-binding protein Rbp4 and its co-factor Fest repress cycB translation, mediated by sequences in the 130nt cycB spermatocyte 3’UTR. Subsequent work has revealed that Fest acts as a scaffold protein, binding Lutin (CG1690) and Syp as well as Rbp4. Lut, like Rbp4, is required for repressing cycB translation, although the premature expression of CycB protein in a lut mutant appears to begin later than it does in an rbp4 mutant. We confirmed this using the heat-shock time-course developed in the lab, where bam mutant spermatogonia are subjected to a pulse of wild-type Bam protein under the control of a heat-shock promoter, then differentiate into...
spermatocytes and later stages in synchrony. Preliminary data indicate that CycB protein is high by 54 hours post-heat-shock (PHS) in rbp4, by 92h PHS in lut, and by 104h (but not 100h) PHS in wild-type. This result raises the possibility that Rbp4 and Lut, while both belonging to the same complex, could be repressing cycB translation via two distinct mechanisms, perhaps at sequential steps. In contrast, we have found that Syp is required for CycB accumulation in mature spermatocytes. Syp, like Rbp4, binds the 130nt cycB spermatocyte 3'UTR. Curiously, Syp binds to Fest and can co-precipitate with both Rbp4 and Lut in the presence but not absence of Fest. Spermatocytes in the rbp4 syp double mutant show a transient, early expression of CycB, whereas CycB is never detectable in lut syp spermatocytes, suggesting that Syp may act by countering Rbp4 function (but not Lut function). The dynamic off-to-on effect on cycB translation is not dictated by any changes in the core interactions within the complex between 72h PHS and 104h PHS.

417B Cellular and molecular basis of transcriptional regulation during spermatogenesis in Drosophila Saurabh Chaudhary, Sabrina Williams, Shrinivas Dighe, Katia Jindrich, Helen White-Cooper School of Biosciences, Cardiff University, Cardiff, UK

Spermatogenesis provides an excellent model system to study many biological processes, including co- and post-transcriptional gene regulation. The adult Drosophila male germline is highly transcriptionally active during the primary spermatocyte stage. The transcriptional complexity in Drosophila spermatogenesis is well known, however, the underpinning molecular mechanism regulating the expression of more than 1000 genes, specifically in testes in Drosophila remains largely undetermined. Using predominantly D. melanogaster as a model, we are employing various state-of-art techniques in the field to understand the cellular and molecular basis of testes specific gene regulation. Testis Meiotic Arrest Complex (tMAC) is required for the activation of transcription of a large number of genes in spermatocytes. This complex comprises several proteins with DNA binding activity and several additional non-DNA binding proteins. We are dissecting how this complex interacts with target promoters by identifying in vitro binding motif for each of the DNA-binding domain-containing proteins within tMAC using high throughput SELEX sequencing. For in vivo identification of binding sites of each tMAC subunit, we are generating GFP-tagged lines and using high-resolution Chip-Exo, RNA-Seq, and Chip-qPCR in Drosophila testes. Testis-specific transcription typically depends on short DNA regions immediately flanking the transcription start sites (TSS). We have generated a set of promoter-reporter constructs and are investigating how these short promoter regions interact with tMAC, and other transcriptional regulatory complexes, to enable gene expression. For some genes, a small number of mutations are sufficient to convert a non-functional promoter into a functional promoter. By integrating the in vivo data with the in vitro binding site motif information, we will determine which DNA-binding proteins contribute to the binding of tMAC to any specific promoter sequence. This study will further deepen our understanding of how testes specific gene regulation achieves at co- and post-transcriptional levels in Drosophila.

418C Characterization of test specific sugar transport and glycolysis genes in Drosophila melanogaster Mark Hiller, Julia Gazzola, Elizabeth Hughes, Rylee McDonell, Katie Shannon, Emily Pochet, Katlyn Heneghan, Gianna Graziano Goucher College

Sugar is required to provide energy for cellular metabolism and development. The genome of Drosophila melanogaster contains twenty-five genes that are annotated as SLC2 type sugar transporters, and five of the sugar transporters appear to be expressed only in the testis. Lactate transport into male germ cells could also be used as a source of ATP, and one gene, CG12866, encodes a testis specific Monocarboxylate transporter. Likewise, there are several genes that function in glycolysis that have testis specific homologs or testis specific splice forms. Each of these genes could function during the generation the cellular energy necessary for spermatogenesis or be required for sperm function during fertilization. Mutations in the sut3, sut4, and CG14605 putative transporters are fertile. To assess the role of glycolysis and lactate transport, we are using RNAi in the germline and cyst cells of the testis to knockdown function of the lactate transporter and glycolysis genes.

419A Rethinking cyst formation during Drosophila spermatogenesis Rocky Diegmiller1, Tomer Stern1, Yukiko Yamashita2,3, Stanislav Shvartsman1,4 1) Princeton University, Princeton, NJ, USA; 2) Massachusetts Institute of Technology, Cambridge, MA, USA; 3) Whitehead Institute, Cambridge, MA, USA; 4) Flatiron Institute, New York, NY, USA

The process of rapidly and properly producing germ cells is essential for life. In Drosophila melanogaster females, germline clusters of 16 cells are formed as the result of four maximally-branched, synchronous divisions from a single germline stem cell. This same pattern of divisions has also been thought to govern Drosophila spermatogenesis, where cells in the cyst additionally undergo a round of meiosis to produce 64 identical spermatids. Here, using 3D reconstructions of structural elements in developing male germline clusters, we provide evidence to suggest that maximal branching can be lost at the fourth division. We additionally show that the fusome, which allows for intercellular communication and transport within the cyst, is fragmented more frequently in the male germline, suggesting it plays a less vital role than in oogenesis. Overall, this work establishes an important distinction between male and female germline development and introduces future areas of research to investigate the mechanisms governing gametogenesis across species.
Using FIB-SEM to create a 3D model of early oogenesis

The fruit fly serves as an excellent model for studying oogenesis, or female gamete formation. Each egg develops from an egg chamber, which progresses through the stages of oogenesis as part of a developmental array known as an ovariole. At the anterior of each ovariole lies the germarium, which contains the germline and somatic stem cells which will divide to give rise to each newly formed egg chamber. Much has already been learned about oogenesis using a combination of fluorescence and electron microscopy to study specific structures and proteins; however, a complete picture of the organization of germline and somatic cells and their intracellular structures and organelles is lacking. To learn more about the early stages of oogenesis, we are rendering a large data set collected using Focused Ion Beam Scanning Electron Microscopy (FIB-SEM) to generate a complete 3D model of the germarium and an early-stage egg chamber. Our lab is primarily interested in the structure of the germline ring canals, or intercellular bridges, which allow nurse cells to transfer their cytoplasmic contents into the growing oocyte. Electron microscopy has previously revealed the presence of extensive membrane protrusions, or microvilli, surrounding the germline ring canals. This microvilli meshwork is thought to maintain the anchoring and stability of the ring canals during egg chamber growth, yet the stage of oogenesis when these protrusions first appear and their spatial distribution within the germline is not known. Therefore, we hope to characterize changes in the size and structure of the germline ring canals and the organization of germ cell membranes during early oogenesis. Once completed, we envision that this dataset can become a valuable resource that can be further mined by other researchers in the field.

The neurodegeneration gene iPLA2-VIA is required for mitochondrial maintenance in the Drosophila melanogaster female germline, with autonomous and non-autonomous components

Neurodegenerative disease (ND) is a major health issue worldwide, yet the underlying mechanisms are still ill-defined. Loss of function mutations in the gene PLA2G6, encoding the group 6A calcium-independent phospholipase A₂, iPLA2-VIA, are associated with severe ND in humans, including autosomal recessive dystonia-parkinsonism. The orthologous Drosophila melanogaster gene encodes a highly similar protein to the human gene product, and loss of function mutations in flies lead to neuron loss and age-dependent locomotor defects. We have discovered a novel role for iPLA2-VIA in the Drosophila female germline. In wild-type flies, iPLA2-VIA is highly expressed in adult germ cells, and a transgenic HA-tagged iPLA2-VIA protein (PB isoform) localizes to the mitochondria of nurse cells. Furthermore, a null mutation in iPLA2-VIA, generated in our lab, causes age-related mitochondrial aggregation and loss of mitochondrial membrane potential in the female germline, symptoms of mitochondrial damage, with eventual germ cell apoptosis. The mitochondrial localization of the protein and the cellular effects in the mutant mirror the behavior of this gene in neurons, suggesting possible common underlying mechanisms of action in these two cell types. Because iPLA2-VIA is expressed broadly in somatic tissues as well as in germ cells, we explored the tissue autonomy of the germline mitochondrial defects using our technique to quantitatively analyze nurse cell mitochondrial aggregation. Surprisingly, germline iPLA2-VIA RNAi knockdown only mildly phenocopied the null mutation, while ubiquitous somatic knockdown led to strong mitochondrial aggregation and death of female germ cells, suggesting a strong non-autonomous component to the germline mitochondrial defects. We currently are knocking down iPLA2-VIA in select somatic tissues, including neurons, muscle, fat body, and ovarian follicle cells, to determine which specific tissues contribute to the germ cell phenotype. In complementary studies, we are conducting tissue-specific rescue experiments using a wild-type cDNA transgene or a catalytically-inactive cDNA transgene, both of which rescue age-dependent locomotor decline. Finally, we have shown that iPLA2-VIA does not interact genetically with pink1, another neurodegeneration gene important for germline mitochondrial integrity, suggesting parallel pathways. We expect our findings to have relevance to the mechanisms underlying PLA2-associated neurodegeneration.

Spargel/dPGC-1 is a closer ancestor to mammalian PRC-1 with an RRM domain that is functionally essential for oogenesis

The transcriptional coactivators Peroxisome Proliferator Activated Receptor-1 (PGC-1) play a key role in wide range of physiological processes in mammals including mitochondrial biogenesis, oxidative metabolism, and adaptive thermogenesis. There are three mammalian PGC-1 paralogs: PGC-1α, PGC-1β, and PRC-1. Spargel/dPGC-1 is an ancestral PGC-1 homolog in Drosophila that is predominantly expressed in the ovaries during adult life. Ovary-specific knockdown of srl induces complete sterility indicating that Spargel plays an essential role in oogenesis and ovarian growth. Spargel shares many structural features with PGC-1 proteins including the Serine-Arginine rich repeats (RS), RNA recognition motif (RRM) and a nuclear localization signal (NLS). Due to their functional redundancy, attempts to make domain specific deletions in PGC-1 failed to demonstrate the role of RRM in mammalian system. We generated two srl mutant lines using CRISPR/CAS9 system: (1) A 3336 bp deletion was created within the srl beginning from Exon 2 (srl"del"), and (2) RRM domain specific deletion (ΔRRM). Homozygous srl"del" embryos complete the embryonic development but can’t...
proceed beyond 1st instar larval stage as these larvae failed to emerge out of the chorion. This suggests that Spargel function is not limited to oogenesis, but it is also essential for postzygotic development. Mitotic clones of srlΔRRM established it as an amorphic allele which may be the reason why most srlΔRRM egg chambers don’t survive beyond previtellogenic stages. Incidentally, germ line homozygous knockouts of PGC-1α or PGC-β are viable and fertile with no global changes in mitochondrial number or morphology, however a germ line knock-out of the PPRC1 gene in mice results in neonatal lethality. Phylogenetic analysis supports that Spargel is actually a closer ancestor to mammalian PRC-1 as compared to PGC-1α or PGC-β. Homozygous srlΔRRM females are viable but carries very few mature eggs in their ovaries. Further analysis shows the infertility of srlΔRRM females stems from disrupted actin cable and cortical actin loss in nurse cells resulting in dumpless phenotype. Together, this collection of spargel mutants is helping us to address (1) the role of Spargel on ovarian growth and development, and (2) to determine the role of Spargel in postzygotic development.

**423B Exploring the role of Oatp74D, an Ecdysone Importer, in the Drosophila ovary.** Amanda Powell, Elizabeth Ables East Carolina University

Oogenesis is the progression of germ cells through mitotic expansion, differentiation into an oocyte, and successful completion of meiosis. Steroid hormones play critical roles in this process in diverse organisms. In *Drosophila*, the main steroid hormone, ecdysone, facilitates female fertility in part by promoting germline stem cell self-renewal. Foundational studies demonstrated that ecdysone is synthesized primarily in egg chambers during mid-oogenesis; however, more recent data suggests that somatic escort cells, which support germline stem cell differentiation, may also produce ecdysone. Understanding how ecdysone is transported and received in the ovary would help resolve these disparate results. Recent studies by the Yamanaka lab support the hypothesis that ecdysone needs Oatp74D, a membrane transport protein, to import ecdysone into cells, challenging the popular assumption that steroids passively transport through membranes. Oatp74D is well-characterized in the *Drosophila* blood brain barrier but may have roles in steroid hormone uptake in other cells. Here, we explore whether Oatp74D is necessary for ecdysone-mediated processes during oogenesis. We use transgenic and immunofluorescence approaches to examine localization of Oatp74D in the ovary. We then test whether Oatp74D knockdown in escort cells effects oogenesis and fecundity. Our preliminary results suggest that Oatp74D is not necessary in escort cells for maintenance of germline stem cells but may promote germ cell differentiation. This data will further our understanding of how ecdysone signaling regulates optimal oocyte production.

**424C Regulation of Delta-Notch pathway by mitochondrial signaling during drosophila oogenesis.** Yipeng Du, Matthew Sieber UT Southwestern Medical Center

Development in all organisms requires the coordination of cellular metabolism with the proper regulation of signaling pathways. For example, changes in mitochondria metabolism have been associated with cell specification and differentiation in many developmental systems. However, despite this association, very little is known about how mitochondrial metabolism regulates developmental pathways. Our lab utilizes *Drosophila*, mammalian cells, and mice to examine the conserved metabolic mechanisms that drive development and disease progression.

To understand the role for mitochondria in developmental signaling, we conducted a genetic screen of mitochondrial proteins. Using this approach, we discovered that a specific subset of mitochondrial genes, when disrupted, cause defects in development of the *Drosophila* ovary and intestine. In particular, we found that inhibition of prohibitin (*PHB in mammals*) in germ cells caused a block in follicle cell differentiation. Loss of prohibitin in the mitochondria causes: a reduction of membrane potential, decreased ATP production, and increased cellular ROS. These defects in mitochondrial oxidative metabolism cause an increase in follicle cell number, reduced size of follicle cell nuclei, and defects in follicle cell differentiation in *prohibitin* RNAi egg chambers. When we examine the localization of the Notch ligand, Delta, we find that it accumulates in large puncta in *prohibitin* RNAi egg chambers and fails to reach the plasma membrane. Interestingly, lowering ROS by overexpression Catalase and Sod2 genes partially rescue the Delta aggregation developmental phenotypes in ovaries. In addition, we have found that overexpression of Rab11 which is a key protein of recycling vesicles rescue the phenotype of Delta aggregation in *prohibitin* RNAi egg chambers. These data suggest that mitochondrial ROS regulates Delta trafficking in Rab11 dependent manner. Overall, our work provides intriguing new evidence that identifies the regulation of receptor trafficking as a novel mechanism for how mitochondria regulate differentiation. Moreover, this work provides a foundation for future studies of how changes in mitochondria metabolism promotes cancer progression by regulating Notch signaling.

**425A A Cytological F1 RNAi Screen for Defects in Drosophila melanogaster Female Meiosis** William Gilliland, Amanda Bowen, Kelly Conger, Doreen Elrad, Marcin Marciniak, Denny May, Gabrielle Presbitero DePaul University

Forward genetic screens induce mutations, make the mutated chromosomes homozygous, and then homozygotes for the phenotype of interest. When studying female meiosis, the phenotype is usually nondisjunction from chromosome segregation errors. This means that mutant females must be viable and fertile, and any meiotic genes that are lethal or sterile when homozygous cannot be recovered by this approach. Our lab has screened the VALIUM22 collection produced by the Harvard TRIP Project, which contains RNAi constructs targeting genes known to be expressed in the germline in a vector optimized for germline expression. By driving RNAi with GAL4 under control of a germline-specific
promoter (nos or mat-alpha), we can test genes that would be lethal if knocked down in all cells, and by examining unfertilized metaphase-arrested mature oocytes, we can identify defects associated with genes whose knockdown results in sterility. We screened this collection to identify genes that disrupt either of two phenotypes: the ability of meiotic chromosomes to congress to a single mass at the end of prometaphase, and the sequestration of Mps1-GFP to unknown structural filaments in response to hypoxia. After screening the ~1500 lines in the collection, we obtained multiple hits for both phenotypes, identified novel meiotic phenotypes for genes that had been previously characterized in other tissues, and found novel phenotypes for several previously uncharacterized genes.

426B Structural changes in centrosomes correlate with activation of a checkpoint that triggers germline stem cell loss Isabella Perales1, Tingting Duan1, Pamela Geyer1,2 1) University of Iowa; 2) NIH

Homeostasis of Drosophila germline stem cells (GSCs) depends upon the integrity of the nuclear lamina (NL). Indeed, loss of the NL protein emerin blocks germ cell differentiation and causes GSC death due to activation of two DNA damage response kinases ATM- and Rad3-related (ATR or Mei-41) and Checkpoint kinase 2 (Chk2 or Loki). Previous studies suggested that checkpoint activation in emerin mutants results from a thickening and lobulation of the NL, due to insertion of enlarged interphase centrosomes that carry excess pericentriolar material (PCM) to nucleate microtubules. To understand whether NL distortion or centrosome structure is the primary activator of the ATR/Chk2 checkpoint, we used the Gal4-UAS system to overexpress two NL proteins, the inner nuclear membrane protein Kugelkern (Kuk) and the A-type lamin (Lamin C), chosen based on previous findings that their increased accumulation caused NL distortion. Although over-production of both proteins was achieved, only Kuk caused NL deformation in GSCs, demonstrating that Lamin C over-expression has cell type specific effects on NL structure. Whereas Kuk overexpression caused NL distortion, interphase centrosome structure was unaffected, with centrosomes localizing outside of the nuclear envelope and retaining low levels of PCM. Notably upon Kuk overexpression, GSCs were maintained and oogenesis was sustained. Based on these findings, we conclude that NL distortion alone is not sufficient for activation of the ATR/Chk2 checkpoint. Instead, our findings suggest centrosome structural changes might drive checkpoint activation in GSCs.

426C Stonewall promotes germ cell to oocyte transition by promoting heterochromatin maintenance during Drosophila oogenesis Noor Kotb1,2, Ankita Chavan3, Lydia Proskauer4, Elliot Martin7, Madhav Jagannathan3, Prashanth Rangan2 1) Department of Biomedical Sciences/ Wadsworth Center, University of Albany, Albany, NY; 2) Department of Biology/RNA institute, University of Albany, Albany, NY; 3) Department of Biology, Institute of Biochemistry, ETH Zürich, Zurich; 4) Colgate University, Hamilton, NY

During oogenesis, germ cells differentiate into an oocyte, which upon fertilization gives rise to a totipotent zygote. In Drosophila, the transition from germ cell to an oocyte requires SETDB1 dependent heterochromatin formation to silence germ cell differentiation genes after the oocyte specification. These silenced germ cell genes are often interspersed between euchromatic genes that are actively transcribed to promote oocyte fate. As heterochromatin is prone to spreading to euchromatic regions, how such heterochromatic territories remain confined during oogenesis remains poorly understood. Here, we find that the DNA binding protein Stonewall (Stwl), is required to establish a boundary between heterochromatic and euchromatic domains. We find that stwl binds to promoters of the silenced germ cell differentiation genes and promotes the maintenance of heterochromatin at these loci. Loss of stwl results in loss of heterochromatin spreading, dysregulation of transcription and loss of oocyte fate resulting in mid-oogenesis death. Thus, during germ cell to maternal transition, Stwl acts a boundary element to maintain heterochromatin to promote an oocyte fate.

428A fs(1)K741 is a female sterile allele of the gene Sxl and disrupts Sxl splicing Jillian Gomez1,2, Myles Hammond2,3, Stephen Kucera2, Brian Oliver1, Leif Benner1,4 1) National Institute of Diabetes and Digestive and Kidney Diseases, NIH, Bethesda, MD; 2) Department of Biology, University of Tampa, Tampa, FL; 3) Integrative Genetics and Genomics, University of California-Davis, Davis, CA; 4) Department of Biology, Johns Hopkins University, Baltimore, MD

Sxl is the master sex-determination gene and acts via a sex-specific splicing mechanism. The locus has often been discussed as two simple steps in females: the production of an early protein involved in establishing autoregulatory splicing and the production of a late protein maintaining the female-specific splice. We mapped a female sterile allele, fs(1)K741, to the Sxl locus through deficiency mapping and duplication rescue. We used whole genome sequencing to determine fs(1)K741 to be a single point mutation (C>T) in the male-specific exon 3 of Sxl. CRISPR-Cas9 and a ssOligo donor directed to the point mutation in fs(1)K741 were employed to recreate the mutation in wild-type females. This point mutation failed to complement fs(1)K741, confirming it to be the cause of sterility and that fs(1)K741 is an allele of Sxl, thus called SxlK741-1. RT-PCR of SxlK741-1 ovaries showed both male and female-specific Sxl splicing products, while RT-PCR on SxlK741-2 carcasses showed only female-specific Sxl splicing. This indicates that aberrant splicing is restricted to the ovary. We wanted to determine how this mutation in the male exon was disrupting Sxl splicing. Homozygous SxlK741-1 females have a temperature-sensitive sterility and lethality effect. At 18°C, homozygous females are 28% viable and 80%
At 29°C, homozygous females are 37% viable and completely sterile. Two duplications, Dp(1;3)DC489 and Dp(1;3)DC490, rescued viability of Sxl<sup>K741-1</sup> females. However, only Dp(1;3)DC490 rescued fertility. The difference between the two duplications is that Dp(1;3)DC490 contains the full Sxl reading frame while Dp(1;3)DC489 only contains the early Sxl reading frame, possibly indicating that the viability defect in Sxl<sup>K741-1</sup> is due to aberrant early protein function. We created loss-of-function frameshift mutations specific to the early, late, and all proteins denoted as, Sxl<sup>Early</sup>, Sxl<sup>Late</sup>, and Sxl<sup>Full</sup>, respectively. Sxl<sup>Full</sup> failed to complement viability of Sxl<sup>K741-1</sup> at both 18°C and 29°C. Sxl<sup>Early</sup> failed to complement viability at 18°C, but was subviable and subfertile, at 29°C. Sxl<sup>Early</sup> complemented viability and fertility of Sxl<sup>K741-1</sup> at both 18°C and 29°C. These results are somewhat at odds with the duplication results since the duplication only containing the early protein rescued viability. Still, it is clear from these new mutants that Sxl<sup>K741-1</sup> is not deficient in the establishment step of Sxl splicing, and the mutation is likely disrupting the maintenance of Sxl autoregulation. Although Sxl has previously been thought of and described as a simple step-wise autoregulatory cascade initiated by the early protein and maintained by the late protein, Sxl<sup>K741-1</sup> clarifies that the autoregulatory process of Sxl is much more complex. Understanding how this mutation in the male exon of Sxl leads to subviability and sterility will help researchers further understand the complex genetics of Sxl.

**429B** The expression of OVO isoforms throughout *Drosophila* development

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An important component in the continuity of life is sexual reproduction, yet the genetic determinants of germline differentiation to create a sex-specific gamete remain largely uncovered. Exploring the activity of transcription factors essential to these processes give needed insight to this concept. In *Drosophila*, OVO is required for female germ cell viability. ovo is a self-regulating gene that encodes two antagonistic isoforms, OVO-A and OVO-B, with OVO-A acting as a repressor and OVO-B acting as an activator. To reveal the behavior of OVO, we chose to visualize its activation during germline development and oogenesis. A sequence containing a 3x-FLAG-HA tag was placed at 4 locations in the open reading frame of the ovo locus, allowing us to tag different isoforms and visualize its expression and cellular localization. This tag was placed at the N-terminus of the OVO-B transcript (ovo<sup>ovo-Cterm</sup>, tagging both OVO isoforms), the N-terminus of the OVO-A transcript (ovo<sup>ovo-A</sup>, tagging OVO-A), the C-terminus of all transcripts (ovo<sup>ovo-Cterm</sup>, tagging both OVO isoforms), and the N-terminus of an OVO-B transcript in which the promoter of OVO-A was deleted (ovo<sup>ovo-Cterm</sup>, tagging OVO-B). Staining in adult ovaries of these different alleles revealed the presence of nuclear OVO in the gerarium and differentiating germ cells of the adult ovaries. ovo<sup>ovo-Cterm</sup> had a persistent staining pattern from germline stem cells throughout developing egg chambers, while ovo<sup>ovo-Cterm</sup> showed a weaker staining intensity in the germline stem cells but stronger in region 2a of the gerarium and stage 4 egg chambers. ovo<sup>ovo-ΔAP</sup> replicated staining patterns of ovo<sup>ovo-Cterm</sup>. ovo<sup>ovo-A</sup> appears to have a decreased staining intensity than the other alleles but follow the same staining pattern as ovo<sup>ovo-Nterm</sup>. We also looked at the expression of OVO in the adult male germline and were able to detect the presence of OVO in ovo<sup>ovo-Nterm</sup> and ovo<sup>ovo-Cterm</sup> in germ cells near the hub, but staining was lost in differentiating spermatogonia, while ovo<sup>ovo-A</sup> staining was not detectable. Immunoblotting analysis of the alleles show that ovo<sup>ovo-Nterm</sup> and ovo<sup>ovo-ΔAP</sup> have identical banding patterns, but the ovo<sup>ovo-Cterm</sup> had different protein bands, while ovo<sup>ovo-A</sup> protein was not detectable. The localization and banding patterns between ovo<sup>ovo-Cterm</sup> and ovo<sup>ovo-Nterm</sup> suggest that OVO may be regulated at the protein level. We are continuing to determine the expression pattern of these alleles during embryogenesis and larval development in males and females to gain a better insight into ovo expression and regulation.

**430C** Analysis of RNA Helicase Me31B’s Molecular Mechanism in Germline Development by Motif Mutations

Carol Dils, Aidan McCambridge, John Eshak, Brooke Pumnea, Yousif Mukatash, Joseph Jansky, Noor Malik, Ming Gao Indiana University Northwest

*Drosophila* Me31B (DDX6) is a conserved ATP-dependent RNA helicase that plays role in post-transcriptional RNA regulation to ensure the correct spatial and temporal expression of germline mRNAs. This, in turn, is crucial for the proper germline development in many animals. However, Me31B’s role and molecular mechanism are not clear. To study this, we aim to first understand the functions of important domains/motifs of the protein. Therefore, we used CRISPR gene-editing technique and generated *Drosophila* strains mutant for functionally important motifs in Me31B including the two RecA-like domains, N-terminal domain and C-terminal domain, motifs involved in ATPase and helicase activities such as DEAD-box motif and HRIGR motif, motifs that cause human developmental defects if mutated such as QxxR motif, and protein-binding motifs such as FDF-pocket motif and W-pocket motif. Further, we are conducting a series of phenotype characterization experiments on the mutants including quantification of the expressed mutant proteins, confocal microscopic examination of the mutant proteins’ localization in the ovaries, and fertility assays of the mutant strains. Considering that Me31B/DDX6 family proteins are also expressed in certain soma like neurons and other organisms such as yeast, worm, mouse, and human, the results from this study could further help us understand Me31B/DDX6 family proteins’ likely conserved roles in different cell types and species.

**431A** Size Regulation within the Germline of the Developing Egg Chamber.

*Zoe Herdman,* Umayr Shaikh, Lindsay
The relationship between the size of a cell and the size of its intracellular structures must be precisely regulated for proper development and functioning of the mature organism; however, the mechanisms that establish and maintain this relationship are not known. Much of the work on organelle size scaling has focused on individual cells or organelles, with some looking at how size scaling is maintained during development. However, it is not known whether the size of multiple organelles is coordinately regulated, and how cell size and organelle size are regulated within a syncytium or group of connected cells. Because many cell types exist within a syncytium, understanding size scaling in this context is important. We have used the developing fruit fly egg chamber as a model system to study the size relationship between multiple structures within the germline syncytium throughout oogenesis. The developing egg chamber contains a cluster of 16 germ cells connected by intercellular bridges, or ring canals. It has been observed that the size of the germline ring canals exhibits spatial variation with the smallest ring canals at the anterior of the egg chamber and the largest ring canals at the posterior near the oocyte. Recent work has also confirmed that nurse cells and their nuclei show a similar size distribution within the germline. Therefore, it is interesting to consider whether the size of these structures may be coordinately regulated, and whether altering the size of one structure (either throughout the germline or clonally) would impact the size of the other structure. We have used a combination of RNAi (Pendulin, Dacapo, or Ctf4), overexpression (Pendulin) and mosaic analysis (myc, arpC1, dock) to try to alter either nuclear size or ring canal size in some or all cells within the germline. We used Fiji to measure the diameter of the ring canals, the size of the nurse cell nuclei (which has been used as a proxy for cell size), and the overall dimensions of the egg chamber. Preliminary data suggest that altering nuclear size within the entire germline does impact ring canal size, and that in mosaic tissue, there is typically a correlation between cell size, nuclear size, and ring canal size. In the future, we hope to further explore whether the size of these structures is coordinately regulated, and if so, to identify the underlying mechanisms that are involved.

432B Nuclear and ring canal growth in the germline of the developing egg chamber Kathleen Sherlock, Julia Wilson, Umayr Shaikh, Daniel Adan, Lindsay Lewellyn Butler University

The size of intracellular structures must be tightly controlled in order to maintain normal cell function. This is especially important when cells are part of growing tissues or organs. However, despite its importance, few studies have analyzed how the size of organelles or other structures changes during development. The developing egg chamber provides a unique model in which to study the relative growth rates of multiple structures. The egg chamber is composed of a cluster of germ cells (an oocyte and 15 supporting nurse cells) surrounded by a layer of somatic epithelial cells. In addition to the growth of the overall egg chamber, it is also known that multiple structures within the egg chamber grow as well. For example, the nurse cells, their nuclei, and nucleoli increase in size as the egg chamber develops, and the intercellular bridges, or ring canals, that connect the germ cells also significantly increase in diameter. Here, we are able to take advantage of differences in egg size both within D. melanogaster and between Drosophila species (D. pseudoobscura, D. melanogaster, D. santomea, D. yakuba, and D. virilis) to test whether the relative rates of growth of the germline ring canals and nurse cell nuclei are consistent in developing egg chambers of different sizes. To determine whether these growth rates are impacted by the location of the structure within the egg chamber, we specifically focus on two sets of nurse cells and their associated structures — the nurse cells directly connected to the oocyte and those located at the anterior of the egg chamber. In addition, we test whether altering oocyte growth impacts the growth of the germline ring canals or nurse cell nuclei. In the future, we hope to determine how these scaling relationships are maintained during dramatic tissue growth and to test whether the same scaling relationships are observed in more distantly related species.

433C Mob family proteins and Tricornered kinase are required to form dorsal appendages of the Drosophila eggshell Keala Watson1, Juan Carlos Duhart2, Laurel Raftery1 1) School of Life Sciences, University of Nevada, Las Vegas, Las Vegas, NV; 2) Current Address: Dept. of Neuroscience, Jefferson University, Philadelphia, PA

Formation of complex tissue structures involves multiple signaling pathways to coordinate cells. In Drosophila, epithelial cells form tubular structures to shape the dorsal appendages on the eggshells. We found that components of a Hippo-like intracellular signaling pathway are involved in shaping dorsal appendages, in particular, the core NDR kinase tricornered (Trc) and several Mob proteins. In humans, mice and flies, Mob proteins bind to NDR kinases and increase their activity. There are 4 Mob genes in Drosophila: mats (encoding Mob1), mob2, mob3, and mob4. We made a mob2 knockout allele by CRISPR/Cas9-aided homologous recombination, and found that homozygous females lay eggs with variably shorter dorsal appendages than control females. Knockdown of Tricornered kinase throughout the follicle cells was predominantly associated with defects in tube formation, with a minority of eggshells showing short dorsal appendages. These data suggest that another Mob protein associates with Tricornered for early dorsal appendage tube formation. We are testing the phenotypes of follicle cell knockdown for each of the other three Mob proteins, to identify a candidate Mob partner for Tricornered function in early dorsal appendage tube formation.

434A Physiological and functional implications of differentially enriched transcripts on eRpl22-family polysomes Caroline Pritchard, Vassie Ware Lehigh University, Bethlehem, PA
The *Drosophila melanogaster* eRPL22 ribosomal protein family contains two structurally divergent & developmentally essential paralogues: eRPL22 and eRPL22-like - the latter exhibits tissue-specific expression across development; the former is ubiquitously expressed. Multi-tissue co-localization comparison of eRPL22-like and core ribosomal components indicates eRPL22-like may have functional roles both within the ribosome itself and apart from ribosomal processes.

Sequencing of RNAs enriched on eRPL22 and eRPL22-like polysomes in adult testes revealed differential enrichment of mRNAs suggesting that paralogue-specific “specialized ribosomes” translate specific mRNAs. Functional enrichment analysis guided investigation into specific tissues by physiology (where to look) and function (what processes, pathways, and programs function differently in Rp mutants). KEGG pathway analysis revealed transcripts differentially enriched on eRPL22-like polysomes were uniquely implicated in pathways not canonically associated with ribosomal functions, including endocytosis, autophagy, and mTOR signaling. Some pathways were over-represented in both polysome types, but these were derived from unique transcripts.

Transcripts of genes functionally implicated in human disease, given by Human Phenotype Ontology (HPO) term association, were over-represented on eRPL22-like polysomes (961:496 terms). Grouping each term into broad categories revealed HPO terms within the genitourinary, musculoskeletal, and nervous systems, and the brain, eye, head and jaw regions, were most functionally enriched on eRPL22-like polysomes (181:8 terms). We have previously shown differential expression of eRPL22-like protein and specific knock-down phenotypes in several analogous regions in the fly.

Conditional knock-out of eRPL22-like resulted in many morphological defects within the ovary, including disruption of the germline stem cell niche, ectopic rounded follicular epithelium cells, oocytes with dual (bifurcated) nuclei, double-anteriorized eggs, and specific spatiotemporal patterns of cell death & oogenesis arrest. This constellation of phenotypes suggests eRPL22-like has a role in ensuring proper cell polarity in early development and the ovary is a viable system to investigate the cellular and molecular basis of these defects.

Taken together, these data broaden the context for essential roles of eRPL22-like across multiple developmental processes.

**435B Identification of E2 ubiquitin-conjugating enzymes required in Drosophila male meiosis** Andrea Binder, John Tomkiel Dean University of North Carolina at Greensboro

Ubiquitination is a post-translational modification in which a small protein called Ubiquitin is covalently attached to Lysine residues of target proteins to specify their degradation, alter their activity or localization. In a multistep process, an E1 ubiquitin ligase activates Ubiquitin and transfers it to an E2 ligase. The activated E2 then interacts with an E3 ligase which specifies the target. In *D. melanogaster* there is a single E1 ligase (Uba1), 29 predicted E2 ligases and 156 predicted E3 ligases, allowing for a multitude of potential ubiquitin ligase complexes. This suggests that this complexity is needed for temporal or tissue-specific ubiquitination, however relatively few complexes and their targets have been defined. Here we sought to identify which E2 ligases are required in the male germ line. We used four *GAL4* drivers (*bam*, *C135*, *nanos*, *T110*) in combination with UAS-RNAi constructs to knock down expression of all 29 E2 ligases and Uba1 in the male germ line. Test males were assayed for fertility, were examined cytologically for visible defects in spermatogenesis and monitored genetically for fourth chromosome missegregation. Knockdowns of both *taf1* and *ubc6* produced the most severe phenotype of male sterility resulting from failure to enter meiosis. This phenotype was also observed for the knockdown of *uba1*. Meiotic chromosome segregation defects were observed in the knockdowns of six E2 ligases (*bruce*, *CG7656*, *CG8188*, *ubc2*, *ubc10* and *ubcE2H*). Fourth chromosome missegregation (which may not be detectable cytologically) was observed via genetic crosses for knockdowns of four additional E2 ligases (*CG4443*, *CG9602*, *CG10862* and *CG17030*). Our results reveal that multiple ubiquitin ligase complexes are necessary for entry into meiosis and meiotic chromosome segregation in males. Several of these have not formerly been implicated in meiosis and future work towards understanding their roles will be focused on identification of their E3 ligase partners and their protein targets.

**436C A Borealin-HP1 Interaction Regulates Chromosome Passenger Complex Binding to Chromosomes and Movement to Microtubules** Manisha Persaud, Kim McKim, Janet Jang Rutgers University -- New Brunswick

Mitosis and meiosis are the most exciting and elaborate processes that occur during the life of dividing cells however, defects in these processes can lead to aneuploidy and lethal phenotypes. Errors in chromosomal segregation in oocytes lead to infertility and birth defects. We use Drosophila to understand the mechanisms of homologous chromosome bi-orientation that occurs during meiosis I, and the features of the oocyte-spindle that make it susceptible to segregation errors. The chromosome passenger complex (CPC) is a highly conserved master regulator that is required for oocyte spindle assembly, kinetochore assembly and homologous chromosome biorientation. It is a complex consisting of the proteins INCENP, Aurora B kinase, Survivin and...
Borealin. The CPC is recruited by the chromosomes and then moves to the microtubules. The mechanisms of this movement and the CPC signaling pathway that recruits and promotes the activity of kinetochore and spindle proteins remains largely unknown. Our previous research has led to a model that Borealin promotes spindle assembly in Drosophila oocytes by interacting with nucleosomes or the chromatin protein HP1. It was hypothesized that a Borealin-HP1 interaction is necessary for CPC binding to the chromosomes and movement to the microtubules. We are testing this model by mutating two domains within Borealin that may support this function: (1) a domain for binding HP1 and (2) a domain for binding microtubules. For these experiments, we have generated two Borealin shRNA lines and found that depletion of Borealin by RNAi showed phenotypes similar to aurB and or Incenp RNAi; complete loss of kinetochore and spindle assembly. Interestingly, many of the oocytes exhibited fragmented karyosomes, suggesting a role for the CPC in oocyte chromatin organization. We are currently generating RNAi resistant transgenes to that express borr mutants that are defective interacting with either nucleosomes / HP1 or microtubules. Parallel to these studies, we are examining the phenotype of Incenp mutants predicted to be defective in microtubule interactions and trying to directly detect interactions between HP1 and either INCENP or Borealin in oocytes.

437A Regulation of Meiotic Kinetochore-Microtubule Attachments by the RZZ Complex Joanatta Shapiro, Janet Jang, Kim McKim Waksman Institute of Microbiology, Rutgers University, Piscataway, NJ

Meiosis is a conserved process of cell division occurring in sexually reproducing eukaryotes which produces gametes with half the number of chromosomes. Errors in this process result in infertility, miscarriages, and genetic disorders. One poorly understood protein in meiosis, ROD, is a component of the ROD-ZW10-Zwilch (RZZ) complex and localizes to the kinetochores. The RZZ complex is involved in attachment error correction and the spindle assembly checkpoint (SAC) during mitosis. Error correction involves the selective stabilization and destabilization of kinetochore-microtubule (KT-MT) attachments while the SAC delays entry into anaphase until accurate attachments form. Mitotic evidence suggests that RZZ at the kinetochores prevents the formation of premature stable attachments. Attachment stabilization requires RZZ removal, a process called streaming which is suggested to depend on the microtubule motor protein Dynein and its recruitment factor Spindly. However, the role of ROD in these processes is not well characterized in female meiosis, which is unique in that it lacks the centrosomes and involves the segregation of homologous chromosomes rather than sister chromatids.

To analyze the function of ROD in female meiosis I, we observed ROD behavior in mutant or RNAi oocytes for proteins hypothesized to interact with RZZ. Female Drosophila depleted of ROD in the ovaries are sterile and have chromosome orientation defects, suggesting that ROD is required for correct KT-MT attachments. We found that ROD recruitment to the kinetochores is mediated by amino acids 154-840 of the kinetochore protein SPC105R. Furthermore, while ROD is not required for the creation of the initial KT-MT attachments, it may have a role in preventing or correcting erroneous attachments. Surprisingly, ROD removal may not be required for creating stable attachments, as in mitosis. Therefore, meiotic RZZ streaming may depend on the cell cycle stage rather than attachment status. We also found that ROD streaming in oocytes is at least partly Spindly-independent even though Spindly is required for meiosis I, suggesting potential Dynein-independent mechanisms of RZZ removal in meiosis. In the absence of microtubules, ROD expands around the kinetochores, likely to facilitate microtubule capture and contribute to SAC signaling. Determining how ROD regulates microtubule attachments will provide valuable insight into how pairs of homologous chromosomes segregate to maintain meiotic fidelity.

438B Genome-wide RNAi screen for new meiotic genes in Drosophila melanogaster Joel Sop, Tyler DeFosse, Ayla Boyd, Faith Verderose, Joanatta Shapiro Rutgers University

The sex cells, egg and sperm, are formed through a form of cellular division known as meiosis, resulting in four genetically unique daughter cells. During meiosis, accurate chromosome segregation is vital for reproductive success. Occasionally, chromosomal segregation errors occur, resulting in gametes with an incorrect number of chromosomes, a condition known as aneuploidy. Aneuploidy is known to be the leading cause of infertility in women. Aneuploids that survive cell division can cause spontaneous abortions or genetic disorders including Down’s Syndrome. Aneuploidy in women rapidly rises after the age of 35, a phenomenon known as the Maternal Age Effect. While several genes required for meiosis have been identified in Drosophila, most of these were identified in limited forward genetic screens or based on conservation in other organisms. Many meiotic genes remain to be identified, particularly those required for fertility or if they are poorly conserved. The goal of this project was to identify and characterize new meiotic genes in the Drosophila melanogaster genome. Using tissue expression patterns from the FlyAtlas and modENCODE databases, we identified 542 Drosophila genes of interest because their expression levels are elevated in meiotically active tissues. RNAi approaches were then used to make a gameline targeted knockdown with the UAS/ Gal4 system to observe which of the genes of interest caused significant errors in meiosis (aneuploidy) or fertility. Out of the 265 genes we have tested
so far, 67 were identified due to increased nondisjunction or sterility. We are currently prioritizing these genes and selecting a subset for further study.

439C Characterization of the Immune Deficiency Pathway during female meiosis in *Drosophila melanogaster* Sarah Mashburn, William Gilliland DePaul University, Chicago IL

Organisms can defend against pathogens by significantly increasing the diversity of their progeny, so that some progeny are more likely to survive infection. This leads to the prediction that infection should cause an increase in recombination rates. This prediction was confirmed in a published study where female *Drosophila melanogaster* that were infected with the gram-negative bacteria *Providencia rettgeri* had increased recombination rates compared to control flies. However, the mechanism of this rate modulation is unknown. Our lab conducted an RNAi screen to identify genes that cause defects during chromosome congression, which identified *mustard (mtd)* as causing ~40% congression failure. We show this is caused by a 95% reduction in recombination rates, which overwhelms the distributive segregation pathway similarly to mutants that block recombination like *c(3)G* or *meiW68*. The *mtd* gene had previously been shown to be part of the Immune Deficiency (IMD) pathway, which primarily defends against gram-negative bacteria. In that study, they showed that a gain-of-function allele increased the survival rate of flies that were infected with the gram-negative bacteria *Vibrio cholerae*. These results suggest that the IMD pathway may be what modulates recombination rates in response to bacterial infection. We are testing this hypothesis by measuring if the recombination rate changes caused by *P. rettgeri* infection depends on *mtd* function.

440A Investigating chromosome-specific differences during meiosis Katherine Billmyre1, R. Blake Billmyre1, R. Scott Hawley2,3 1) Stowers Institute for Medical Research, Kansas City, MO, United States of America; 2) Department of Molecular and Integrative Physiology, University of Kansas Medical Center, Kansas City, KS, United States of America

Inheritance of a complete set of chromosomes is critical for fertility and production of viable offspring. However, all chromosomes are not the same. Cells face the challenge of building a system that can accurately segregate chromosomes with vast structural and size differences (as much as 5x in humans and 46x in *Drosophila*). However, the mechanisms underlying recombination and segregation of specific chromosomes are not understood. In *Drosophila melanogaster*, a set of partial loss-of-function synaptonemal complex (SC) mutants exhibit chromosome-specific defects in recombination and pairing during meiosis. The SC is a conserved meiotic structure that holds homologous chromosomes together and is necessary for the repair of double-strand breaks (DSBs) into crossovers. We are currently using a two-pronged approach of super-resolution imaging and genomics to examine the relationship between the SC, DSBs, and recombination on different chromosomes both in wildtype flies and those that exhibit chromosome-specific defects. Using next generation sequencing, we have found in a partial loss of SC background, that DSBs are able to be repaired as non-crossover gene conversions, even when they are unable to be repaired as crossovers. To further examine the importance of synopsis, pairing, DSB placement, and crossovers on the behavior of different chromosomes, we are combining the LacO/LacI system with super-resolution imaging and chromosome tracing to assess the meiotic landscape on individual chromosomes. Together these approaches will provide a one-of-a-kind map to elucidate the role of the SC in regulating meiosis.

441B The Effect of Heterozygous Inversion on Crossover Frequency near Inversion Breakpoints by High-Res Whole Genome Sequencing Nicole Crown, Haosheng Li Case Western Reserve University

Heterozygous inversions disrupt the meiotic crossover (CO) distribution by suppressing COs locally both within and outside of the inversion, and also triggering the interchromosomal (IC) effect, where the CO frequency on structurally normal chromosomes dramatically increases. We previously showed that these changes in the CO distribution are mediated by CO patterning mechanisms that alter the CO/NCO decision in *Drosophila melanogaster*. However, the exact decision making process of how the patterning mechanism exert its influence on a double-stranded break (DSB) to suppress COs outside of inversion breakpoints, as well as the pattern of suppression in terms of distance and severity, are still unknown. Traditionally, this analysis can be done by phenotypic scoring of recombinants, but we are currently analyzing the recombination mechanics of DSB repair by creating a fine-scale recombination map of the region near the breakpoints of an X chromosomal inversion dl-49 through whole genome sequencing of individual *Drosophila melanogaster* offspring. Our preliminary data show that from the distal breakpoint to telomere region, almost no COs were recovered. From the proximal breakpoint to centromere region, a bell-shaped pattern was observed, where the further away the location is from the breakpoint, the higher the CO frequency is (compared to the CO profile of completely wild type flies). To reduce variations in genomic background among tested flies, we are currently creating isogenized full-sibling stocks, and the experiment will be repeated using these new stocks for more robust data.

442C Meiotic Crossovers on Chromosome 4 induced by the Interchromosomal Effect in *Drosophila melanogaster* Joseph Terry, Savana Hadjipanteli, Nicole Crown Case Western Reserve University

Meiosis is essential for the production of genetically diverse gametes in all sexually reproducing eukaryotes. During
meiosis, genetic material is exchanged between chromosomes through homologous recombination, driving genetic diversity amongst gametes while simultaneously ensuring proper chromosome segregation. Recombination is initiated in response to programmed DNA double-stranded breaks (DSBs) across the genome. These DSBs are then repaired, resulting in a crossover (CO) or non-crossover (NCO).

When looking at genome-wide DNA DSB repair distribution, NCOs make up the majority of repair outcomes while a limited number of DSBs are repaired as COs. This imbalance between the total number of DSBs across the genome and the small number of those repaired as COs introduces the question: what dictates crossover patterning across the genome? Accumulating evidence has implicated three key phenomena to be involved in crossover patterning within Drosophila: crossover assurance, crossover interference, and crossover suppression. Interestingly, nearly all chromosomes are affected by these three CO patterning mechanisms, with the exception of chromosome 4 where meiotic COs never occur.

Disturbances in CO patterning, such as the attenuation of CO interference and suppression have been suggested to occur during the interchromosomal effect (IC). During the IC effect, the presence of a heterozygous inversion inhibits COs locally, while increasing CO frequency across unbalanced chromosomes. Here, we propose that CO patterning mechanisms indeed, are attenuated during the IC effect by generating a fly model that undergoes meiotic COs on chromosome 4 in the presence of a multiply inverted balancer chromosome. Using a fly model that undergoes meiotic COs on chromosome 4, we aim to generate CO distribution maps as a method to assess the impact of CO patterning mechanisms on recombination during the IC effect.

443A  Identification of Meiotic Recombination Nodule Proteins Utilizing Proximity Labeling Oscar Bautista, Haosheng Li, Gabrielle Bais, Nicole Crown Case Western Reserve University, Cleveland, OH

During meiosis, recombination between homologous chromosomes ensures genetic diversity among haploid products, but errors in this mechanism are a major source of human infertility. Recombination events, initiated by enzymatic-directed double-stranded breaks can result in crossovers (COs), involving reciprocal exchanges between homologues, or noncrossovers (NCOs), where a portion of DNA is copied from the donor homologue without altering it. Protein complexes known as recombination nodules are thought to reside around future crossovers and promote and catalyze recombination events. Although their function has been hypothesized, its protein composition has not been elucidated, nor has the mechanism through which it facilitates recombination outcomes. Additionally, due to their insoluble nature, previous attempts at isolating recombination nodules using classical biochemical purification techniques have failed. To address these issues, I will generate several fly lines containing proximity biotin-labeling protein, APEX2, conjugates of known recombination proteins. Using these fly lines and mass spectrometry, we will be able to identify novel protein interactors and create interaction maps to elucidate the proteome that is necessary for processing recombination.

Utilizing our model system, I eliminate the need to isolate recombination nodules and, instead, can reliably identify protein components and interactions to understand the mechanisms that facilitate meiotic recombination. The protein-protein interactions uncovered will also provide insight into the complex regulatory mechanisms that are required to initiate double-stranded breaks and subsequently repair them as NCOs or COs.

444B  Mechanism of bruno-mediated tolerance to P-element activity in Drosophila melanogaster germline Modupeola Bolaji, Erin Kelleher, Vanessa Marcano University of Houston

Transposable elements (TEs) are mobile genetic elements whose proliferation in the germline induce germ cell loss and sterility. Hosts reduce this fitness cost through resistance, which involves suppression of TE proliferation, or tolerance, where the germline withstands the damaging effects of transposition. While host resistance to TEs by piRNAs is studied extensively, little is known about host factors that confer tolerance. Our lab identified bruno as a source of natural variation in tolerance to P-element DNA transposon induced germ cell loss in Drosophila melanogaster. Bruno is an RNA binding protein and a translation repressor that plays important roles in the regulation of oocyte development in adults. However, the role of bruno in determining tolerance to transposition is unknown.

Recent studies have shown that P-element induced dysgenesis triggers loss of germ cells in the larval stage. While bruno previously had no known function in larvae, we have discovered that Bruno protein is expressed in larval primordial germ cells (PGCs) and that Bruno function impacts P-element induced germ cell loss specifically in the larval stage. We are currently testing two hypotheses to explain the relationship between Bruno function and PGC loss: 1) hybrid dysgenesis increases bruno expression in PGCs, resulting in activation of bruno-dependent differentiation pathways and 2) bruno increases transposition of P-element, leading to increased DNA damage and loss of PGCs.

445C  Characterizing the composition and morphology of the germ plasm in the wasp Nasonia vitripennis Allie Kemph1, Jeremy Lynch1, Alexey Arkov2, Kabita Kharel2, Samuel Tindell2 1) University of Illinois at Chicago; 2) Murray State University

Specification of germ cell fate during embryogenesis is an essential process in sexually reproducing organisms to ensure the correct transmission of parental genetic information to offspring. In many cases it involves germ plasm, a specialized cytoplasmic organelle composed largely of mRNA and RNA-binding proteins that drives germ cell fate determination.
There is great variability in morphology and composition among organisms that contain germ plasm. For example, in *Drosophila* the germ plasm consists of many relatively small granules that remain associated with the posterior pole of the egg until they are taken into individually budding pole cells. In contrast, in the wasp *Nasonia vitripennis*, germ plasm assembles into an extremely large, dynamic structure, called the “oosome”. The oosome migrates anteriorly to 50% egg length, before returning to the posterior pole where a single large bud containing multiple nuclei emerges during pole cell formation. While much is known about the composition of the fly germ plasm, how the structure of the oosome compares is as yet unknown. Here we describe our progress in characterizing the dynamic oosome morphology and determining the spatial arrangement of several oosome mRNA and protein components. These results will be compared to the known homotypic clustering and dynamics of the fly polar granules.

446A  **bourbon** interacts with known germline sex determination regulator *otu* and promotes the expression of *sxl* in the *Drosophila* female germline Marianne Mercer, Michael Busczak University of Texas Southwestern Medical Center, Dallas, TX

Sexually reproducing species exhibit a wide variety of mechanisms to establish male and female identity at the cellular level. In *D. melanogaster*, *sex lethal* (*sxl*) is the master switch of sex determination in somatic cells; however, the process of sex determination in the germline remains poorly understood. Previous studies show that *ovarian tumor* (*otu*) acts genetically upstream of and promotes the expression of *Sxl* in the female germline, which ultimately leads to a female fate. How Otu protein controls Sxl expression is less clear. Through IP-mass spec analysis we find that Otu physically interacts with Bourbon (*Bbn*), a previously uncharacterized protein. *bbn* displays enriched expression in germ cells, and *bbn* null mutant females exhibit agamic ovarioles and cystic germline tumors. These germline tumors do not express Sxl protein and resemble *sxl* sterile mutants, displaying expanded and overlapping expression of Bam and Nanos. Transgenic expression of *sxl* can partially rescue this phenotype. We tagged *bbn* endogenously with an HA tag and found it is expressed in germline stem cells, cystoblasts and two cell cysts, similar to the expression pattern of cytoplasmic Sxl protein. In addition, Bbn is enriched in developing oocytes. Because the phenotypes and localization pattern of *bbn* are strikingly similar to those of *otu* and they physically interact, we are testing how Bbn impacts the ability of Otu to promote Sxl expression. We hypothesize that Otu, the founding member of a family of deubiquitinases, deubiquitinates Sxl to prevent its degradation by the proteasome. Future experiments will help us determine if Bbn regulates Otu expression levels, localization, substrate specificity or catalytic activity.

447B  **Targeted mutagenesis of orco disrupts fertility in the second gonotrophic cycle in the Aedes aegypti** mosquito Olayinka David, Kevin Sanchez, Anthony Bellantuono, Andre Costa-da-Silva, Matthew DeGennaro Florida International University

Mosquitoes are the deadliest animals to humans. They are responsible for over half a million deaths annually and get more than 200 million people around the world sick every year. Female mosquitoes of many species require vertebrate blood for egg development. A female mosquito undergoes multiple discrete rounds of reproductive cycles also known as gonotrophic cycles. A gonotrophic cycle spans the period from bloodmeal intake to the time the mature oocytes are deposited. Mosquitoes are capable of transmitting diseases to humans and other vertebrate hosts during bloodmeal intake. Transmission of mosquito-borne diseases to hosts typically occurs during the second round of blood ingestion, an event that often coincides with the start of the second gonotrophic cycle. Mosquitoes rely on odorant receptors (ORs) for detecting chemical molecules in their surroundings. This chemosensory potential is necessary for processes such as vertebrate host localization, nectar seeking, identifying suitable mates, and finding egg-laying sites. Odorant receptor *orco* is an insect-specific obligate counterpart necessary for the function of tuning ORs. Studies have reported the expression of *orco* in ovaries and testes across different insect taxa. These receptors have been shown to activate both human and mosquito sperm. To determine the role of *orco* in *Aedes aegypti* reproduction, we employed individual mosquito fecundity and fertility assays to quantify the reproductive output of *Aedes aegypti orco*−/− mosquitoes across two gonotrophic cycles. Surprisingly, we found that eggs produced by mutant mosquitoes have a significantly reduced hatch rate in the second reproductive cycle compared to controls. Histochemical assays revealed that the unhatched eggs were fertilized but the resulting embryos failed to attain full development. Additionally, using mixed genotype mating experiments, we found that the embryo developmental defect is a maternal effect, suggesting that *orco* is required in the female mosquito for optimum embryo viability. In control experiments, we quantified the amount of ingested blood, the number of sperm stored as well as the number of eggs laid per female mosquito and found no difference between wild-type and mutant mosquitoes. Lastly, when the *orco* gene was reconstituted into the *orco*−/− background, the reproductive phenotype was reverted. Our findings inform a novel vector control approach that targets mosquito reproduction through halted embryo development.

448C  **Searching for the female receptor for the D. melanogaster** seminal fluid protein *ovulin* Mengye Yang1, Melissa White1, Jennifer Ager-McGlaughon1, Geoffrey Findlay1, Mariana Wolfner1 1) Cornell University, Ithaca, NY; 2) College of the Holy Cross, Worcester, MA

Males transfer hundreds of seminal fluid proteins (Sfps) along with sperm to the female reproductive tract during...
449A RNA–protein interaction mapping via MS2-based APEX2 targeting in the Drosophila ovary Kwan Yin Lee, Elizabeth R. Gavis Department of Molecular Biology, Princeton University, Princeton, NJ

Mechanisms of mRNA regulation commonly involve the engagement of protein factors capable of influencing transcript biology. Proteins can engage with mRNAs in stable ribonucleoprotein complexes or form more fluid and dynamic structures by coalescing into phase-separated condensates. Currently, how the physical features of these complex structures are associated with mRNA regulation remains poorly understood. We are combining proximity-dependent labeling by APEX2 with the MS2/MCP system to spatially define the molecular environment proximal to specific mRNAs in the Drosophila oocyte. In late-stage oocytes, nanos mRNA is incorporated into phase transitioned condensates known as germ granules that are a conserved feature of germline development. The nanos mRNAs occupying germ granules form spatially distinct clusters within the granule compartment, each containing multiple copies of the mRNA. We aim to understand the role of these clusters in mRNA regulation by defining their surrounding molecular environment using the MS2-based APEX2 targeting system described here. To test the method, we have tethered MCP-APEX2 to nanos mRNA. APEX2 dependent biotinylation is detected together with nanos mRNA at multiple sites across Drosophila egg chambers. Biotinylated proteins nearby nanos mRNA will be isolated and identified by mass spectrometry.

450B Genetic interactions between new bag-of-marbles mutants and the endosymbiont bacteria Wolbachia in D. melanogaster Miwa Wenzel, Charles Aquadro Cornell University

The D. melanogaster protein-coding gene bag of marbles (bam) plays a key role in early male and female reproduction by forming complexes with partner proteins to promote differentiation in gametogenesis. A bam null mutant and a bam partial loss-of-function hypomorphic mutant result in sterility and reduced fertility, respectively. Like another germline gene, Skl, bam genetically interacts with the endosymbiont Wolbachia, as Wolbachia rescues the reduced fertility of the hypomorphic mutant. Here, we explored the specificity of the bam Wolbachia interaction by combining an alanine scanning approach with phylogenetic sequence data to efficiently generate new mutants to test for Wolbachia rescue. We have generated several new mutants with reduced fertility in one of bam’s documented binding regions, highlighting the functional importance of certain residues for bam’s activity. All of these new reduced fertility mutants are rescued by Wolbachia in females. Additionally, we find that Wolbachia also rescues the reduced fertility of a bam transgenic allele in male D. melanogaster, revealing the first genetic interaction between Wolbachia and germline genes in males. Additional mutants in other documented binding regions of bam as well as bam RNAi knockdowns are currently being evaluated to further elucidate the specificity of the bam Wolbachia interaction. If Wolbachia rescue remains specific to this single binding region, it would suggest that the Wolbachia interaction occurs in this pathway of bam’s function. Better understanding the nature of bam’s interaction with Wolbachia will allow us to evaluate hypotheses about the potential contribution of Wolbachia as a driver of bam’s rapid evolution in the D. melanogaster group.

451C Nuclear actin is a critical regulator of Drosophila germline stem cell maintenance Nicole Green, Tina Tootle Carver College of Medicine, University of Iowa, Iowa City, IA

While actin was observed in the nucleus decades ago, nuclear functions of actin dynamics have only recently become widely acknowledged. Indeed, nuclear actin regulates the activity of RNA polymerases and transcription factors, chromatin organization via remodeling complexes and histone deacetylases, and nuclear integrity. Furthermore, nuclear actin is involved in determining differentiation state, deciding cell identity, and reprogramming cells to a pluripotent state. Our studies investigate the roles of nuclear actin in regulating stemness using a model tissue, the Drosophila ovary. The Drosophila ovary is made up of 15-20 ovarioles of sequentially developing follicles. Germline stem cells (GSCs) reside in a niche at the anterior tip of each ovariole in a structure known as the germarium and give rise to all germline cells. GSCs are maintained by asymmetric divisions that retain a stem cell and produce a daughter cell, which continues to incompletely divide to produce 2-, 4-, 8-, and 16-cell germline cysts. Our lab previously defined several distinct pools of nuclear actin in the ovary by screening established actin labeling tools. One of these actin pools stained by anti-actin C4 is found in both the nucleoplasm and nucleolus of GSCs. These C4 positive nuclear actin pools are present in a dynamic...
pattern across the germarium and are lost as cells become more differentiated. This trend suggests nuclear actin may play a role in regulating stemness. To test this hypothesis, we overexpressed NLS-actin constructs in germline cells of the germarium. When we increased monomeric nuclear actin (NLS-Act5C
\textsuperscript{G13R}), GSC maintenance is disrupted and there is progressive germline loss which results in empty germaria. As nuclear actin is enriched in the nucleolus and previous studies uncovered that nucleolar functions are essential for GSC maintenance, we were curious whether nuclear actin regulates nucleolar functions. Using nucleolar structure as a readout for nucleolar function, we find that increasing clear monomeric actin results in increased nuclear size and hypertrophic nucleoli. These nucleoli are also deformed and fragmented suggesting altered nucleolar functions. These results indicate that nuclear actin dynamics regulate key nuclear functions, including nucleolar activity, which are necessary for maintaining stem cells.

452A  Evaluating the Effect of Architectural Features on Border Cell Migration in Drosophila Alexander George, Bradford Peercy, Michelle Starz-Gaiano University of Maryland Baltimore County

Collective cell migration is paramount throughout a multicellular organism’s life from embryogenesis to adult tissue maintenance. Many cell migration studies are conducted in vitro, but to understand the full complexity of the living tissue environment, more in vivo studies are required. Using the border cells, which navigate as a cluster through the 3-D terrain of the Drosophila egg chamber during oogenesis, we can investigate the impact of physical architecture on collective cell migration and migration-related signaling in vivo. We observe small, acellular gaps at cell-cell junctures along the border cell migration route and hypothesize that these architectural features directly affect the migration behaviors of the border cells, in part by altering local distributions of secreted chemical attractants (chemoattractants). In silico, we have demonstrated that these gaps can affect the distribution of the morphogen that specifies border cell fate, as well as the distribution of secreted chemoattractants. Using staining and imaging techniques, we characterized the geometry and distribution of these gaps in fixed tissue and observed changes in border cell behavior at cell-cell junctures where gaps are observed in live tissue. Furthermore, flooding these gaps with above-endogenous levels of chemoattractant delayed border cell migration, and thus we are considering the effect of receptor dynamics on cluster migration. Additionally, we are using two complementary methods to reveal the chemoattractant gradient and evaluate the influence of the egg chamber’s terrain on signal distribution and consequently, border cell migration. First, by inducing extra border cells throughout the egg chamber and examining their speed and persistence, we infer the positions of local hotspots of concentrated chemoattractant. And second, by tagging chemoattractants to visualize them, we can examine signal distribution in real-time. This data will further inform and allow us to refine our in silico models on how border cells might respond to the chemoattractant gradient. Our studies emphasize the importance of considering physical features of the tissue architecture when examining collective migration and migration-relevant signaling in vivo.

453B  A Genetic Screen Identifying E2s and E3s Involved with Maternal Protein Clearing During the Maternal to Zygotic Transition Calvin Bleskan\textsuperscript{1,2,3}, Chloe Briney\textsuperscript{2,3}, Jesslyn Henriksen\textsuperscript{2,3}, Olivia Rissland\textsuperscript{2,3}, Hector Cobian\textsuperscript{1,2,3}  
1) Metropolitan State University, Denver, CO, USA; 2) Department of Biochemistry and Molecular Genetics, University of Colorado School of Medicine, Aurora, CO, USA; 3) RNA BioScience Initiative, University of Colorado School of Medicine, Aurora, CO, USA

The maternal-to-zygotic transition (MZT) is an essential process during early animal embryogenesis where developmental control shifts from maternally deposited gene products to a newly made zygotic one. Although the MZT typically considered from the perspective of RNA, we and others have found that maternal protein clearance is a key part of this transition. As part of a multi-year undergraduate research project, we are performing an RNAi screen to identify E2s and E3s required for oogenesis and embryogenesis. This RNAi screen consists of expressing RNAi constructs using an ovariyspecific UAS driver and of quantifying various phenotypes from these female flies (including egg laying, egg viability, and other phenotypes). Here, we will describe the results for ten randomly selected genes: skp2, cul-1, Elo-C, Gus, SAK, CSN5, cul-2, skpA, UBC-4, and UBC-D6[BC1]. Preliminary results indicate that knock down of cul-1 resulted in lethality for a majority of embryos. Because Cul-1 is a component of the SCF E3 complex and the SCF complex regulates cell cycle progression and the removal the Smaug RNA binding protein, this result is consistent with Cul-1 performing an essential role during embryogenesis, but not oogenesis. Interestingly, embryos with UBC-4 depletion showed a developmental delay in pupae eclosion, although the underlying mechanisms are unknown. Together, these results demonstrate the potential of this screen to provide a foundation for future research on protein clearance during the MZT.

454C  A Genetic Screen for Identifying E2s and E3s Involved in Protein Clearance During the Maternal-to-Zygotic Transition Hector Cobian\textsuperscript{1,2,3}, Chloe Briney\textsuperscript{1,2}, Jesslyn Henriksen\textsuperscript{1,2}, Calvin Bleskan\textsuperscript{1,2,3}, Olivia Rissland\textsuperscript{1,2}  
1) RNA BioScience Initiative, University of Colorado School of Medicine, Aurora, CO; 2) Department of Biochemistry and Molecular Genetics, University of Colorado School of Medicine, Aurora, CO; 3) Metropolitan State University of Denver, Denver, CO

The maternal-to-zygotic transition (MZT) is an essential developmental process whereby maternal gene products deposited into the oocyte are degraded, and zygotic gene products take their place. The degradation of maternal RNA has been well studied and is known to be an important part of the MZT. However, the removal of maternally deposited proteins is far less studied. The main mechanism for protein degradation pathway works through the
ubiquitin-proteosome system, which uses E1s, E2s, and E3s and culminates in the ubiquitylation of target proteins, thus marking them for degradation. The *Drosophila melanogaster* gene encodes dozens of E2s and hundreds of E3s, but we know very little about their involvement in the MZT. As part of a multi-year undergraduate research program, we are performing an RNAi screen to identify E2s, E3s, and related proteins that are essential for oogenesis and embryogenesis. Here, we will describe the results of ten randomly selected E2 and E3 genes: CG11658, Cul-3, Cul-5, EloB, jaf, slmb, l(3)73Ah, CSN6, Nedd4, and Pros alpha s. We will present results on the effectiveness of RNAi knockdown by RT-qPCR and western blotting. We will describe the extent to which each gene affects: egg laying; embryonic viability; other developmental effects. In the future, we will be able to use unbiased techniques, like mass spectrometry, to identify putative targets of maternally required E2s and E3s. Together, this screen will help reveal the factors required for proper protein degradation in the MZT.

455A Octopaminergic/tyraminergic Tdc2 neurons regulate sperm preference in female *Drosophila melanogaster* Dawn Chen, Andrew Clark, Mariana Wolfner Cornell University, Ithaca, NY

In polyandrous internally fertilizing species, a multiply-mated female can exert preference on which stored sperm she uses to fertilize her eggs. The female’s ability to assess sperm quality and compatibility is essential for her reproductive success, and represents an important aspect of postcopulatory sexual selection. In *Drosophila melanogaster*, previous studies demonstrated that the female nervous system plays an active role in influencing progeny paternity proportion, and suggested a role for octopaminergic/tyraminergic Tdc2 neurons in this process. Here, we report that inhibiting Tdc2 neuronal activity causes females to produce a higher-than-normal proportion of first-male progeny. This difference is not due to differences in sperm storage or release, but attributable to the suppression of preference for second-male sperm that normally occurs in control females. We further show that a subset of Tdc2 neurons innervating the female reproductive tract was largely responsible for the progeny proportion phenotype that was observed when Tdc2 neurons were inhibited globally. On the contrary, over-activation of Tdc2 neurons does not further affect sperm storage and release or progeny proportion. These results suggest that octopaminergic/tyraminergic signaling allows a multiply-mated female to exert sperm preference, and identify a new role for the female nervous system in postcopulatory sexual selection.

456V Transcriptional and mutational signatures of the aging germline Evan Witt, Christopher Langer, Li Zhao Laboratory of Evolutionary Genetics and Genomics, The Rockefeller University, New York, NY

Aging is a complex biological process which is accompanied by changes in gene expression and mutational load. In many species including humans, old fathers pass on more paternally-derived de novo mutations, however, the cellular basis and cell types driving this pattern are still unclear. To understand the root causes of this phenomenon, we performed single-cell RNA-sequencing (scRNA-seq) on testes from young and old male *Drosophila*, as well as genomic sequencing (DNA-seq) on somatic tissue from the same flies. We found that early germ cells from old and young flies have similar mutational loads, but older flies are less able to remove mutations during spermatogenesis. This indicates that germline mutations arise from primarily non-replicative factors, and that the increased mutational load of older males is due to differences in genome maintenance activities such as repairs to DNA damage. We also found that T:A mutations are enriched in older flies, and transcription-related enrichment terms are depleted in older males. Early spermatogenesis-enriched genes have lower dN/dS than late spermatogenesis-enriched genes, supporting the hypothesis that late spermatogenesis is the source of evolutionary innovation. This transcriptional disruption is reflected in the decreased expression of genome maintenance genes in early germ cells of older flies, as well as potentially aberrant transcription of transposable elements in the aging germline. Our results provide novel insights into the transcriptional and mutational signatures of the male germline.

457V Modeling effects of human disease variant of Barrier-to-Autointegration on oogenesis Felipe Rodriguez1, Tingting Duan1, Katherine Mathews2, Pamela Geyer1 1) University of Iowa. Department of Biochemistry and Molecular Biology; 2) University of Iowa. Stead Family Department of Pediatrics

The nuclear envelope is a complex structure that defines the cellular nucleus. Lying beneath the inner nuclear envelope is the nuclear lamina (NL), a protein network comprised of lamins and lamin-associated proteins. One abundant protein in this network is the Barrier-to-Autointegration Factor (BAF), a small dimeric protein that binds the NL LEM domain proteins, histones, and DNA, all properties that contribute to the organization of chromatin within the nucleus. Mutations in the amino terminus of BAF cause human diseases associated with age-related declines in tissue function, including Nestor Guillermo progeria (NGP) and neuromuscular disease (NMD). To gain insights into the mechanism of these diseases, we are modeling disease associated mutations in Drosophila BAF, a protein with 63% identity to human BAF. We are investigating the effects of the disease variants of BAF on the ovary, as BAF is essential for sustained oogenesis. We find that oogenesis is impaired in NGPs flies, with females laying few eggs that do hatch. Our data suggest that reduced egg production is linked to altered GSC mitosis, leading to increased DNA damage and death of transit amplifying cells, defects associated with activation of the DNA damage response kinase Checkpoint kinase 2. We are initiating studies of the NMD variant. Our Drosophila studies of the NMD variant will build from our observations that
NMD patient fibroblasts display age-dependent increases in nuclear ruffling, higher levels of heterochromatin, and increased DNA damage, defects linked to changes in the cell cycle that lead to faster proliferation. Taken together, our studies will provide an understanding of the role of BAF in development and tissue homeostasis.

**458V**  
Polycomb group (PcG) proteins prevent the assembly of higher order repetitive structures during meiosis  
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The synaptonemal complex (SC) is a proteinaceous scaffold that is assembled between the paired homologous chromosomes during the onset of meiosis. The SC stabilizes chromosome pairing and is important for crossovers formation, recombination and accurate segregation of meiotic chromosomes. Timely expression of SC genes is essential for SC assembly and successful meiosis. However, SC components have an intrinsic tendency to self-organize into alternative higher order repetitive structures, which are not assembled between the paired homologs and whose formation is potentially deleterious for meiosis and gametogenesis. This creates an interesting conundrum, where SC genes need to be robustly expressed during mitotic to meiotic transition, but their expression must be carefully regulated to prevent the formation of abnormal SC structures. In this manuscript, we show that the Polycomb group protein Sfmbt, the *Drosophila* ortholog of human MBTD1 and L3MBTL2, is required to avoid excessive expression of SC genes during prophase I. Although SC assembly is normal after Sfmbt depletion, SC disassembly is abnormal with the formation of a complex network of alternative SC structures (polycomplexes) within the oocyte. Overexpression of the SC gene corona and depletion of other polycomb group proteins are similarly associated to polycomplexes formation during SC disassembly. These polycomplexes are highly dynamic and have a well-defined periodic structure. Further confirming the importance of Sfmbt for female gametogenesis, germ line depletion of this protein is associated to significant metaphase I defects and a reduction of female fertility. Polycomb group proteins are therefore crucial for coherent gene expression during prophase I.

**459V**  
Identification of factors regulating individualization. *Sepideh Dadkhah,* Douglas Harrison University of Kentucky, Lexington, KY

Sperm development in *Drosophila* has many similarities to spermiogenesis in other organisms. Individualization is one of the later stages of spermiogenesis in which the 64 interconnected spermatids in a cyst are separated and excess cytoplasm eliminated. The synchronous differentiation of spermatids from interconnected spermatocytes is evolutionarily conserved. While JAK/STAT pathway activation in the somatic cyst cells is required for individualization, the mechanism by which it controls the process is not known. The aim of this project is to identify effectors regulated by JAK/STAT signaling that initiate individualization.

Using RNA-seq, we have examined expression profiles from testes in which JAK/STAT signaling has been impaired prior to individualization by expressing the negative regulator Eye-transformer/Latran (ET/Lat) and compared to testes dissected from wild type flies. Estimates of transcript abundance of the control and experiment profiles were calculated. Differential expression analysis identified more than 400 genes that were differentially expressed upon the arrest of JAK/STAT signaling at elongation. The differentially expressed genes identified in the RNA-Seq analysis were prioritized based on being associated with relevant GO terms and relation to the JAK/STAT pathway. Knockdown of 38 prioritized genes was carried out by utilizing the UAS-Gal4 system to express corresponding RNAi constructs with somatic and germine drivers. Functional analysis was done by quantitative analysis of elongated spermatid nuclei bundles and investment cones, both hallmark events of progression of individualization. From this functional analysis, genes involved in extracellular matrix and cell signaling were found to influence individualization.

**460V**  
Distinct downstream effectors downstream of InR activity control multiple aspects of oogenesis. *Tancia Bradshaw,* Alissa Armstrong University of South Carolina

Approximately 40% of adults in the United States are obese. Adipocytes, the primary cellular component of fat tissue, are nutrient-sensitive and secrete adipokines that control physiology, including metabolism, appetite, and insulin sensitivity. Despite the intricate relationship between dietary input, adipose tissue, and peripheral organ function, we are at the tip of the iceberg regarding our understanding of the cellular and molecular mechanisms underlying adipose communication to other tissues. The *Drosophila* ovary receives nutritional signals from the fat body with multiple nutrient-sensing pathways functioning in adipocytes to control oogenesis. Insulin/insulin-like growth factor signaling (IIS) within adult adipocytes remotely controls oocyte production at distinct stages of oogenesis. The PI3K/Akt1 axis in...
adipocytes promotes germline stem cell (GSC) maintenance via SGG, *Drosophila* GSK3beta, yet its targets in this context remain to be identified. Here, we measured the expression of candidate SGG/GSK3beta substrates in adipocytes and assessed their role in regulating GSC number. Using the UAS/Gal4 system for adipocyte-specific knockdown of four putative targets, *ATPCL*, *chb*, *porin*, and *SREBP*, we find that *SREBP* acts within adipocytes to regulate GSC maintenance. In addition, we find that a second axis downstream of the insulin receptor, the Ras/MAPK signaling pathway, acts cell autonomously to control adipocyte size and remotely to regulate the survival of early and late germline stem cell progeny – germline cysts and vitellogenic egg chambers, respectively. Future studies will identify how SREBP activity within adipocytes controls GSC maintenance as well as additional SGG/GSK3beta substrates within adipocytes required to communicate to the ovary. These studies highlight the complex mechanisms that underlie inter-organ communication.

### 461V Warm and cold temperatures have distinct germline stem cell lineage effects during *Drosophila* oogenesis

*Ana Caroline Gandara*, Daniela Drummond-Barbosa Johns Hopkins University, School of Public Health, Baltimore, MD

Reproduction of all organisms is shaped by their external environments. As climate change alters the abundance and distribution of many types of organisms around the globe, investigating how temperature impacts the reproduction of organisms has become a major area for scientific investigation. Insects are cold-blooded animals and therefore particularly sensitive to environmental temperatures. Oogenesis is highly sensitive to a variety of environmental and physiological factors; however, it remains largely unknown how exposure of *Drosophila melanogaster* adult females to chronic thermal stress affects oogenesis. To directly address this question, we incubated newly-eclosed *yw* adult flies (raised at room temperature, 23°C) at 18°C (cold), 25°C (optimal), or 29°C (warm) for 20 days at constant humidity (>70%) and found that the rate of egg production was reduced in females at both warm and cold temperatures, albeit through distinct cellular mechanisms. Chronic exposure of females to 18°C improved the maintenance of germline stem cells, survival of early germline cysts, and hatching rates but reduced the rate of follicle growth with no obvious effect on vitellogenesis. By contrast, in females at 29°C, germline stem cell numbers and follicle growth were comparable to those in females at 25°C, while early germline cyst death and degeneration of follicles were markedly increased, and hatching rates plummeted over time. Finally, we uncovered a novel role for the warm temperature receptor TrpA1 in mediating the effects of 29°C on oocyte quality. These findings are widely relevant not only to cold-blooded organisms, which exhibit limited thermoregulation, but also potentially to warm-blooded organisms, which are subject to hypothermia, heatstroke, and fever.

### 462V Obesity and oogenesis in *Drosophila*: Increased fat storage is not sufficient to impair fertility

*Rodrigo Dutra Nunes*, Daniela Drummond-Barbosa Johns Hopkins Bloomberg School of Public Health

Obesity is highly correlated with infertility; however, the underlying mechanisms remain largely unknown. Our previous research revealed complex roles of adipocytes in multiple steps of *Drosophila* oogenesis, and others have shown that a high sugar diet (HSD) causes obesity and reduces egg production. To carefully investigate how obesity affects oogenesis, we increased dietary sugar content from 5% (ND) to 30% (HSD). In accordance with published studies, females on HSD had a 3-fold increase in triacylglycerol content and lipid droplets, a 2.5-fold increase of glycogen content, and a drastic reduction in egg production and quality compared to ND controls. In addition, we found that these obese females ingested 5 times less food and had increased death of early germline cysts and vitellogenic follicles. In these experiments, however, HSD drives obesity, and it is not clear whether changes in oogenesis are due to HSD, obesity, or both. To test how obesity alone impacts oogenesis, we induced adult adipocyte-specific RNAi against the *brummer* or *adipose* genes (both of which inhibit excess fat accumulation). Adipocyte-specific *brummer* knockdown led to a marked increase in triacylglycerol content and adipocyte lipid droplet size, comparable to those observed in HSD-induced obese females. Strikingly, these highly obese females did not show any significant differences in food ingestion, glycogen content, specific steps of oogenesis, rates of egg production, or oocyte quality relative to non-obese control females. We obtained similar results in obese females generated by adipocyte-specific RNAi against *adipose*. These results indicate that increased fat storage is not sufficient to reduce fertility. It remains unclear, however, if obesity is required for the reduced fertility of HSD-induced obese females: it is possible that either HSD alone or in combination with obesity is responsible for their fertility defects. Our future studies will investigate the underlying mechanistic basis for the oogenesis differences we observe in these different types of obese females.

### 463V Validation of candidate genes influencing egg size in cold-adapted *Drosophila melanogaster*

*Cecelia Miles*1, Allison Hoefakker1, Intisar Koch2, Rachel Flynn2, Rosanna Beraldi2 1) Augustana University, Sioux Falls, SD; 2) Mount Marty University, Yankton, SD

Egg size is a highly polygenic trait that is closely related to fitness in *Drosophila melanogaster*. Cold-adapted *D. melanogaster* are known to produce larger eggs (Azevedo et al. 1996). The Miles lab previously identified 22 candidate genes influencing egg size based on whole-genome sequence data in multiple cold-adapted lines. Here we used semi-quantitative PCR to try and validate five of our 22 candidates (*cappuccino*, *stathmin*, *dreadlocks*, *lilliputian*, and gene CG31690) by comparing expression in cold-adapted inbred lines to the standard lab line OregonR. Expression differences in *stathmin* were found in a majority of the cold-adapted lines and appear to represent the absence of a
splice variant. This is the first validation of a gene from our candidate list. We also report a significant upregulation in actin observed in half of our cold lines. We are currently following up on these results using qPCR.

464V  Genetic Requirement of IC effect Bowen Man, Nicole Crown Case Western Reserve University

While the interchromosomal effect (IC effect) has been well characterized at the phenotypic level, almost nothing is known about the mechanism. Crossovers that occur during meiosis require a specific set of proteins to form and are subject to crossover control mechanisms such as interference (these are the so-called Class I COs). However, COs can also form using mitotic-like CO pathways, but these use a different set of proteins and are not subject to crossover control mechanisms. I will be determining if the extra COs that form during the IC effect are Class I meiotic COs or the mitotic-like COs. To do this, I’ll measure the IC effect in mei-9 mutants and blm mutants. Evidence suggests that mei9 is playing a critical role in resolving recombination intermediates into COs during meiotic recombination by cleaving double Holiday junctions into COs. Therefore, the mei9 mutant would not be able to produce class I crossover. By comparing the CO distribution of mei9 mutant and mei9;+/TM6B, the identity of extra crossovers from IC effect can be determined. Blm mutant would produce crossovers that lack chromosomal interference, so those crossovers found in blm mutants are considered to be class II crossovers. Therefore, if the blm mutants with balancer are scored, I can further insight about those extra crossovers from IC effect.

465V  Broad is sex and cell type specifically required in the Drosophila gonads for gametogenesis and fertility. PRADEEP BHASKAR, Brian Oliver NIDDK, NIH

Broad (Br, CG11491, FBgn0283451) is a BTB domain (Broad-Complex, Tramtrack and Bric a brac) containing protein that has a homodimerization domain at the N terminus and multiple copies of either zinc fingers of the C2H2 type or Kelch repeats. Br is best known as a modulator of the ecdysone-response and has been attributed to cause chromatin changes by puffing of chromosomes to affect gene expression. We examined Br expression and function in third instar larval gonads when ecdysone levels are high.

In our whole gonad RNA-seq experiments, we detected 11 out of 15 Br transcripts encoding 4 different protein isoforms of Br (Z1-Z4), on the basis of distinct Zinc-fingers at C-terminus. Further, Single Cell RNA sequencing (scSeq) and immunostaining with antibody show that, Br, was expressed only in somatic cells of testis and ovary. In the testis, Br expression was enriched in cyst cells that enclose the spermatogonia, which are mitotic germline cells near the apex, but not the cyst cells surrounding spermatocytes. Br was also expressed in terminal epithelium, which ultimately attaches to the reproductive tract derived from the genital disc, and the pigment cells that ensheath the testis. In ovary, we found Br expression in all somatic cells (Sheath cells, Terminal Filament, Cap cells, Intermingled cells, follicle cell and Swarm cells). To determine the function of Br in these cell types we knocked down all of its isoforms (by targeting common BTB domain) in the enclosing the germ cells (male cyst cells and female intermingled cells) using a traffic jam driver. This led to female-specific sterility due to germline defects including loss of germ cells. Br expression was lost from the spermatogonia cyst cells, but did not result in an overt phenotype. However, knockdown of Br using a doublesex driver resulted in sterility in both sexes. Knockdown of individual isoforms (RNAi against C-terminus) using doublesex driver results in testes that failed to elongate and attach to the reproductive tract, while in females egg retention phenotype was observed. Both males and females from these crosses were sterile. This indicates that Br is required in terminal epithelium to mediate attachment to the reproductive tract as well as egg laying in females and leads to sterility when Br is knocked down. Interestingly we found an enhancer that drives Br expression in both sexes, indicating dual role of enhancer in regulating Br function in sex and cell type specific way. In summary, we show that Br is required in both sexes, but in fundamentally different somatic cell types. This highlights the context-dependency of gene expression in sexual development and fertility.

466V  Nucleoporin107 mediates female sexual differentiation via Dsx Offer Gerlitz2, Tikva Shore1, Tgst Levi1, Rachel Kalifa 1, Amatzia Dreifuss 1, Dina Rekler1, Ariella Weinberg-Shukron2, Yuval Nevo 1, Tzofia Bialistoky 1, Victoria Moyal 1, Merav Yaffa Gold 1, Shira Leebhoff 1, David Zangen 1, Girish Deshpande 1 1) IMRIC, The Hebrew University- Faculty of Medicine; 2) Weizmann Institute of Science, Rehovot, Israel; 3) The Hadassah Hebrew University Medical Center, Jerusalem, Israel; 4) Division of Pediatric Endocrinology, Hadassah Hebrew University Medical Center, Jerusalem, Israel.; 5) Princeton University, Princeton, NJ, USA

We previously identified a missense mutation in the nucleoporin-107 (Nup107; D447N) as the cause for XX-ovarian-dysgenesis in five female cousins from a consanguineous family. All men in the family had normal pubertal development and those married have multiple children. Nup107 is an essential component of the nuclear pore complex expressed ubiquitously in a sex-non-specific manner. Modelling of the human mutation in Drosophila or specific knockdown of Nup107 in the gonadal soma resulted in ovarian-dysgenesis-like phenotypes, while male flies were fully fertile. To uncover the targets of Nup107 that mediate its sex and tissue specific activity, we conducted a genome-wide transcriptomic analysis on larval gonads compromised for Nup107. We identified 82 candidate genes which displayed significant changes in mRNA expression, including doublesex (dsx), the somatic sex-determination gene. Either loss or gain of Dsx in the gonadal soma is sufficient to mimic or rescue the phenotypes induced by Nup107 loss, establishing Dsx
as a primary relevant target of Nup107. Importantly, the aberrant phenotypes induced by compromising either Nup107 or dsx are reminiscent of BMP signaling hyperactivation. Remarkably, in this context, the metalloprotease AdamTS-A, a transcriptional target of both Dsx and Nup107, is necessary for the calibration of BMP signaling. We also uncovered the cellular impairments underlying BMP signaling dysregulation. As modulation of BMP signaling is a conserved critical determinant of soma-germline interaction, the sex and tissue specific deployment of Dsx-F by Nup107 seems crucial for the maintenance of the homeostatic balance between the germ cells and somatic gonadal cells.

467V  Tudor5-like promotes post-transcriptional regulation of maternal RNAs Caitlin Pozmantar, Sydney Kelly, Harrison Curnutte, Mark Van Doren Johns Hopkins University

Tudor-domain containing proteins are conserved across the animal kingdom for their necessary functions in germline development including post-transcriptional gene regulation. Recent work in our lab identified Tudor5-like (Tdrd5l), which promotes male germ line identity in germline stem cells (GSCs) in the testis, but is repressed by the RNA binding protein Sex lethal (SXL) in female GSCs. Interestingly, Tdrd5l is also expressed in the differentiating germline in both sexes, indicating that it may also act to control germline differentiation in both sexes. Previously we reported that Tdrd5l localizes to an RNA granule. To understand what RNA regulatory pathway Tdrd5l functions in, we conducted RNAi against the deadenylase twin in mutant gonads, which revealed a genetic interaction between Tdrd5l and the CCR4-NOT deadenyllation complex. Recent investigation into the role Tdrd5l plays in the female germline showed a decreased hatch rate and dorsal appendage defects in eggs laid by Tdrd5l mutants. Since Grk regulates dorsal appendage development we stained for Gurken(Grk) in Tdrd5l mutant ovaries and wild type ovaries. In wild type flies we see normal Grk translation in the anterior dorsal corner of the oocyte, while in tdrd5l mutant ovaries we see translation of Grk in the nurse cells. A similar nurse cell expression phenotype is observed when immunostaining for Orb in Tdrd5l mutants which is a known activator of Grk translation. This suggests the loss of maternal RNA regulation by Tdrd5l could result in patterning defects. Additionally, both osk mRNA and protein are mislocalized in Tdrd5l mutant oocytes where we observed them localized to the center of the oocyte instead of the posterior. One possibility is Grk fails to specify the posterior before migrating to the dorsal anterior corner. To investigate if this was due to failure to specify the posterior pole we used kin-lacZ and found in Tdrd5l mutants, kinesin does not strictly localize to the posterior pole of the oocyte in stage 9 or later egg chambers. Together our results indicate that Tdrd5l functions in regulating maternal RNA repression. To determine the mechanism by which this occurs we are currently conducting proximal biotinylation to identify other proteins in the Tdrd5l granule.

468V  The bHLH-PAS transcriptional complex Sim::Tgo plays active roles in late oogenesis to promote follicle maturation and ovulation Rebecca Oramas1, Elizabeth Knapp 1, Jianjun Sun1,2 1) Department of Physiology & Neurobiology, University of Connecticut; 2) Institute for Systems Genomics, University of Connecticut

Across species, ovulation is a process induced by a myriad of signaling cascades that ultimately results in activation of proteolytic enzymes and degradation of follicle cells to release encapsulated oocytes. In order for follicles to ovulate successfully, follicles need to become mature and gain ovulatory competency. Our previous work showed that upregulation of Zinc finger transcription factor Hindsight (Hnt) in follicle cells of stage-14 Drosophila egg chambers induces the expression of otopamine receptor in mushroom body (OAMB) in all follicle cells and matrix metalloproteinase 2 (MMP2) in posterior follicle cells, both of which are essential for ovulation. We also found that NR5A-family nuclear receptor Ftz-f1 is upregulated in stage-10 follicle cells and promotes follicle maturation via bHLH/ PAS-family transcription factor Single-minded (Sim). It is largely unclear how Sim, a critical player in central nervous system development, promote follicle maturation and ovulation. In this work, we discovered that Tango (Tgo), a class-II bHLH/PAS-family transcription factor and cofactor of Sim, is also expressed in follicle cells during late oogenesis. In addition, Tgo relocates to the nucleus when Sim is co-expressed in follicle cells from stage 10-12 and stage 14. Genetic manipulation suggests that Tgo is essential for follicle cell differentiation as Sim. Furthermore, we discovered that upregulation of Sim in stage-14 follicle cells is also essential for ovulation in addition to its role in stage 10-12 follicle cells with Tgo. This late Sim not only induces expression of Hnt, OAMB, and MMP2, but also NADPH oxidase (Nox), another factor involved in superoxide production and ovulation. Together, our work indicates that Sim and Tgo play multiple roles in late stage follicle cells to promote follicle maturation and ovulation.

469V  Explore the roles of steroid hormone signaling mediated Drosophila oogenesis Chueh Wen Wang, Anna C.-C. Jang Biotechnology and Bioindustry Sciences, National Cheng Kung University, Tainan, TW

Steroid hormone is temporal control in development process. The p160 steroid receptor coactivators (SRC) are shown being amplified and overexpressed in various cancers to promote proliferation and metastasis. However the molecular mechanism by which SRC induces dissemination of tumors has not been well established. Therefore, use border cells that are originated from follicle epithelial cells and invade through nurse cells that migrate to oocyte boundary in Drosophila oogenesis be study collective cell migration model. This migration process is also regulated by steroid hormone signaling and required SRC, named Taiman (Tai) in Drosophila. Tai interact with hormone receptor complex via LXXLL domain in a ligand dependent manner. Without any of these components, border cells fail to migrate and egg...
472C

Examining binding to sperm, although Hatch secretory cells (SSCs), or mutation of female molecules participate in facilitating SP's and other SFPs' binding to sperm. Ablation of the female's spermathecal to sperm, the effects of female factors on the timing of their (transient) sperm-binding differ. This, in turn, suggests that within females, suggesting that priming of sperm improves inside the female reproductive tract. Moreover, SP binding the female reproductive tract are male derived. We wished to know whether female factors (protein/non-protein) modulate post mating responses long-term. All factors currently known to account for binding of SP to sperm within proteins (LTR-SFPs) "prime" sperm to bind the seminal Sex Peptide (SP). This allows SP to persist in the female and needed for efficient fertility. These post-mating changes last for ~10-14 days, because some male-derived seminal "Sex Peptide" (SP) remains bound to sperm long-term, allowing it to be retained to modulate female behavior and physiology for ~10-14 days. To determine the generality of this response, we collaborated with the Dorus and Pitnick labs* to identify sperm-bound SFPs in an unbiased mass spectrometry screen. We are using RNAi to determine the fertility functions of sperm-bound SFPs detected in this screen. In addition, we are using two tagging methods (TAP, TurboID) to identify the binding targets of one of these SFPs (SP) on sperm.

*Whittington, Murphy, Singh, Pitnick, Wolfner, and Dorus, The life history of Drosophila sperm involves molecular continuity between male and female reproductive tracts. In review

470V

Functions and interactions of sperm-bound seminal proteins in Drosophila melanogaster Sarah Allen, Snigdha Misra, Mariana Wolfner Cornell University, Ithaca, NY

Sperm enter mating Drosophila females accompanied by secretions from the male reproductive tract. These secretions include seminal fluid proteins (SFPs) that modify the physiology of the female to enhance the fertility of the mated pair. While most seminal proteins are present transiently within the mated female, a few bind to sperm. In particular, the seminal “Sex Peptide” (SP) remains bound to sperm long-term, allowing it to be retained to modulate female behavior and physiology for ~10-14 days. To determine the generality of this response, we collaborated with the Dorus and Pitnick labs* to identify sperm-bound SFPs in an unbiased mass spectrometry screen. We are using RNAi to determine the fertility functions of sperm-bound SFPs detected in this screen. In addition, we are using two tagging methods (TAP, TurboID) to identify the binding targets of one of these SFPs (SP) on sperm.

471V

Female factors are important for the seminal Sex Peptide’s association with sperm, in mated D. melanogaster Snigdha Misra, Akanksha Singh, Mariana Wolfner 1) Cornell University; 2) National Heart Lung and Blood Institute, NIH, Bethesda, MD

Seminal fluid proteins (SFPs) induce a myriad of physiological and behavioral changes in mated female flies that are needed for efficient fertility. These post-mating changes last for ~10-14 days, because some male-derived seminal proteins (LTR-SFPs) “prime” sperm to bind the seminal Sex Peptide (SP). This allows SP to persist in the female and modulate post mating responses long-term. All factors currently known to account for binding of SP to sperm within the female reproductive tract are male derived. We wished to know whether female factors (protein/non-protein) also play roles in associating SP with sperm. We found that sperm in ejaculate bind SP much more weakly than sperm within females, suggesting that priming of sperm improves inside the female reproductive tract. Moreover, SP binding to sperm increases with time or transit within the female reproductive tract. Interestingly, while some LTR-Sfps also bind to sperm, the effects of female factors on the timing of their (transient) sperm-binding differ. This, in turn, suggests that female molecules participate in facilitating SP’s and other SFPs’ binding to sperm. Ablation of the female’s spermathecal secretory cells (SSCs), or mutation of Hr39, which leads to defective SSCs and/or parovaria, did not affect SP’s initial binding to sperm, although Hr39 mutant females retained more sperm and SP long-term.

472C

Examining essential functions of KDM5 via a novel truncation allele (kdm5Q19) Melissa Castiglione, Hayden Hatch, Julie Secombe, Andreas Bergmann 1) Albert Einstein College of Medicine, Bronx, NY; 2) UMass Chan Medical School, Worcester, MA

The lysine demethylase 5 (KDM5) family of transcriptional regulators are important for normal development, and their dysregulation is a key driver of intellectual disability and several forms of cancer. Most work to-date has focused on the histone demethylase activity of KDM5 proteins, which targets the active chromatin mark H3K4me3. However, KDM5 proteins can also regulate transcription through non-enzymatic mechanisms. While KDM5 is essential for development, its demethylase activity is not required, as is demonstrated by the viability of demethylase dead adult flies. In this work, we examine essential functions of KDM5 via a novel truncation allele, kdm5Q19, which does not alter demethylase activity. kdm5Q19 inserts a stop codon in a previously unrecognized, evolutionarily conserved, motif within an intrinsically disordered region of KDM5 at the C-terminus. kdm5Q19 animals do not survive to adulthood, which is distinct from null, demethylase dead, and other mutants generated in our lab, suggesting that the motif disrupted by the truncation has an essential as-yet-unknown role in normal KDM5 function. To further dissect the molecular activities of this region of KDM5, we generate additional alleles of kdm5 to refine the critical region(s) of the protein and assess viability and changes to transcription.

473A

Use of transformants bearing deletions in the 5’ upstream region of the Hdc gene to identify regions required
for CNS expression of Hdc Collin Louis1, Martin Burg1,2 1) Dept. of Biomedical Sciences, Grand Valley State University, Allendale, MI; 2) Dept. of Cell & Molecular, Grand Valley State University, Allendale, MI

The study of histamine and its function in Drosophila melanogaster has primarily been focused on its role as a neurotransmitter, both in photoreceptors and neurons of the CNS, leading to the identification of histamine’s role in a number of functions such as vision, grooming behavior, thermal preference, and sleep. Histidine decarboxylase is the enzyme that synthesizes histamine, and is encoded by the Hdc gene. Mutations in the Hdc gene have been identified that result in the inability to synthesize histamine in the entire fly. We have set out to determine how tissue-specific expression of Hdc could be regulated by creating deletions in the 5’ region of the Hdc gene where 3 distinct transcription start sites (TSSs) have been reported for Hdc. A 9.4 kb genomic Xba1 fragment was cloned into the P(CaSpeR-3) vector and used to generate germline transformants (P(CaSpeR3-Hdc9.4+)) in a mutant Hdc0930 background. Histamine immunostaining of the P(CaSpeR3-Hdc9.4+) transformant indicated that all CNS neurons contained histamine in both larvae and adults, suggesting that the Hdc gene was intact. A series of deletions, in ~300 bp steps, were generated in the 9.4 kb Xba1 Hdc genomic fragment from the 5’ end (~4.3 kb from the start of the HDC coding region) towards the coding sequence and transformed back into Hdc0930 mutant flies. Various P(CaSpeR3-Hdc9.4+) transformants were subjected to histamine immunodetection to determine which histaminergic cells in the CNS were no longer detected as each successive deletion step was taken in the transgene. Larval and adult stages were examined for alterations in the CNS and PNS pattern of histamine staining to correlate expression in specific cells with the removal of genomic regions containing predicted TSSs. Deletions that removed the Hdc-RD TSS had minimal effect in the tissues examined, while the removal of the TSS associated with the Hdc-RB, -RC isoforms appeared to cause loss of histamine staining in most of the CNS cells, while leaving PNS expression (photoreceptors) intact. Additionally, we have identified 3 consecutive deletions that disrupt expression in single pairs of neurons that span between the Hdc-RD and Hdc-RB, -RC isoform TSSs. Elimination of these specific regions in the 5’ region of the Hdc gene do have incremental effects on Hdc expression, suggesting that some of the TSSs identified are tissue-specific. These transgenic deletion-bearing flies could be useful as tools for examining the contribution of some histaminergic CNS neurons to a variety of behaviors ascribed to histamine in the CNS as photoreceptor expression appears to be retained as the Hdc-RA TSS was not disrupted.

References:
1. Oh et al., 2013, PLoS ONE 8(7):e68269
3. Larkin, A. et al. (2021), Nucleic Acids Res. 49(D1) D899–D907
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474B Investigating the role of intrinsic protein disorder in transcription factor dynamics and function Colleen Hannon1, Michael Eisen1,2 1) University of California, Berkeley, Berkeley, CA; 2) Howard Hughes Medical Institute, Berkeley, CA

Modern microscopy has revealed that core nuclear functions, including transcription, replication, and heterochromatin formation occur in spatially restricted clusters. Our lab is investigating the role that the sub-nuclear localization of transcription factors, target DNA sequences and transcriptional machinery plays in regulating RNA synthesis in the early Drosophila embryo. A nearly ubiquitous feature of eukaryotic transcription factors (TFs) is tracts of low complexity amino acid sequence, which result in intrinsically disordered regions (IDRs) within the protein. It has been proposed IDR-containing TFs mediate the co-localization of transcriptional machinery and target genes. However relatively little is known about the contribution of IDRs to TF localization, molecular interactions, and transcriptional activation. Using a novel algorithm to identify IDRs in the Drosophila proteome, we designed a library of IDRs from TFs expressed in the early Drosophila embryo. We used this library to conduct a broad survey of the nuclear sub-localization of TFs, using a high throughput imaging screen in Drosophila S2 cells. We found that while subnuclear clustering does not occur when the IDRs are expressed alone, it is frequently seen in full length TFs. These results are consistent in Drosophila embryos, suggesting that IDRs are insufficient to drive the sub-nuclear clustering behavior of transcription factors.

475C Nuclear Function of the protocadherin fat in Drosophila Janette Rusch1, Chikin Kuok2, Joe Thanintorn1, Yonit Tsatskis1, Helen McNeill1 1) Washington University St Louis School of Medicine; 2) Lunenfeld-Tanenbaum Research Institute, Mount Sinai Hospital, Toronto, Canada

The conserved atypical cadherin fat (ft) controls cellular processes such as growth control via the Hippo pathway, planar cell polarity, and mitochondrial function, in organisms ranging from fruit flies to mammals. The intracellular domain of the Ft protein, FtICD, regulates a variety of partners to execute these functions. Our lab has found that FtICD is present in the nucleus in tissue culture cells, and we have identified both nuclear localization and nuclear export signals in Ft. Moreover, a membrane-bound version of FtICD, fused to a Gal4VP16 transactivation domain, is able to activate a reporter construct in imaginal discs, demonstrating that FtICD can be cleaved and enter the nucleus in vivo. These observations suggested that ft may have a nuclear function in addition to its cellular signaling functions. ChIP experiments on Drosophila larval tissues identified putative ft targets, including the anti-apoptotic gene, Diap1, which is also a target of the Hippo pathway. Analysis of a small fragment of the Diap1 enhancer, HRE (Hippo Response
Element), in vivo suggests that FtICD can activate this element, further supporting the novel notion of a nuclear function for Ft. Remarkably, FtICD ChiPs to Yorkie sites across the genome. We hypothesize that the nuclear function of Ft functions to modulate Hippo pathway activity, complementing its established function as an upstream regulator of Hippo signaling.

476A Establishing the Role of the Conserved TN Domain in Tinman Cayleen Bileckyj, Richard Cripps San Diego State University

Congenital heart disease (CHD) is a major factor in mortality and morbidity in children and adults. Even though there has been substantial progress in detection and treatment, not all congenital heart defects can be identified early on through physical screenings. To advance our abilities to identify and manage CHD, we need to further understand the key genetic factors involved in causing these maladies. Since Drosophila melanogaster shares similar cardiac developmental mechanisms with humans, it has been an essential model for human heart development. One similarity occurs between Tinman (Tin), a transcription factor in Drosophila necessary for the differentiation of cardiac cells, and its mammalian ortholog NK2 Homeobox 5 (Nkx2.5). These proteins share two conserved regions: the homeobox domain and the tin (TN) domain. Although the TN domain is completely conserved between these two proteins, there is little known about its significance. By utilizing CRISPR/Cas9 gene editing, I established a line of Drosophila containing an in-frame deletion of the TN domain. Staining Drosophila embryos for NMR, a marker of cardinal cells, revealed mutant embryos generate a significantly higher number of cardinal cells when compared to wild type embryos (p<0.01). I also stained embryos for Tin and Svp, a protein important for heart development in Drosophila, to determine if the deletion of the TN domain affected cardinal cell specification. Mutant embryos contained more cells expressing tin and more cells expressing svp than wild type embryos. These results support the importance of the TN domain in heart development and indicate the increase in cardinal cells occurs before cardinal cells differentiate into cells exclusively expressing tin or svp. Tin and Nkx2.5 are vital to proper heart development. Characterizing the role of the conserved TN domain in these transcription factors provides the opportunity to vastly improve our understanding of the mechanisms involved in cardiac formation and maturation.

477B Initiating and Maintaining the Histone Locus Body: Two Sides of the Same Coin? Greg Kimmerer, Leila Rieder Emory University

Histone proteins play an essential role in nearly all genomic activities in eukaryotes, from gene regulation to DNA damage repair. In order to perform these functions, the ratio of DNA to histone levels must be kept constant, as deviations from this ratio cause cell cycle arrest and genomic instability. Consequently, a cell must double its histone content during each cell cycle. To meet these regulatory requirements, the histone genes are targeted by a conserved pantheon of chromatin proteins, known collectively as the Histone Locus Body (HLB). Most of the dozens of HLB components have no DNA-binding activity, yet they are able to faithfully locate the histone genes in the early embryo. The HLB must also be maintained at histone genes in adult tissues. How the HLB is initiated in the early embryo and maintained in the adult is a crucial gap in our knowledge of how proper histone gene regulation is achieved. Recently, our lab discovered that a single regulatory element located in the histone3/histone4 bidirectional promoter directs HLB initiation along the entire histone array. However, it is not clear if this cis-element, which consists of GA-dinucleotide repeats, is required to maintain the HLB after initiation. To answer this question, I have designed and cloned a novel transgene that takes advantage of the FLP/FRT recombination system in Drosophila. This transgene allows me to wait for the HLB to initiate in the embryo, then heat shock larvae to induce recombination, removing the GA-repeats. If the ectopic HLB is lost following heat shock, this will suggest that the GA-repeats plays an active role in maintaining the HLB. If the ectopic HLB survives heat shock, this will show that the HLB can be maintained even without the cis-element that initiated it, an equally illuminating result.

478C Developmental regulation of histone genes by pioneer factor Zelda Thomas O’Haren, Leila Rieder Emory University, Atlanta, GA

The zygotic genome is mostly unstructured and quiescent, so embryos are maternally loaded with proteins and RNAs that control early development. However, at some point, the zygotic genome must activate and begin transcribing necessary genes to assist in the early cellular divisions, a process known as zygotic genome activation (ZGA). The initial organization and transcriptional activation of the zygotic genome is facilitated by pioneering transcription factors that are able to bind and open regions of nucleosome-bound DNA. The histone proteins, which are critical to chromatin organization and regulation of the cell cycle, are some of the earliest expressed proteins from the zygotic genome. The five canonical, replication-dependent histone genes cluster in the genomes of metazoans, forming histone loci. A suite of factors known as the Histone Locus Body (HLB) act upon histone loci and regulate histone expression. The genome of Drosophila melanogaster carries a single histone locus of 107 tandem copies of the 5kb histone gene array. It remains unclear how the histone locus is activated and the HLB is established and maintained. The pioneer factor Zelda, a maternally deposited, zinc-finger protein, is the “master regulator” of ZGA and binds across the genome to “TAGteam”
sites, opening chromatin and marking surrounding regions for activation and transcription. We discovered the Zelda targets TAGteam sites in the histone locus as early as embryonic nuclear cycle 8. A histone array transgene in which the TAGteam sites are ablated fails to attract HLB factors suggesting that Zelda is necessary for HLB formation in the early embryo and is involved in activating zygotic histone gene expression. In the future, we will restore Zelda to the transgenic histone array using dCas9 and assay rescue of HLB formation. Overall, our results indicate that one indirect mechanism through which pioneer factors remodel the zygotic genome is through zygotic histone gene expression.

479A  Fruitless modulates the threshold of Notch target gene transcription during asymmetric neuroblast division Arjun Rajan1, Lucas Anhezini1, Megan Neville2, Elizabeth Larson3, Stephen Goodwin2, Melissa Harrison3, Cheng-Yu Lee1,4,5 1) Life Sciences Institute, University of Michigan, Ann Arbor, MI; 2) Centre for Neural Circuits and Behaviour, University of Oxford, Oxford, UK; 3) Department of Biomolecular Chemistry, University of Wisconsin, Madison, WI; 4) Department of Cell and Developmental Biology, University of Michigan Medical School, Ann Arbor, MI, 48109; 5) Division of Molecular Medicine and Genetics, Department of Internal Medicine, University of Michigan Medical School, Ann Arbor, MI

A defined threshold of gene transcription prevents the accumulation of excess transcripts and proteins, allowing cells undergoing developmental transitions to rapidly rewire their gene regulatory circuits and assume new cell identities. The delay negative feedback loop that maintains the oscillatory pattern of Notch target gene expression in vertebrates exemplifies how post-transcriptional regulatory mechanisms function as a timer to indirectly regulate gene identities. The delay negative feedback loop that maintains the oscillatory pattern of Notch target gene expression in cells undergoing developmental transitions to rapidly rewire their gene regulatory circuits and assume new cell identities. We investigated if repressive inputs could function in synchrony with transcriptional activators to modulate gene transcription thresholds in fly larval brain type II neuroblasts, which divide asymmetrically to generate a NotchON neuroblast and a NotchOFF differentiating progeny. We hypothesize that factors eliciting repressive inputs should (1) be specifically expressed in type II neuroblasts, (2) bind Notch target gene loci and (3) repress Notch target gene expression. We performed single cell RNA-sequencing to establish a transcriptomic atlas of a wild-type type II neuroblast lineage that includes neuroblast progeny undergoing commitment to differentiation. We discovered that fruitless (fru) gene transcripts are uniquely detected in type II neuroblasts, and the C isoform of Fru (FruC) is specifically expressed in type II neuroblasts. Loss of fru function enhanced the supernumerary neuroblast phenotype in numb- or brat-hypomorphic brains in which reduced function of either gene leads to aberrant upregulation of Notch signaling in neuroblast progeny driving their reversion. We performed CUT&RUN to profile FruC-bound loci in type II neuroblast enriched brains and found that FruC binds most Notch target gene loci in the fly genome. These data suggest that FruC functions to antagonize Notch target gene expression in type II neuroblasts. Consistent with this model, reducing fru function increases Notch target gene expression levels in mitotic type II neuroblasts. Furthermore, overexpressing wild-type FruC or a constitutive repressor form of FruC is sufficient to downregulate Notch signaling and partially restore differentiation in brat-null brains. We propose that the balance between FruC-mediated chromatin modifications and Notch-induced histone acetylation specifies Notch target gene transcription thresholds in type II neuroblasts.

480B  Out of the shadows: Co-acting cis-regulatory elements control T-box transcription factors midline and H15 during development. Cody Stevens, Nir Yakoby Rutgers University Camden

Tissue development involves highly coordinated expression of transcription factors controlled by complex cis-regulatory elements (CREs). The posterior end of the follicular epithelium is patterned by Midline (MID), the Drosophila homolog of the mammalian Tbx-20 transcription factor. Here, we identify two CREs that recapitulate the endogenous pattern of mid. It has been genetically shown that mid is negatively regulated by the Iroquois transcription factor mirror and the bone morphogenetic protein signaling pathway, while positively activated by the ETS transcription factor pointed. In this study, we corroborate these trans regulators through cis regulation of mid, highlighting regulatory regions within the CREs. Furthermore, through CRISPR/Cas9 genome editing, we illustrate a redundancy of mid CREs. However, deletion of both CREs resulted in a significant reduction, but not loss of mid, indicating additional CREs are possible. Interestingly, unlike the published trans regulatory connection between MID and a second Tbx-20 homolog H15, we found co-regulation between one mid CRE and H15, whereas deletion of this CRE eliminated H15 transcription. These results suggest that both T-box transcription factors mid and H15 share regulatory capacity that may act as safeguards for determination of posterior fate during egg development.

481C  Necessity versus sufficiency: furthering understanding of ftz cis-regulatory elements in Drosophila melanogaster Matthew Fischer, Patricia Graham, Leslie Pick University of Maryland, College Park

Spatiotemporal regulation of transcription is essential for directing complex patterns of gene expression during embryogenesis. This is achieved through the action of cis-regulatory elements (CREs), which have been studied extensively in Drosophila with reporter genes that permitted the in vivo dissection of CREs. How these CREs dynamically interact with each other and specific promoters in the context of native chromatin remain as unresolved questions. The Drosophila Hox complex includes the pair-rule gene fushi tarazu (ftz) nested between the homeotic genes Antennapedia and Sex combs reduced (Scr). Interestingly, ftz and Scr CREs are intermixed, but the genes have
distinct expression patterns: *ftz* is expressed in seven stripes during the blastoderm stage while *Scr* is expressed later in the embryonic labial and prothoracic segments. Our investigation is twofold; one, to assess the necessity of previously identified *ftz* CREs in the native context of the genome using CRISPR-Cas9 induced deletions; two, to identify the location of potential structural elements in this genomic region through the use of reporter transgenes. To this end, we independently deleted the zebra and upstream CREs, two classic seven-stripe *ftz* CREs, from the endogenous context of the *ftz* locus using CRISPR/Cas9. These deletion mutants are homozygous viable and fertile, contrary to expectations from historic rescue experiments. While some of these homozygotes develop all segments, other progeny develop abdominal abnormalities including specific segmental deletions and/or fusions. Quantification of the *ftz* signal using in situ hybridization chain reaction showed that *ftz* expression is reduced to 25% of wildtype in stripe four of the zebra deletion mutant. This refines the lower threshold of *ftz* transcript that is both necessary and sufficient for segmentation to proceed. Similar deletions have been made of several stripe-specific CREs, and these homozygotes are viable and fertile. Their survival demonstrates redundancy of multiple CREs directing *ftz* expression. Analysis of reporter transgenes generated in both orientations has identified two putative insulators. These regions will be targeted for deletion by CRISPR-Cas9 to determine if removing them causes crosstalk between *ftz* and *Scr*. Overall, this work provides insight into how insulators organize specific enhancer-promoter interactions in complex genomic regions, allowing for precise spatiotemporal regulation.

### 482A Defining the mechanisms underlying how enhancer binding sites regulate Notch signal strength

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Notch signaling regulates distinct cellular fates and processes during the development of nearly every organ in the body. Moreover, Notch-dependent processes are often highly sensitive to Notch signal strength as evidenced by genetic haploinsufficiency across multiple species. For example, decreases in Notch signal strength have been associated with developmental syndromes such as Alagille Syndrome and Adams Oliver Syndrome. However, we still do not fully understand how changes in Notch signal strength affect cell-specific responses. In general, Notch signal strength is determined by ligand induced production of the Notch Intracellular Domain (NICD), which forms a complex with the CSL transcription factor, Suppressor of Hairless (Su(H)) in flies, to induce gene expression via two types of binding sites: monomeric (CSL) and dimeric (Su(H) paired sites, SPS). Unexpectedly, we found that simply inserting synthetic enhancers that couple dimeric SPSs, but not monomeric CSL sites, with binding sites for the Grainyhead pioneer transcription factor resulted in flies that develop wing nicks, a classic Notch-sensitive phenotype. Moreover, this enhancer induced phenotype is highly sensitive to genetic changes in both the Notch pathway and the Cdk8-Mediator complex, which has been previously shown to promote NICD turnover. These findings suggest that Notch dimers preferentially promote NICD turnover and that Grainyhead can aid this process by promoting chromatin accessibility. To test this model, we generated a series of enhancers that couple SPS sites with binding sites for other pioneer transcription factors or transcription factors not known to have pioneer activity. Importantly, we found that pioneer factors, such as Zelda and Trithorax-like, could produce wing nicks when placed next to SPS sites, whereas non-pioneer sites such as E-box and UAS sites failed to do so. Our results also revealed that some, but not all, endogenous dimeric SPSs produced wing nicks, suggesting that other factors influence this activity. Genetic dosage experiments in *Drosophila* further revealed that removing a copy of each of the Cdk8-Mediator components significantly reduced wing nicking. Ongoing studies are focused on testing whether Grainyhead promotes the opening of these synthetic enhancers and on identifying the additional sequences and factors that allow for a subset of SPSs to induce wing nicking phenotypes.

### 483B *brinker* gene promoter-proximal element drives ovary expression and supports sequential action of distal enhancers

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Limiting BMP signaling range in the stem cell niche of the ovary protects against germ cell tumors and supports germline homeostasis. In both the embryo and wing disc, the Brinker (Brk) transcription factor serves as the canonical repressor of Dpp, the main *Drosophila* BMP ligand, and its downstream targets. In the germline stem cell niche within the gerarium of the ovary, Dpp provides a pivotal cue for maintaining stem cell identity. Despite this well-established role, Brk’s function in the gerarium has not previously been described. Here we find that *brk* expression throughout the ovary requires a promoter-proximal element (PPE), a DNA region upstream of and distinct from the gene promoter. In the gerarium, the PPE acts as an enhancer to regulate *brk* expression levels to support germline stem cell homeostasis and, surprisingly, to positively regulate Brk’s canonical antagonist, *dpp*. This PPE has been previously described in the embryo where it does not itself drive expression, but is required for the sequential action of two distal enhancers to support continuous *brk* expression. In the egg chambers of the ovariole, at least two enhancers perform a similar temporal handoff and their activities require the PPE, indicating that this element may serve a general function of managing gene regulation by multiple enhancers over time by coordinating shifts in local chromatin conformation.

### 484C Investigating the genome-wide cooperativity between the pioneer factor Zelda and patterning transcription factors in the early embryo

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#Dros22 | 154
Enhancers, or cis-regulatory DNA sequences that control spatiotemporal gene expression programs, have intrinsically high nucleosome barriers that prevent the nonspecific binding of transcription factors (TFs). For an enhancer to become active, TFs must overcome this barrier, access their binding motifs, and evict the nucleosome, but the mechanisms by which these steps occur are not clear. Current models suggest that specialized TFs called pioneer factors can access their motifs in the presence of nucleosomes and foment nucleosome depletion through cooperativity with additional TFs. Even still, how pioneer and non-pioneer TFs cooperate to generate chromatin accessibility at enhancers is not yet known. To understand how TFs overcome the nucleosome barrier, we are using Zelika, a pioneer TF for the Drosophila maternal-to-zyotic transition, and a set of TFs that drive pattern formation in the early embryo (patterning TFs) as a model system. To dissect Zelika’s cooperativity with the patterning TFs, we have integrated ChIP-nexus TF binding data with our novel deep learning model, BPNet, which learns the relationship between DNA sequence and TF binding in an inherently combinatorial way. BPNet reliably predicts the genome-wide binding of Zelika and the patterning TFs, and successfully discovers known and novel motifs at well-studied enhancers. Through employing the trained BPNet model as an in silico oracle, we reveal that Zelika and the patterning TFs exert directional cooperativity within nucleosome-range distances. Using time course ATAC-seq measurements, we find evidence that Zelika alone is insufficient to generate accessibility, rather chromatin accessibility in the early embryo relies on cooperativity between Zelika and the patterning TFs. Taken together, these results suggest a nucleosomal basis for cooperativity between Zelika and the patterning TFs and present a model where Zelika establishes a more permissive chromatin state for TF binding but relies on cooperative interactions with the patterning TFs to bestow accessibility. To further dissect the hierarchical nature of TF cooperativity for chromatin accessibility, we are training a BPNet model on our time course ATAC-seq data, which will learn the relationship between DNA sequence and accessibility and will identify the sequence features that are predictive of chromatin accessibility.

485A  Enhancer hijacking leads to flies with no thorax Taylor Crawford, Anna Horacek, Victoria Blake, Judith Kassis NICHD/NIH

The invected (inv) and engrailed (en) genes are expressed throughout development in Drosophila melanogaster. Distinct enhancers drive the co-expression of inv and en in discrete parts of the embryos including in stripes, parts of the head, and the CNS. Inv/En are also co-expressed in the anterior/posterior compartment of larval imaginal discs. The inv and en promoters are separated by 154 kb and their expression is regulated by enhancers distributed across a 100 kb region. One enhancer that regulates expression in imaginal discs is located 40kb and 90kb upstream of the en and inv promoters. How does this imaginal disc enhancer find the en/inv promoters? Previous results from our lab suggested that there is a promoter-proximal tethering element (PTE) that facilitates this activation. We are interested in understanding how this PTE functions. Our lab isolated a transgenic line containing a 2 kb en regulatory fragment, including the PTE, fused to a Beta-galactosidase reporter gene, in a P-element vector including the mini-white reporter gene, inserted 250bp upstream of the endogenous en promoter. Flies heterozygous for the transgene show expression of Beta-galactosidase and en in the posterior compartment of the wing imaginal discs. However, homozygotes show a reduction in en expression in wing imaginal and die as pharate adults with no thorax. These results indicate that the imaginal disc enhancers are hijacked by the transgene, causing a tissue-specific loss-of-function phenotype. We are investigating which sequences are required to mediate enhancer hijacking by the transgene using the Cre-loxP recombination system to insert various transgenes 250bp upstream of the en promoter. We have identified a 181bp DNA fragment that, when combined with a minimal promoter, captures imaginal disc enhancers and disrupts en expression. Interestingly, this fragment also acts as a Polycrumb response element (PRE) in which it can recruit Polycrumb group proteins. In addition to identifying sequences necessary for enhancer-promoter communication, our study shows that interfering with promoter-enhancer communication is another way to generate a tissue-specific mutant phenotype.

486B  Investigating the role of Notch signalling in the development of the ventral mesoderm in Drosophila melanogaster Marvel Megaly, Gregory Foran, Aleksander Necakov Brock University

Notch signaling is a critical regulator of multiple developmental processes through its ability to control gene expression, and thereby influence cell fate specification and cell proliferation through direct cell-cell communication. Notch signaling is activated through binding of the transmembrane ligand, Delta, to the Notch extracellular domain (NECD) of the transmembrane Notch receptor. Ligand engagement and trans-endocytosis into the signal-sending cell drives removal of the labile NECD, and concomitant cleavage of the Notch intracellular domain (NICD). Upon subsequent translocation to the nucleus, the NICD regulates target gene expression through its interaction with Suppressor of Hairless Su(H). Although Notch signaling has been shown to play a role in regulating single minded (Sim) expression in the embryonic mesectoderm, Notch activity and function in the directly-adjacent cells of the ventral mesoderm remain unknown. Considering that Delta endocytosis and NECD trans-endocytosis are required for Notch signal activation, and are restricted to the ventral mesoderm in the early embryo, we investigated the role of Notch signaling in the ventral mesoderm. To test our hypothesis, we have used a combination of Optogenetics, quantitative RNA fluorescent in situ hybridization (FISH), and qPCR. Through in silico analysis, we identified 13 potential Notch target genes based on the following three criteria: 1) expression in the embryonic ventral mesoderm, 2) promoter-resident Su(H) binding site(s),
and 3) differential expression during ventral mesoderm formation. We have validated these candidate Notch target genes, which include WntD, Asph, Heartless, Traf4, Tinman, Stumps, Mef2, Mes2, mir-1, Neurotactin, and NetrinA, by comparing the level of gene expression between loss-of-function Notch mutants and wild-type embryos through two orthogonal techniques; FISH and qPCR. Consistent with a role for Notch signaling, expression of these mesoderm-specific genes is reliant upon Notch. These results prompted us to ask whether Notch signaling is sufficient to drive expression of these mesodermal target genes. To address this, we have developed and validated a set of novel Optogenetic tools to ectopically activate Notch signaling in a precise spatio-temporal manner. Consistent with previous findings, we observed that ectopic activation of Notch signaling is sufficient to drive Sim, a Notch target gene normally restricted to the mesectoderm. Interestingly, we have observed a stage-dependent expansion of Sim expression into both the ectoderm and the ventral mesoderm of the early embryo, along with a change in the position of the ventral mesodermal boundary, demonstrating an effect of precocious Notch activity on embryonic patterning.

487C The synergistic roles of Glass and EGFR signaling in the differentiation of multiple retinal cell types Hongsu Wang1, Raja Komal2, Kelvin Yeung2, Graeme Mardon2, Jessica Treisman1 1) Skirball Institute for Biomolecular Medicine and Department of Cell Biology, NYU School of Medicine; 2) Department of Pathology and Immunology and Department of Molecular and Human Genetics, Baylor College of Medicine

Light-detecting photoreceptors and supportive cone cells differentiate from identical precursor cells in the eye disc during late larval and early pupal stages. Reiterative Epidermal Growth Factor Receptor (EGFR) signaling induces the specification of photoreceptors R1-7 and cone cells. Glass (Gl) is an eye-specific transcription factor that is necessary for the differentiation of all cell types in the retina. Gl is expressed in all cells posterior to the morphogenetic furrow, but only turns on photoreceptor-specific genes in cells that receive an EGFR signal. We hypothesized that the cell-intrinsic transcription factor Gl and the external EGFR signal synergistically induce accurate and robust differentiation. Misexpression of Gl in the wing disc leads to the activation of some eye-specific genes, but only coexpression of Gl with an activated form of the EGFR signaling component Ras (RasV12) could induce expression of the neuronal markers Elav and Futsch. We used RNA-Seq to identify genes that are synergistically induced by Gl and RasV12 in the wing disc and to determine which of them are dependent on the presence of the transcription factor Pointed (Pnt), which acts downstream of Ras. A large proportion of these Pnt-dependent synergistically induced genes are characteristic of neurons. We are using Targeted Dam ID with Gl and PntP1 to determine which of these genes are direct targets of Gl and/or EGFR signaling. Alternatively, transcription factors that are regulated by the combination of Gl and RasV12 could mediate their downstream effects, and we are testing several candidates. Single-cell RNA-Seq revealed that gl mutant eye discs lack both the distinctive differentiation trajectory profile of photoreceptors R2, 5, 3, and 4 present in wild type white prepupal eye discs (accuracy of differentiation) and any clusters representing R1, 6, or 7 (robustness of differentiation). Studying the synergy between Gl and EGFR signaling will help us understand how the cell intrinsic factor Gl cooperates with external signaling to induce accurate and robust differentiation of key retinal cell types.

488A Tissue-specific diversity of the Muscleblind expression in adult flies Davron Hanley Kennesaw State University

The muscleblind (mbl) family of RNA-binding proteins regulates alternative splicing, determining mRNA transcript composition for various types of tissue, and has been implicated in myotonic dystrophy. The mbl gene is subject to alternative splicing in Drosophila, leading to multiple isoforms, and has several paralogs in humans. Mbl proteins vary significantly in length, although the significance of such diversity and the role of specific isoforms have not been fully explored. Using immunofluorescence microscopy and polyclonal serum, we analyzed Mbl protein expression across adult Drosophila tissues. Mbl was detected in various locations, including the brain, gonads, muscle, and gut epithelium. Skeletal muscles demonstrated the greatest diversity in Mbl expression, with other tissues showing more homogenous expression. Mbl was present at low levels in flight and jump muscles, while other thoracic muscles and abdominal muscles showed high Mbl levels. Intracellular localization of Mbl was typically nuclear, however in the nervous system the protein was strongly expressed in the cytoplasm. During early adult development in the pupa, in various tissues, Mbl was initially detected in discrete nuclear bodies, before it assumed more general nuclear staining. Our study reveals natural locations that have drastically different levels of Mbl as well as its intracellular localization. Our further aim is to supplement these findings with molecular analysis of Mbl isoforms to create the basis for functional interrogation of the Mbl diversity and its relation to tissue-specific regulation of alternative RNA splicing.

489B Lingerer interact with FMRP to promote FMRP target translation KAICHENG MA, Al Rohet Hossain, Kayla Judson, Ethan Greenblatt The University of British Columbia

Fmr1 encodes an RNA binding protein FMRP necessary for proper ovary and neuronal development. Mutations in Fmr1 lead to fragile X syndrome, the most commonly inherited intellectual disability (ID) autism-associated disorder. By studying Fmr1’s role in transcriptionally silent mature Drosophila oocytes, we recently discovered that Fmr1 functions primarily to promote the translation initiation of genes encoding large proteins, many of which are orthologs of human genes associated with ID and autism.
In order to identify FMRP binding partners and elucidate the mechanism of Fmr1-dependent translation, we used CRISPR to tag Fmr1 at its endogenous locus and performed IP-mass spectrometry experiments from ovarian extracts. We also determined which FMRP binding partners are required for its activity by using a novel single molecule fluorescence in situ hybridization (smFISH)-based assay, in order to assess Fmr1’s activity in the absence of its binding partners.

Our preliminary data identify a critical role for the RNA and ubiquitin binding protein lingerer (lig). We find that lig physically associates with FMRP, and the loss of lig appears to strongly diminishes FMRP’s activity towards one of its targets. Previous studies of the mammalian lig ortholog UBAP2L in cultured cells have shown that it can bind to ribosomes to promote translation. In flies, lig is required for proper copulation behavior and short-term memory. Our data indicate that lig acts in vivo with FMRP to promote the production of FMRP targets, a function which is likely to be critical for proper neuronal and ovarian function.

490C Development of a novel molecular assay to sensitively detect Fmr1’s translational function in Drosophila ovarian follicles. Kayla Judson, Al Rohet Hossain, Kaicheng Ma Department of Biochemistry and Molecular Biology, UBC, Vancouver, BC

Loss of function mutations in the Fragile X Mental Retardation 1 (Fmr1) gene are associated with fragile X syndrome and fragile X-associated primary ovarian insufficiency, leading causes of autism and female infertility respectively. Although Fmr1 has been highly studied, its biological role is still subject to debate. A major challenge in studying Fmr1 is that its molecular function is to weakly promote the translation of its targets ~2-fold, making it difficult to assess its activity in vivo. Here we present a novel assay to detect Fmr1’s function at a molecular level with high sensitivity in developing Drosophila ovarian follicles.

Using single molecule fluorescence in situ hybridization (smFISH), we observed that the mRNA of the Fmr1 target Poe forms microscopically visible granules that depend on ongoing translation. These “Poe particles” appear to be sites of Fmr1 translational activation, as Fmr1, Poe mRNA, and fully translated Poe protein are each enriched in Poe granules. Fmr1 knockdown, or treatments that reduce translation (puromycin or heat shock) resulted in a robust dispersal in Poe mRNA granules. Using the presence of Poe particles as a sensitive readout of Fmr1 activity, we conducted a genetic screen to identify potential partners of Fmr1, by labeling Poe mRNAs in either wild type egg chambers or those lacking Fmr1 or candidate genes. We screened over a hundred different candidate genes using germline-specific RNAi. We found that only a few RNAi lines targeting RNA binding proteins and/or proteins which physically bind to Fmr1 exhibited a strong reduction in Poe particles. Our findings suggest that Fmr1 functions together with a small set of additional factors to promote the expression of genes essential for normal ovarian and neuronal function.

491A Precocious expression of Zelda does not initiate early zygotic genome activation Elizabeth Larson1; Zoe Fitzpatrick1, Hideyuki Komori2, Cheng-Yu Lee2, Melissa Harrison1 1) Department of Biomolecular Chemistry, University of Wisconsin School of Medicine and Public Health, Madison, WI; 2) Department of Cell and Developmental Biology and Life Sciences Institute, University of Michigan, Ann Arbor, MI

During the first stages of development, the fertilized germ cells rapidly transition to totipotency. Maternally deposited mRNAs encode the proteins necessary for reprogramming the transcriptionally quiescent zygotic genome during this maternal-to-zygotic transition (MZT). The transcription factor Zelda is essential for this reprogramming in the Drosophila embryo. Zelda is necessary for transcriptional activation of the zygotic genome, and the absence of Zelda leads to embryonic lethality during the MZT. Excess Zelda activity is also lethal to the embryo, demonstrating that Zelda levels must be precisely controlled during early development. Because Zelda is encoded by a maternally deposited mRNA, Zelda levels in the embryo are controlled at the level of translation. To understand how levels of this essential reprogramming factor were regulated to allow for embryonic development and zygotic genome activation, we investigated the factors that regulated translation of zelda. Brain Tumor (BRAT) is a translational regulator that was previously shown to bind to zelda mRNA in the embryo. We showed that BRAT functions to repress zelda translation, as embryos deficient for maternal BRAT activity prematurely express Zelda. We further showed that in the larval brain, BRAT similarly regulates Zelda levels and identified specific BRAT-binding sites that mediate these effects. Thus, BRAT regulates Zelda in multiple tissues. Because both too little and too much Zelda are lethal to the embryo, we hypothesized that precocious expression of this transcriptional activator might be capable of driving precocious activation of the zygotic genome, leading to embryonic lethality. To test this hypothesis, we performed single embryo RNA-seq at distinct nuclear cycles throughout zygotic genome activation (NC10, NC12, NC13, and NC14) in control and brat-mutant embryos. Our results conclusively demonstrated that in embryos lacking functional BRAT, Zelda target genes were not prematurely
activated. Rather, these genes were expressed normally, but become significantly upregulated at nuclear cycle 14, when the division cycle slows. Our data support a model in which zygotic genome activation requires precise coordination between expression of reprogramming factors, such as Zelda, and the slowing of the cell cycle.

492B  Tet (Ten-Eleven Translocation) Regulates Axonal Development in the Drosophila Pupal Brain via Transcriptional Repression Hiep Tran, Badri Nath Singh, Joseph Kramer, Ruth Steward Waksman Institute, Rutgers University

Tet (ten-eleven translocation) is one of the most important epigenetic regulators. In vertebrates, TET1/2/3 control demethylation DNA to activate gene transcription. Both vertebrates and Drosophila Tet can also hydroxymethylate cytosine in mRNA (5hmC). Our recent results show that Drosophila Tet is essential for viability and that the protein plays crucial roles in brain and muscle development. Further Tet binds to 2,500 genes and appears to deposit the 5hmC modification to their mRNAs thereby enhancing their translation levels. In this study, we found that Tet is required for mushroom body axon outgrowth in the pupal brain. Lack of Tet resulted in β lobe axon crossing the brain midline. By comparing CRISPR/Cas9-induced point mutations in the Tet DNA binding domain (TetαXXC) and catalytic domains (TetγRA), we discovered that in the DNA binding domain mutant TetαXXC the β lobe axons crossed the brain midline. This phenotype was not observed in the catalytic domain mutant TetγRA. Surprisingly, in the DNA binding domain mutant, the expression of 1,597 genes in developing pupal brains were increased while the catalytic domain mutant only showed 28 genes up- and 16 genes down-regulated. Comparing these results with Tet ChIP-seq data from our recent study shows that 448 Tet-bound genes are upregulated in the DNA binding domain mutant. The RNA of 444 out of 448 genes do not show 5hmC modification, indicating a distinct function of the Tet DNA binding and catalytic domains. The genes whose RNA level is increased in TetαXXC exhibit low levels or the absence of expression in wild-type brains demonstrating that Tet can suppress gene transcription via its DNA binding domain. Our study depicts a new function of Tet protein; it regulates axon outgrowth via an epigenetic mechanism in which Tet binds DNA and suppresses unwanted gene transcription in the brain, independent of cytosine methylation.

493C  (E)close but no cigar: Essential developmental programs transcriptionally regulated by the chromatin modifier KDM5 Michael Rogers1, Coralie Drelon1, Helen Belalcazar1, Owen Marshall1, Julie Secombe1 1) Albert Einstein College of Medicine, Bronx, NY; 2) Menzies Institute for Medical Research, University of Tasmania, Hobart, Australia

Chromatin-regulating proteins are powerful supervisors of the genome that are critical for the precise control of gene expression. Developmental processes depend on these proteins for the coordinated regulation of transcriptional programs that are essential for proper tissue formation and organismal maturation. However, there still exists significant gaps in knowledge about the mechanisms of this fine-tuning aspect of gene expression, and tissues like the prothoracic gland provide fantastic settings to dissect these dynamics during development. Here we show that lysine demethylase 5 (KDM5, formerly Lid), a histone modifier capable of dynamically regulating chromatin through multiple functions, plays critical roles in the prothoracic gland. Interestingly, animals lacking canonical KDM5 histone demethylase activity are viable, but dm5 null mutants exhibit delayed development and a failure to eclose after metamorphosis. Although KDM5 is ubiquitously expressed across the organism throughout development, we have demonstrated that restoring KDM5 expression specifically within the prothoracic gland of dm5 null mutant animals rescues both the larval developmental delay and pupal lethality. Our studies show that KDM5 regulates MAPK signaling within prothoracic gland cells that functions in promoting endoreplication and ecdyson hormone production. Leveraging Targeted DamID technology, we have generated the first genomic-binding data set (akin to ChIP-seq) for prothoracic gland cells and identified the genomic targets of KDM5 localization across the genome. We are currently integrating these data with transcriptome, interactome, and cellular analyses to characterize KDM5-mediated mechanisms of gene expression regulation in these cells. Taken together, our studies provide key insights into the chromatin regulation critical to prothoracic gland function as well as expand our understanding of the functions of the transcriptional regulators like KDM5 that coordinate development.

494A  Extracellular neuronal stimulation promotes Tip60 histone acetyltransferase mediated epigenetic neuroplasticity gene control in the Drosophila brain. Christina Thomas, Ellen Armour, Felice Elephant Drexel University

Experience-dependent (ED) plasticity is central for establishing memories and can influence the severity of AD progression. In this regard, we demonstrated that extracellular stimulation of rat hippocampal neurons in vitro triggers epigenetic gene regulator Tip60 nuclear import with concomitant genomic reorganization and synaptic gene induction. We further show that Tip60 nuclear import is recapitulated in the fly brain in vivo in response to stimulating neurons indubitably or by exposure to ED living conditions. Strikingly, Tip60 is found excluded from hippocampal nuclei in AD patients. Here we test the hypothesis that Tip60 extracellular mediated subcellular dynamics functionally impact synaptic gene control and cognition in vivo and that this process goes awry during early stages of AD. Using genetically modified Drosophila to inhibit the Shaker potassium pump within the neuronal membrane, we induced neuronal stimulation in the fly brain. We then used qPCR on RNA isolated from dissected brains in induced versus non-induced flies to assess expression levels of activity dependent plasticity genes we previously demonstrated to be bona-fide Tip60 targets. Strikingly, we found that as Tip60 increased within the nucleus, expression of specific Tip60 target neuroplasticity
genes including dlg, dsh, and futsch were induced. These results suggest that Tip60 shuttles into the nucleus in vivo to activate gene expression of synaptic plasticity genes. Future directions will focus on exploring the colocalization of Tip60 with transcription factors (TFs) to further elucidate the mechanism underlying Tip60’s role in activity dependent gene control and ED plasticity.

495B Extracellular stimulation triggers Tip60 HAT nucleocytoplasmic transport in the Drosophila brain with concomitant induction of Tip60 target neuroplasticity genes Ellen Armour, Gabrielle Greco, Felicity Khoa, Felice Elefant Drexel University

Experience-dependent (ED) plasticity is central for establishing memories and is driven by epigenetic mechanisms that regulate dynamic gene transcription in response to neuronal stimulation. Our lab has previously shown the histone acetyltransferase (HAT) Tip60 shuttles into the nucleus in vitro to regulate activity dependent gene control of synaptic plasticity genes. Furthermore, we also found that Tip60 is largely excluded from the nucleus in hippocampal neurons from Alzheimer’s disease (AD) patients. However, if and how Tip60 shuttles in the in vivo brain to regulate the expression of activity dependent synaptic genes is yet to be elucidated. Here, we used immunohistochemistry (IHC) to compare the subcellular localization of Tip60 HAT in the Drosophila brain under normal conditions and in response to stimulation of fly brain neurons in vivo either by inducibly activating potassium channels using genetic manipulation or by exposure to natural ED living conditions. Strikingly, we found that both inducible and ED living condition mediated neural induction triggered changes in Tip60 subcellular localization evidenced by a significant increase of Tip60 in the nucleus with concomitant induction of previously identified Tip60 target genes. Our results support a model by which neuronal stimulation promotes nuclear import of Tip60 to promote induction of activity-dependent neuroplasticity gene transcription. Future work will focus on labeling Tip60 with a GFP tag to track its in vivo nucleocytoplasmic transport in real-time and determine whether this processes is disrupted in brains from Drosophila that model AD. We will also mutate the nuclear localization sequences and nuclear export sequences to determine the functional impact of disrupting Tip60 nucleocytoplasmic transport on cognition and use chromatin assays to explore underlying mechanisms.

496C Regulation of Polycomb silencing initiation during nurse cell development Steven DeLuca Brandeis University

Polycomb group (PcG) proteins repress transcription and maintain a molecular memory of gene inactivity that is important for embryonic patterning and cell differentiation. Where and when Polycomb silencing initiates is therefore critical for producing correct gene expression patterns during later development. While biochemical and genetic approaches have identified many PcG proteins, how these proteins collaborate to correctly initiate silencing in genomic space and developmental time and poorly understood. We developed a new model system to study Polycomb silencing initiation. We recently discovered that nurse cells in the Drosophila ovary initiate Polycomb silencing on common target genes similarly to somatic cells in the early embryo. During silencing initiation, accessory proteins remodel PRC2 mobility across the transcriptionally inactive B-compartment. Faster migrating PRC2 complexes methylate and silence most inactive regions and slower migrating complexes focus PRC2 activity on traditional Polycomb target genes. By combining genetics and live imaging, we determined how individual PcG proteins contribute to this transition. A single protein, Scm, is critical for establishing high concentrations of PcG proteins on target genes during the initiation step. We will present our recent findings for how Scm is developmentally regulated and how it contributes to the site-specificity of PcG protein targeting. We will also present our ongoing efforts to use single molecule tracking to measure the kinetic properties of PcG protein complexes before and after silencing initiation.

497A Developmental ethanol exposure causes changes in the expression of histone modifying enzymes and results in long-term changes in gene expression Joshua Marsh, Madeleine Mok, Rachael French San Jose State University

Ethanol is a teratogen. Developmental alcohol exposure (DAE) in humans leads to a Fetal Alcohol Spectrum Disorder (FASD). Individuals with FASD can exhibit a variety of deleterious phenotypes, including slow growth, metabolic changes, behavioral difficulties, and intellectual disabilities. Recent findings in mammals indicate that metabolic changes associated with DAE involve long-term changes in gene expression mediated by epigenetic effects.

We have established Drosophila as a model for FASD. Using this model, we have found that flies exposed to ethanol during larval stages display phenotypes similar to mammals exposed during fetal development, including developmental delay, reduced adult size, smaller brains, CNS dysfunction, reduced sensitivity to ethanol sedation, impaired insulin signaling and lipid metabolism, and reduced survival. Some of these phenotypes, including impaired lipid metabolism and sedation resistance, persist into adulthood. Consistent with this observation, we have shown that DAE causes long term changes in gene expression.

Ethanol exposure in adult flies alters the expression of histone modifying enzymes. We hypothesize that some of the long-term changes in phenotype and gene expression caused by DAE are due to epigenetic alterations in gene expression due to similar effects on the expression of histone modifiers. To test this hypothesis, we tested the DAE sensitivity of flies mutant for a variety of histone modifiers. In addition, we used qPCR to measure the expression of genes encoding...
histone modifiers in ethanol-reared larvae.

We will present data showing that mutations disrupting the histone modifiers Sirt1, Lid, dG9a, JHDM2, and NO66 result in changes in sensitivity to DAE. Additionally, DAE causes reduced expression of G9a, Sirt1, and Lid in larvae. Thus, DAE alters epigenetic regulation of gene expression, and it is likely that some DAE-induced phenotypes are due to these effects.

We will additionally present the results of experiments to test the effects of DAE on additional histone modifying enzymes, as well as the role of those proteins in sensitivity to developmental ethanol. In addition, we are testing whether persistent DAE-induced phenotypes are due to changes in epigenetic regulation of gene expression.

498B Epigenetic regulation of energy homeostasis by the RNA adenosine methylation Daniel Wilinski1, Tahrim Choudhury1,2, Monica Dus1 1) Department of Molecular, Cellular and Developmental Biology, The University of Michigan, Ann Arbor; 2) PREP Program, The University of Michigan, Ann Arbor

Epigenetic mechanisms control the proper function of beta cells and alterations in these pathways have been linked to the development of type 2 diabetes (T2D), a debilitating disease that affects ~30% of Americans and that has no cure. Among these epigenetic mechanisms, the methylation of N6-adenosines in mRNAs (m6A) –which controls mRNA stability, turnover, and translation– has recently emerged as a novel regulator of ß-cell biology. In mammals, many of the mRNAs important for insulin secretion and signaling, including the insulin mRNA, are marked with this epigenetic mark and many are hypomethylated in the ß-cells of people with T2D. Consistent with this, mutations in the methyltransferase enzymes impair insulin secretion and glucose homeostasis in mammals, suggesting that m6A directs essential aspects of ß-cells physiology. However, the molecular underpinnings of these phenotypes are not known and the precise function of this epigenetic pathway in the biology of the ß-cells remains largely a mystery. This lack of knowledge has hindered progress in uncovering the underlying causes of T2D and in developing new therapeutic interventions based on this epigenetic pathway. Here we propose to exploit the unique advantages of the fly D. melanogaster model to identify the molecular mechanisms through which the m6A pathway regulates the physiology of ß-cells. We discovered that in flies mutations in the m6A pathway give rise to similar cellular and physiological phenotypes measured in mammals. Our hypothesis is that this epigenetic pathway regulates the responses of the insulin cells to glucose and amino acids and the release of insulin. We are currently taking a multidisciplinary approach that includes molecular biology, genomics, ex vivo imaging, and metabolic measurements to test these hypotheses and will present the result of these experiments at the conference. Discovering how specific epigenetic mechanisms direct distinct ß-cells processes is relevant to public health because it will advance our understanding of the biology of these cells and the etiology of T2D.

499C Pleiotropic fitness effects at the Uhg4-Boot locus in Drosophila melanogaster Rebecca A MacPherson1,2, Robert R. H. Anholt1,2, Trudy F. C. Mackay1,2 1) Center for Human Genetics, Clemson University, Greenwood, SC; 2) Department of Genetics and Biochemistry, Clemson University, Greenwood, SC

U snoRNA host gene 4 (Uhg4) encodes an antisense long noncoding RNA that is moderately expressed throughout the fly, with highest expression in embryos and adult ovaries. Uhg4 is host to seven small nucleolar RNAs (snoRNAs) and is coregulated with a subset of 37 snoRNAs that are differentially regulated in a background-dependent manner in females in response to developmental ethanol exposure. Bootlegger (Boot) is immediately upstream of and in opposite orientation to Uhg4 and encodes a protein critical for nuclear export of noncoding piwi-interacting RNAs. Expression of Boot is almost exclusively confined to ovaries. We have generated multiple deletions spanning the first exon of Uhg4 and the promotor regions of both Boot and Uhg4 using CRISPR-Cas9 and revealed a spectrum of pleiotropic effects on fitness traits, including increased development time, changes to sleep and stress response, background-dependent decreases in viability, as well as sterility in both sexes. RNA sequencing data suggest that the region between Uhg4 and Boot is required for normal expression of Uhg4, two intronic snoRNAs hosted by Uhg4, as well as Boot, leading to 668 differentially expressed genes. Gene ontology enrichment analysis showed enrichment of genes involved in immune response, ribosome formation, and DNA repair. We compared CRISPR-Cas9 deletion mutants of the Boot-Uhg4 regulatory region with RNAi-mediated knockdown of Boot and found that many of the effects on organismal phenotypes can be attributed both to Boot and the IncRNA Uhg4. However, we also observe differences in sexual dimorphism as well as antagonistic effects on fitness traits due to disruption of these genes. Thus, this regulatory region in the Drosophila genome highlights a long noncoding RNA and a protein coding gene that are both essential for reproduction and fitness.

500A Using Natural Variation and Machine Learning to map Gene Regulatory Networks Prasad Bandonkar1, Elle Rooney2, Samiul Haque2, Cranos Williams2, Gregory Reeves1 1) Texas A&M University; 2) North Carolina State University

In most well-characterized Gene Regulatory Networks (GRNs), maps of GRNs remain incomplete. The GRN for regulating anterior-posterior (AP) patterning in early Drosophila embryos, is a well-studied network and most of the major components have been identified. However, the minor components of the network, responsible for compensatory
regulation, remain elusive. Typical molecular techniques, such as a knockout, could result in several lost connections, swamping out compensatory regulation. Instead, we are using natural variation in the ~200 fly lines in the Drosophila Genetic Reference Panel (DGRP) to identify and characterize novel components of the network, using machine learning techniques.

The segmentation gene network that patterns the anterior-posterior axis in the early embryo is hierarchical and we are investigating the time period that is roughly mid-way between the start of nuclear cycle 14 and cellularization. During this time period, maternal coordinate genes such as Bicoid (Bcd), activate and regulate Gap genes such as Krüppel (Kr), and together they regulate pair-rule genes such as even-skipped (eve). The idea is to quantify the spatial expression patterns of the genes of interest to the segmentation GRN and use the subtle differences in spatial positions to map the cause back to specific regions in the genome and to the transcriptome. With 13 fly lines and imaging two genes- Kr and eve, searching within a 20 kb region of the genome, we were able to identify pangolin (pan) as a novel component of the AP patterning network. With data from ~200 fly-lines, we expect to have enough statistical power to identify several new components. We have built image analysis pipelines to reliably extract spatial gene expression data from a large number of stained embryos without manual supervision. Using unsupervised machine learning techniques over dataset from all ~200 fly lines and searching across the genome and the transcriptome, we expect to uncover novel components of the AP patterning network, or at the very least better characterize the existing network.

### 501B Sources of variation in gene expression Siddhant Kaira, Stephen Lanno, Lupita Sanchez, Joseph Coolon Wesleyan University, Middletown, CT

A century of genetic research has revealed that there are numerous causes of organismal trait variation. These include important contributions from an organism’s genotype, environment, age, and even the environment experienced by previous generations referred to as transgenerational effects. While a multitude of studies has demonstrated diverse and abundant effects of each source of variation in a wide variety of organisms, the relative contribution of the different sources of variation remains largely unknown. This has largely resulted from a lack of studies that simultaneously determine the contributions of each source of variation within a single experiment, and this is especially true for transgenerational effects that are in general poorly understood. Here we quantified genome-wide gene expression traits in Drosophila and compared the contribution of genotype (D. simulans vs. D. sechellia), environment (control food vs octanoic acid food), age/developmental stage (L3 larvae vs adult), and previous generation environment (control food vs octanoic acid food) to variation in gene expression levels. We followed this with analysis of genome-wide allele-specific expression in F1 hybrids between D. simulans and D. sechellia to disentangle the cis and trans-regulatory contributions to differences in variation for each source of variation. We found that genotype and developmental stage have much greater effects on gene expression than environmental differences and all were much more abundant than the variation due to trans-generational effects from previous generation exposure to different environments. By simultaneously analyzing major sources of variation in the same experiment, our results suggest a hierarchy of importance of the different sources of variation in traits that answers a long-standing question in genetics.

### 502C Using ISRES+, an evolutionary optimization algorithm to fit experimental data in systems biology models Razeen Shaikh, Prasad Bandodkar, Gregory Reeves Texas A&M University

Modern genetic research focuses on elucidating protein function and pathways. These involve interactions among various entities that influence gene regulation and expression. The field of Systems Biology seeks to devise experimental and computational approaches to understand how these interactions impact the behavior of the whole. However, systems biology models contain many unknown parameters and assumptions, and to improve their accuracy they must be fit to experimental data. One historically successful approach to fit data is to use evolutionary algorithms. These algorithms begin with randomly-selected parameter sets, which are improved in each generation by ranking and selecting the best parameter sets. These parameter sets are then adapted to experimental data using mathematical strategies like “recombination” and “mutation” to reach a global optimum.

Improved Stochastic Ranking Evolution Strategy (ISRES) is one such evolutionary optimization algorithm, developed by Runarsson and Yao, 2005. It uses stochastic ranking and island hopping to solve nonlinearly constrained optimization problems. It is an approach to fit experimental data to a rule-based model and obtain the best-fit parameter set. We modified this algorithm to make it faster and more accurate. We tested this modified algorithm, ISRES-plus, to verify the improved performance using two models from literature for this: Schmierer et. al, 2008 model for TGF- signaling in HaCat Cells and Manu, et. al, 2009 model for gap gene segmentation in Drosophila Melanogaster. We generated a best-fit parameter set with ISRES and ISRES-plus and verified that ISRES-plus is faster, robust, and more accurate than ISRES.

### 503A Genome-wide Effects of the GeneSwitch GAL4 System on Drosophila melanogaster Gene Expression Caroline Pitton, Sara Gregory, Zachary Drum, Joseph Coolon Wesleyan University, Middletown, CT

Our understanding of gene function in model organisms has increased dramatically over the last 30 years utilizing new reverse genetic techniques to manipulate gene expression. Selectively changing the expression of a gene in an
otherwise wild-type organism allows for much greater understanding of that gene’s function and downstream effects on phenotypes than mutant analysis alone. This is especially important for studying essential genes, pleiotropic genes and genes with tissue, stage, sex or environment specific effects. In the Drosophila melanogaster model system, use of the GAL4-UAS system is one such genetic tool for experimental manipulation of gene expression that is used in a large number of studies to date. A recent improvement added a hormone-induced (mifepristone) GAL4 expression system called GeneSwitch making it even more tractable and applicable to many experimental questions. Because the GAL4-UAS system was taken from yeast, mifepristone from mammals and both used in D. melanogaster, it is largely considered to act specifically with few if any off-target effects. To test the effects of GAL4 protein expression and mifepristone exposure on genome-wide gene expression, we performed RNA sequencing (RNA-seq) and identified numerous D. melanogaster genes that are significantly differentially expressed in response to GAL4 and RU486. Our study provides evidence that use of the GeneSwitch GAL4-UAS system requires effective control experiments and provides suggestions for future studies to maximize the utility of this genetic toolkit.

504B  A homeostatic transcriptional response counteracts I-SMAD activity in Drosophila motor neurons Jacqueline Kanzer, Zainab Seyal, Mikolaj Sulkowski Southern Connecticut State University

Given its pleiotropic nature, TGF-β signaling must be precisely regulated both intra- and extracellularly to have the proper effect on target cells. Inhibitory SMADs (I-SMADs) are a potent intracellular regulator of TGF-β signaling. However, there are numerous hypotheses regarding the mechanism by which I-SMADs function. Although biochemical studies show that Dad, the sole Drosophila I-SMAD, inhibits the receptors that phosphorylate the effector molecule Mad, our in vivo results indicate that Dad functions downstream of the receptors. We have found a paradoxical increase in phosphorylated Mad (pMad) in motor nuclei following Dad overexpression. qPCR analysis supports the hypothesis that a homeostatic transcriptional response counteracts I-SMAD activity in motor neurons.

505C  Characterization of a Drosophila Activin signaling network Yisi Louise Lu, Hiroshi Nakato, Michael O’Connor University of Minnesota, Minneapolis, MN

Drosophila is an ideal model system for the study of TGF-β signaling networks and their roles in organism development and homeostasis. In Drosophila, there are three TGF-β/Activin-like ligands, Myogetin (Myo), activin-β (Actβ), and Dawdle (Daw). These factors have been implicated in regulation of different physiological activities in a tissue specific manner. For example, loss of actβ results in small muscles with altered carbohydrate metabolism and changes in NMJ electrophysiology. Loss of daw results in major metabolic dysfunction including altered carbohydrate metabolism and a disruption in the TCA cycle, while loss of myo leads to smaller brain and imaginal disc sizes. Although loss of each Activin ligand gives signal through dSmad2, a common intracellular transcriptional transducer. Our goal is to elucidate the mechanisms by which the three Activin-like ligands produce specific responses and phenotypic outcomes in different tissues. We hypothesize that tissue-specific responses result from several attributes of the signaling network. First, there are cross regulatory interactions between the different ligands. Second, each ligand preferentially signals through one of three isoforms of the Type I receptor encoded by the Babo locus. Third, each tissue expresses its own complement of Babo isoforms. Finally, the transcriptional transducer dSmad2 associates with different tissue-specific cofactors to stimulate an appropriate downstream response for that tissue. The lab’s long-term goal is to elucidate how these three ligands coordinate to produce the appropriate balance of signals to maintain viability and physiological homeostasis.

We have begun to investigate the cross-regulation relationships among the three Activin ligands by qPCR studies as well as by employing transgenic GFP reporter fly lines. Our preliminary data suggested Daw is negatively auto-regulated by dSmad2 while Myo may be positively autoregulated and that these ligands likely cross-regulate each other’s expression in a tissue-specific manner. We have also begun efforts to identify direct downstream targets of Activin signaling through transcriptomic and ChIP-seq analyses. Preliminary analysis of this data will be presented. These studies will provide a deeper understanding of the Activin signaling network and the regulatory interactions that it provides in controlling important physiological activities such as proliferation and energy utilization in several key tissues including muscle, fat and brain.

506A  Determining how antagonistic transcription factors control transcription dynamics for robust cell fate specification by single nuclei imaging of transcription factor and target mRNA dynamics Suzy SJ Hur1, Sebastian Bernasek 2, Nicolás Peláez2, Richard W. Carthew3, Ilaria Rebay1 1) Ben May Department for Cancer Research, University of Chicago, Chicago, IL; 2) Department of Chemical and Biological Engineering, Northwestern University, Evanston, IL; 3) California Institute of Technology, Pasadena, CA 91125, USA. Hanna H. Gray Fellow of the Howard Hughes Medical Institute (HHMI); 4) Department of Molecular Biosciences, Northwestern University, Evanston, IL

During animal development, a highly organized pattern of tissue-specialized cell fates emerges from a population of initially equivalent progenitor cells. This is mediated by transcription factors (TFs) acting downstream of spatially localized extracellular signals to produce spatially refined expression of fate determining genes. Each gene is transcribed in discontinuous bursts with gene-specific temporal dynamics. How TFs control the mRNA expression dynamics of their
targets, and how these dynamics in turn contribute to the specificity and robustness of cell fate transitions remains an important question in developmental biology.

The Drosophila eye development is an ideal system to address this question. In the developing eye disc, a precise complement of eight photoreceptor neuron fates (R1-R8) are specified in each ommatidium from a population of multipotent progenitor cells through the reiterative use of RTK/MAPK signaling. Transition from a progenitor to a specified state is mediated by an antagonistic pair of TFs, Pointed and Yan, which compete for the same binding site and exert opposing functions downstream of RTK/MAPK signaling. Yan represses gene expression to maintain cells in a progenitor state while Pointed activates the expression of distinct photoreceptor fate genes to trigger the transition to a specified state.

We found that increased Pointed-to-Yan ratio in the nucleus, rather than the absolute concentration of either TF, promotes the transition to a specified state. Focusing on the R3/R4 outer and on R7 inner photoreceptor fates, we show that genetic perturbations that elevate the ratio prematurely in the progenitors induce ectopic fate transitions while perturbations that prevent the ratio increase bias cells to remain in the progenitor state. By combining immunostaining, single-molecule RNA fluorescent in-situ hybridization (smFISH), and quantitative imaging, we will simultaneously measure the protein levels of Pnt and Yan and nascent mRNA levels and bursting frequency of the R3/R4 gene salm and the R7 gene pros in individual cells that remain in the progenitor state vs. those entering the specified R3/R4 and R7 states. By comparing tissues with wild type and altered Pnt and Yan levels, we will present our latest mechanistic models for how distinct mRNA transcription dynamics downstream of Pnt and Yan direct precise and reliable tissue patterning.

507B  In vitro identification of critical cis elements in the embryonic Drosophila histone locus Pamela Díaz-Saldana¹, Tsutomo Aoki², Paul Schedl², Leila E. Rieder¹ 1) Dept. of Biology, Emory University, Atlanta GA; 2) Dept. of Molecular Biology, Princeton University, Princeton, NJ

The timing of the nuclear cycles during early Drosophila embryonic development relies heavily upon the availability of histones. The five canonical histone genes are clustered into an array, and ~100 arrays are arranged in tandem at the single histone locus on chromosome 2L. Prior to zygotic genome activation, the histone locus is activated, which includes recruiting a suite of transcription and processing factors collectively known as the histone locus body (HLB). Even though embryonic HLB formation is an evolutionarily conserved event, we do not yet understand the molecular interactions that mark the histone locus for early and unique regulation. Previous in vivo work using transgenes pinpointed specific cis elements (GA-repeats) in the histone3/histone4 promoter that identify the locus for HLB formation by recruiting the transcription factor CLAMP. CLAMP also participates in male dosage compensation. In order to probe the interaction between histone locus and transcription factors more precisely, we tested histone array sequences in vitro for their ability to bind proteins in embryo extract. We performed gel shift assays using histone array probes, confirming the requirement for the GA-repeat cis elements in the histone3/histone4 promoter. In addition, we performed supershifts and observed that both CLAMP and the GA-repeat binding protein GAGA factor bind to this region in vitro. Interestingly, both of these proteins are members of the late boundary complex (LBC), a giant insulator complex that forms in late embryos and participates in both fab7 boundary function and dosage compensation. In the future, we will test the ability of recombinant CLAMP alone to target cis elements in the histone3/histone4 promoter, which it may do prior to LBC formation.

508C  Investigating the Effects of Genetic Distance and Regulatory Elements on Tandem Gene Duplicate Expression in Drosophila Melanogaster Georgia McClain, David Loehlin Williams College

The ways in which a gene’s position in the genome and the surrounding genomic contents can affect its expression is not well understood. Tandem duplicate genes are a good model for studying these phenomena, as previous studies have demonstrated evidence of a nonlinear relationship between gene copy number and protein expression. Furthermore, there are several known examples of human diseases associated with copy number variation and relative gene overexpression, so a better understanding of gene duplication and its effects may provide valuable information in addressing the specific causes of these disorders. The Loehlin lab is attempting to investigate the characteristics of tandem Adh gene duplicates in Drosophila melanogaster that contribute to deviation from two-fold expression. We are investigating two qualities of synthetic tandem Adh duplicates: distance between duplicates, and regulatory element duplication. To do this, we inserted FRT constructs on either side of the Adh gene of Drosophila melanogaster and used them to create recombinant flies. To test whether the distance between tandem Adh copies affects relative protein expression level, we designed recombinants with Adh genes at different distances from one another. To test whether the duplication of Adh regulatory elements affects expression, we keep the distance between the duplicates the same, but vary whether or not the regulatory region was duplicated. We can then measure the ADH enzyme levels in the flies in order to determine whether ADH expression is influenced by distance between the gene duplicates and/or duplication of regulatory elements. Preliminary results show that our method successfully creates recombinant Drosophila melanogaster individuals that have tandem duplicate Adh genes, and that these duplicates have higher expression levels relative to individuals with a single copy.
In the last two decades many studies have utilized the powerful method of knocking down gene expression via RNAi. However, the level of knockdown varied between genes leading to a range of mRNA levels from full knockdown to very minimal decrease in the target mRNA. We found that two post-transcriptionally regulated mRNAs are protected in P-bodies, giving us a possible hint as to why certain mRNAs are not susceptible to RNAi. This hypothesis stemmed from our initial finding that oskar mRNA was only degraded after it localized to the posterior pole. oskar mRNA is transcribed and transported to the oocyte throughout oogenesis and it is localized to P-bodies. Since ribosome entry into P-bodies is inhibited, it is reasonable for us to hypothesize that at the posterior pole, where oskar mRNA is translated, the transcript has to be released from P-bodies, thus allowing access to the transcript for the RNAi machinery. Our finding that mRNAs localized in P-bodies are not degraded by RNAi came as a surprise to us. Numerous proteins that are components of the RNAi machinery localize into P-bodies, and we expected a strong knockdown when we initiated oskar RNAi. During early investigations into P-bodies, it was hypothesized that P-bodies are sites of mRNA decay due to the presence of numerous decay enzymes such as Dcp 1, Dcp 2 and Xrn 1. Recently, experiments carried out in live cells did not find mRNA degradation in P-bodies. Furthermore, it would be also of interest to assess if other translationally repressed mRNAs (i.e. nanos) are also protected from RNAi in P-bodies. Our results indicate that P-body protection is maintained in the nurse cells, but investigation of mRNA levels in the nurse cells could reveal further insights into this intriguing mechanism.

**The mRNA regulatory function of Brat is essential for development and neurogenesis**

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TRIM-NHL proteins share a conserved domain architecture and play crucial roles in stem cell biology, fertility, and development. Recently, multiple TRIM-NHL proteins were shown to recognize specific RNA motifs and structures via their NHL domain. TRIM-NHLs negatively regulate their bound mRNAs. In light of this observation, we sought to determine the biological role of their RNA-binding activity. Additionally, we investigated the mechanism by which TRIM-NHLs repress mRNAs.

We focused on the *Drosophila* TRIM-NHL protein Brain Tumor (BRAT), which controls development and stem cell fate. First, RNA-binding defective mutations were introduced into the endogenous brat locus via CRISPR/Cas9 genome engineering. Our phenotypic analysis demonstrated that the key residues necessary for RNA-binding *in vitro* are essential for survival and larval neurogenesis. RNA-binding defective mutations phenocopy the lethality observed in loss-of-function brat mutations. Additionally, these mutations produce ectopic neuroblasts and brain tumors similar to classical brat mutants. These results demonstrate the essential function of BRAT derives from its ability to bind and regulate mRNAs.

We further used cell-based reporter assays to elucidate the molecular mechanism by which BRAT controls protein expression. We find that BRAT-mediated repression depends on the 3’ poly-adenosine tail of target mRNAs and components of the CCR4-NOT deadenylase complex. We also identified the domains of BRAT that are necessary to repress target mRNAs. Together, our data support a mechanism wherein BRAT recruits deadenylases to reduce translation and promote decay of target mRNAs.

Collectively, our findings provide crucial insights into the molecular mechanism and function of BRAT *in vivo*.

**Protein-RNA interaction drives co-transcriptional regulation and RNA processing**

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Biomolecular condensates are membrane-less compartments formed by the biophysical phenomena of phase transition mainly through Protein-RNA interactions. Both transcriptional complexes and RNA processing units are reported to form such condensates. Several transcription factors and RNA binding proteins that contain IDR (Intrinsically Disordered Regions) have been reported to have a role in phase transition, and thus it is interesting to study how RNA-protein complexes can potentially form and dissolve such condensates. In the present study, we have explored the potential of DNA binding transcription factor protein CLAMP (Chromatin Linked Adaptor for MSL Proteins), which contains IDR, in *Drosophila Melanogaster* as a protein that can form bio-condensates. CLAMP recruits the MSL (Male Specific Lethal) Complex, an RNA-protein complex consisting of long non-coding RNAs roX1 and roX2 along with five other proteins, to the male X-chromosome resulting in the transcriptional upregulation of the male X-chromosome in a process known as dosage compensation. Therefore in this project, we studied the interaction between CLAMP and roX2 RNA to test the hypothesis that these two components might drive the process of dosage compensation via phase separation. Also, CLAMP regulates sex-specific splicing, a process regulated by ribonuceloprotein complexes called
splicing components that coordinate co-transcriptional RNA processing via the mechanism of alternative splicing. In flies and mammals many RNA-binding proteins are part of the alternative splicing spliceosome complex along with non-coding RNA components. In Drosophila, CLAMP binds to several of such RNA binding proteins that are reported to interact with a long non-coding RNA hsrω-n. Thus, we also explored CLAMP’s binding with hsrω-n to test CLAMP’s potential as a legitimate player in phase transition and formation of biomolecular splicing condensates. Using electron mobility shift assays (EMSA) we found that CLAMP and the CLAMP domain that contains the IDR physically bind to both roX2 and hsrω-n non-coding RNAs, reinforcing our hypothesis that CLAMP is a bifunctional protein with DNA/RNA binding capacity and has the potential of regulating transcription and co-transcriptional RNA processing by manipulating bio-physical properties of RNA-protein complexes.

512A  Nonsense-mediated mRNA decay plays an essential role during female germline development in Drosophila melanogaster Omar Omar1,2, Arwa Abdelhamid2, Emily Makowicz2, Diana Bratu1,2 1) The Graduate Center, CUNY; 2) Hunter College, CUNY

Across eukaryotes, nonsense-mediated mRNA decay (NMD) is a vital cellular surveillance process. Its initial role was presumed to be mitigation of truncated proteins by degrading aberrant transcripts encoding for them. On an organismal level, NMD is necessary for proper growth and development but how this is accomplished remains poorly understood. Our studies in flies have found that efficient knockdown of various NMD factors results in embryonic lethality as well as perturbations of essential maternally-derived factors, including localized mRNAs necessary for body axis patterning. Moreover, these knockdowns result in improper germline cyst formation due to problems during early mitotic events initiated in the female germline stem cell niche. Taken together, our data suggest that Upf1 protein functions in regulating mRNA stability of transcripts necessary for both proper maternal embryonic axis patterning and germline development.

513B  Bruno 1 and Cup interdependent regulation of oskar mRNA life cycle Livia Bayer1, Irina Catrina2, Stephen Formel3, Rishi Ravichandran4, Juan Cambeiro1, Lizaveta Slinko1, Diana Bratu1 1) Hunter College; 2) Yeshiva University; 3) Tulane University; 4) Sloan Kettering Institute

In D. melanogaster, a subset of mRNAs essential for embryo development are spatio-temporally controlled during oogenesis. oskar mRNA is a critical transcript for germplasm assembly, which is translationally regulated by Bruno 1 and Cup. The current model suggests that Bruno 1 is the key factor for recruiting Cup to oskar mRNA that enables the formation of the translational silencing complex. Here, we address the spatio-temporal requirements for the formation of the Bruno 1-Cup complex contributing to a more universal understanding of mRNA regulation. Our work reveals that the Bruno 1-Cup interaction, as well as their interdependent influence on each other’s protein expression, lead to a precise oskar mRNA regulation. Cup mediates recruitment of oskar mRNA to P-bodies driving the translational repression and stability of oskar mRNA and inhibits Bruno 1’s role in the activation of translation. When Cup is removed from the complex, Bruno 1 is free to interact with Orb, thus facilitating translational activation.

515A  Exploring the novel role of a putative tRNA methyltransferase in synaptic growth and neuronal development Jennifer Dumouchel2, Kimberly Rose Madhwani2, Caley Hogan3, Jenna Lentini4, Kevin Welle5, Dragony Fu4, Kate O’Connor-Giles6,7 1) Therapeutic Sciences Graduate Program, Brown University, Providence, RI; 2) Neuroscience Graduate Program, Brown University, Providence, RI; 3) Laboratory of Genetics, University of Wisconsin-Madison, Madison, WI; 4) Department of Biology, Center for RNA Biology, University of Rochester, Rochester, NY; 5) Mass Spectrometry Resource Laboratory, University of Rochester Medical Center, Rochester, NY; 6) Department of Neuroscience, Brown University, Providence, RI; 7) Carney Institute for Brain Science, Providence, RI

Dynamic regulation of neuronal genes during transcription and translation is required for proper nervous system development and function. Investigation of transfer RNA (tRNA) modifying enzymes has highlighted the importance of chemical modifications for tRNA stability and protein synthesis. Yeast tRNA methyltransferase 9 (Trm9) methylates wobble uridines in anticodon loops to reinforce cognate codon-anticodon pairings. In metazoa, two Trm9 paralogs, ALKBH8 and TRMT9B have been identified. We identified TRMT9B in an RNAi screen as a negative regulator of synaptic growth, and find that it also plays a key role in regulating neurotransmission. While TRMT9B has been studied for its role as a tumor suppressor, its biochemical role has remained unknown. To study the role of the expanded family, we have generated mutants in both paralogs and assessed changes in bulk tRNA chemical modifications by mass spectrometry. Wobble uridine methylation is absent in ALKBH8 mutants, consistent with studies in mammals. In contrast, we do not observe deficits in wobble uridine methylation in TRMT9B mutants under basal conditions. Thus, TRMT9B may catalyze wobble uridine modification of a small subset of tRNA isodecoders not detectable by bulk nucleotide analysis or evolved a new function. Wild-type, but not methyltransferase-dead, transgenes rescue synaptic overgrowth in TRMT9B mutants, indicating that TRMT9B functions through a methyltransferase-dependent mechanism. Unbiased homology modeling of TRMT9B’s methyltransferase domain revealed a strong conservation of secondary structure with the reported yeast Trm9 X-ray crystallography structure; consistent with a role as a methyltransferase. To assess TRMT9B function, we turned to transcriptomic and proteomic studies, and will present our latest findings. We investigated transcriptome and proteome...
changes in TRMT9B null adult fly heads by RNA sequencing and high resolution mass spectrometry, respectively. Differentially expressed genes and proteins are consistent with TRMT9B’s role at synapses and prior studies of yeast Trm9, which modulates translation of specific mRNAs in response to DNA damage and oxidative stress. Together, our studies reveal a novel role for tRNA methyltransferase family member TRMT9B in the regulation of synaptic growth and neuronal function.


Development is an extraordinarily complex and tightly regulated process. A significant part of developmental regulation depends on gene regulatory networks (GRNs), consisting of interacting transcription factors governing spatial and temporal gene expression. Synthetic biology opens a new way to study GRNs: on top of deciphering natural GRNs, scientists reconstruct synthetic ones using engineered transcription factors and target genes. Existing synthetic GRNs have primarily been established in prokaryotes that benefit from toolkits consisting of ‘easy to assemble’ well-studied genetic elements, such as promoters and transcription factors.

In our project, we are developing a toolkit to reconstruct synthetic GRNs in vivo, in Drosophila melanogaster (D.mel). Our toolkit enables massive combinatorial assembly of synthetic transcription factors (sTFs) using a library of DNA binding domains, activating or repressing domains and a collection of engineered cis-regulatory binding sites that are absent from the (D.mel) genome. We are establishing the gene expression profiles derived from these sTFs individually or in a variety of combinations. Ultimately, sTFs will be used to recreate motifs that occur naturally in GRNs and logic gates. By providing this sTFs library we hope to facilitate the implementation of synthetic GRNs in D.mel and other model organisms, thus enabling in vivo modeling of the native developmental networks.

517V  Determinants of transcription factor function Lauren Hodkinson, Leila Rieder Emory University

Despite the crowded nuclear environment, transcription factors locate and bind cis elements across the genome while performing context-specific functions. To explore this phenomenon, we broadly aim to understand how transcription factors integrate cis sequence and genomic context to function uniquely at different loci, which is critical for development and disease. One example of a context-specific transcription factor is Chromatin-Linked Adapter for MSL Proteins (CLAMP), which targets GA-rich cis elements while performing several distinct functions throughout the genome. CLAMP primes the male X-chromosome for dosage compensation, regulates the accessibility of promoters genome-wide, and promotes formation of the conserved histone locus body (HLB), which regulates expression of the replication-dependent histone genes. Although CLAMP targets similar cis elements in all three contexts, it recruits very different locus-specific transcription factors. Here we investigate how the function of CLAMP at the histone locus is impacted by the origin of its cis binding elements. CLAMP binds a long GA-repeat element in the bidirectional promoter of histone genes 3 and 4 (H3H4p). We engineered flies to carry a transgenic histone locus in which we replaced the natural GA-repeating cis element in the H3H4p with CLAMP-recruiting GA-rich elements from the X chromosome. We assessed how X-linked cis elements impact HLB formation by staining third instar larval polytene chromosomes with antibodies specific to a core HLB protein as well as proteins found in the other CLAMP binding contexts on the X chromosome.

518V  De novo discovery of motifs enriched in promoters of D. ananassae F Element genes Annabelle Laughlin1, Wilson Leung2, Chris Shaffer3, Cindy Arrigo2, Genomics Education Partnership 1) Washington University in St. Louis, St. Louis, MO; 2) New Jersey City University, Jersey City, NJ

The Drosophila melanogaster Muller F Element exhibits mostly heterochromatic characteristics (e.g., high repeat density, low recombination rates), but the distal ~1.3 Mb region contains ~80 protein-coding genes that show expression levels similar to that of euchromatic genes. Interestingly, this region has expanded to ~19.1 Mb in Drosophila ananassae due to an increase in repeat density (particularly retrotransposons). This project seeks to understand the regulatory mechanisms that allow the successful transcription of F Element genes in such repeat-rich domains by conducting comparative analysis of D. ananassae, D. bipectinata, D. takahashii, and D. kikkawai, (where the F Element has expanded to different degrees), focusing on transcription start sites (TSS). The GEP annotation protocol uses experimental data (RAMPAGE, ATAC-Seq, RNA-Seq) and sequence similarity to other Drosophila species to define TSS positions and promoter-flanking regions. Over the past two summers, 97 unique TSSs in D. ananassae have been annotated. The TSS data were used in coding region annotations. For example, the locations of the putative TSSs relative to the available start codons was used as evidence to support the hypothesis that the G isoform of the D. melanogaster Zyx gene...
does not exist in *D. ananassae*. In a preliminary analysis using 26 unique TSSs in the *D. ananassae* F Element scaffold QMES02000012, ~77% of the TSS positions and ~86% of the narrow promoter-flanking regions were defined based on RAMPAGE or ATAC-Seq data. In contrast, ~54% of the wide promoter-flanking regions were defined by manual analyses (e.g., BLAST searches, multiple sequence alignments). A *de novo* motif discovery analysis of the 50 RAMPAGE peaks in QMES02000012 identified three significant motifs. Analysis of these motifs using TomTom showed that the two most significant motifs have no similarity to known *D. melanogaster* motifs, while the third showed similarity to a putative Top2 motif abundant at heterochromatic TSSs in *D. melanogaster*. These motifs are being used in searches of the genomes of the other three species to see if they are enriched near the promoters of F Element genes. Comparative analysis of *Drosophila* F Elements could identify regulatory motifs and promoter architecture that are unique to the F Element and that facilitate expression in heterochromatic domains.

**519V Regulation of gene expression by the HP1 variants**

John Schoelz1,2, Annesha King1,2, Nicole Riddle1,2 1) University of Alabama-Birmingham; 2) Department of Biology

Heterochromatin Protein 1 (HP1) was discovered in Drosophila and is a major component of heterochromatin. In *Drosophila melanogaster*, there are three somatically expressed HP1 proteins: HP1a, HP1b, and HP1c. HP1a is essential in flies and is necessary for genomic integrity. While enriched in heterochromatin, it has binding targets in heterochromatin and euchromatin and it acts as both a repressor and activator of gene expression. HP1b is found in heterochromatin and euchromatin as well and acts as repressor and activator of gene expression depending on the target. HP1c is found in both chromatin compartments as well, but its function seems to be mostly in gene activation. Thus, all three proteins are involved in the regulation of gene expression, and analysis of ChIP-seq data suggests that they share many binding sites. To understand the contribution of the HP1 proteins to gene regulation, elastic net regression models were utilized. In this study, we assessed gene expression changes by recruiting HP1 proteins to endogenous genes. These models suggest that HP1b is of particular importance, but that genomic features including promoter motifs, accessibility, and sequence composition also contribute. To address the model predictions, we used dCas9 to recruit HP1 proteins to specific targets in the genome. We found that dCas9 tethered HP1 proteins have different transcriptional outcomes depending on gene target.

**520V Regulation of PDF neuropeptide production in the central nervous system**

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Pigment-dispersing hormone was first characterized in Crustaceans for its role in the daily rhythms of the retinal pigment dispersion. Insect gene encoding PDH-homolog, PDF, was first characterized in *Drosophila*. PDF is an important output factor for the maintenance of the circadian rhythms via synchronizing various clock-gene expressing neurons. Expression of PDF was found in the three distinct groups of neurons in the central nervous system; s-LNvs and l-LNvs in the brain lobe, and abdominal ganglionic neuron (PDFab). However, regulation of PDF production is not well understood. We analyzed the role of *scro* in PDF expression. We found that overexpression of *scro* in PDF-negative neurons. While loss-of-function mutations caused abnormal development of the optic lobes, ectopic PDF expression in PDF-negative neurons. We also observed that knockdown of *disco-related* (*disco-r*) gene suppressed PDF expression, raising the possibility of *disco-r* as a putative PDF transcription activator. To understand the mechanisms of PDF precursor, we generated an antibody that is intended to detect only PDF-associated peptide. The antibody detected all endogenous cell bodies and axonal projections, suggesting that PDF maturation takes place in the vesicles while transporting to the axon terminals. This reagent will be used to study the genetic and molecular mechanisms of the maturational processes of PDF precursor in the clock neurons.

**521V It's about time: an investigation into the role of abnormal oocyte (abo) in embryonic histone gene regulation**

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*Abnormal oocyte* (*abo*) is a maternal effect gene first characterized by Larry Sandler in 1970. Loss-of-function mutations in *abo* cause defects in embryonic development. Later characterization of *abo* revealed that it negatively regulates histone gene expression—likely through targeting histone promoters. Interestingly, the viability defects prompted by *abo* mutation can be rescued by an overexpression of heterochromatic regions, suggesting that heterochromatin can act as a “sponge” to soak up the excess histones. Despite these important experiments, *abo* has received little attention since the early 2000s, and its precise role in histone repression is unknown. Thus, our goal is to define the molecular role of *abo* in histone gene expression. To that end, we generated V5-tagged *abo* lines via CRISPR. Using these lines, we probe *abo* localization through staining polytene chromosomes and determine abo expression levels by western blot. We also utilize available *abo* mutants and RNAi lines to determine the molecular mechanism of histone gene repression. Overall, our work will contribute to a growing body of knowledge regarding histone gene regulation in early embryo.
Many eukaryotic genes contain alternative promoters with distinct expression patterns. How these promoters are differentially regulated remains elusive. Here, we apply single-molecule imaging to quantify the transcriptional regulation of two alternative promoters (P1 and P2) of the Bicoid (Bcd) target gene *hunchback* in syncytial blastoderm *Drosophila* embryos. Contrary to the previous notion that Bcd only activates P2, we find that Bcd activates both promoters via the same two enhancers. P1 activation is less frequent and requires binding of more Bcd molecules than P2 activation. Using a theoretical model to relate promoter activity to enhancer states, we show that the two promoters follow common transcription kinetics driven by sequential Bcd binding at the two enhancers. Bcd binding at either enhancer primarily activates P2, while P1 activation relies more on Bcd binding at both enhancers. These results provide a quantitative framework for understanding the dynamics of complex eukaryotic gene regulation.

Enhancers are critical for eukaryotic transcriptional regulation. However, just how enhancers should be defined remains unclear. While reporter gene assays, which are function based, have been the traditional benchmark for enhancer identification, next-gen sequencing-based techniques that scan for open chromatin, histone modifications, or specific transcription factors (e.g. ATAC-Seq, ChIP-Seq) have become a new source of powerful, high-throughput methods for defining enhancers. Whether these various enhancer definitions consistently lead to identification of the same sequences is unknown. To compare the functional and the chromatin-level enhancer definitions, we analyzed the overlap between enhancers defined in two enhancer databases, REDfly (Rivera et al. 2019, *NAR* 47:D828) and EnhancerAtlas2.0 (Gao and Quan 2019, *NAR* 48:D58). REDfly uses primarily a functional definition based on reporter gene analysis, while EnhancerAtlas integrates the results of chromatin-level assays using a supervised learning model. We used REDfly’s search capabilities to build tissue-specific enhancer datasets and compared these with tissue-specific EnhancerAtlas datasets. Surprisingly, we found that only 4 of 11 sets (36%) showed statistically significant overlap. From this, we hypothesized that the observed discrepancies could be caused by the way data from multiple techniques/assays are integrated by the EnhancerAtlas method. We took the underlying EnhancerAtlas data subsets and compared them individually with their matched REDfly sets. 66% of the EnhancerAtlas subsets had significant overlap, a substantial increase from the previous, full-set comparisons, although still limited. However, these EnhancerAtlas subsets only had a median intersection with REDfly enhancers of 39%. Thus, even the sets with significant overlap include fewer than half of the expected reporter-gene defined enhancers from the corresponding REDfly set. We derive two conclusions from our findings. First, during the integration of the EnhancerAtlas data sets, enhancers present in the underlying EnhancerAtlas data are being lost, suggesting that a more sensitive learning model may be required. More importantly, the poor overlap between the reporter-gene defined enhancers and the chromatin-assay defined enhancers suggests that one or both of these approaches carries a high error rate. Further investigation will be required to determine which approaches lead to the most accurate enhancer definition.

Animals have different types of muscle fibers with distinct morphologies and contractile properties. These characteristics are established during development through regulation of alternative splicing and gene expression. Patterns of mRNA isoform expression differ between myofiber types, change as muscles differentiate, mature and age, and are often disrupted in muscle disease, suggesting that RNA regulation and in particular alternative splicing helps define muscle functional properties. It is therefore important to examine how RNA processing functions in healthy muscle, to better understand how misregulation contributes to muscle disease. Misregulation of the RNA-binding protein CELF1 is thought to underly myotonic dystrophy (DM1), but its function in muscle development is incompletely understood. Using *Drosophila melanogaster* as a tractable genetic model, we have previously shown using RNAi knockdown that the CELF1 homologue Bruno-1 (Bruno1) controls flight muscle specific alternative splicing, regulating both sarcomere growth and myosin contractility. We generated new CRISPR-mediated alleles in Bru1 that affect all Bru1 isoforms, resulting in stronger phenotypes in mutant flies that revealed structural defects at the earliest stages of myofibril formation, notably disorganization of the actin cytoskeleton that adversely affects myofibrillogenesis in differentiating IMF. Using temporally-restricted RNAi knockdown and rescue experiments, we demonstrate that there are distinct requirements...
for Bru1 mediated splicing during early and late stages of myofibril formation. After sarcomere formation, aberrant actin incorporation arrests growth in thin-filament length, but promotes lateral growth leading to formation of hollow myofibrils. We performed mRNA-Seq and mass spectrometry and identified misregulation of both sarcomeric proteins and other RNA-processing factors. Moreover, our temporal transcriptomics data reveal a progressive misregulation of gene expression and splicing as IFM development proceeds. Taken together, our data indicate that during muscle differentiation, Bru1 regulates cytoskeletal rearrangement necessary for myofibril formation as well as the balance in length versus lateral growth of the thin-filament. Defective RNA processing thus causes sarcomeric structural defects and progressive malfunction of dystrophic muscle.


Drosophila nochte encodes a 250 kDa protein with an N-terminal BAT2 domain, RGG motifs in the middle, and large low-complexity regions. Studies in Drosophila showed that a partial mutation of nochte leads to defects in temperature compensation of the circadian clock. A recent study shows that mouse Nocte orthologs PRRC2A is an m6A reader and its binding can stabilize specific mRNAs; and its inactivation disrupts neural development. However, the molecular functions of Nocte and its orthologs are still largely unknown. Our results from IP-Mass Spec experiments of RNA-binding proteins Top3b and TDRD3 identify Nocte as their strong binding partner. IP-Mass Spec experiment of Nocte shows that it interacts with other RNA-binding proteins and ribosomal proteins, supporting that Nocte is an RNA binding protein. We generated nochte knockouts by CRISPR and found that nochte-KO flies die at larval-pupal stages. Specific knockdown of nochte by RNAi in fly eyes induces rough and small eye phenotypes. The RNA-seq results from WT and nochte-RNAI eye discs indicate that Nocte depletion disrupts chaoptin transcription. Immunostaining, qRT-PCR and RIP-qPCR results show that Nocte is necessary and sufficient to promote the translation of glass mRNA, which encodes a transcription factor regulating chaoptin directly. With reporter assays and CRISPR edited alleles, we demonstrated that Nocte counteracts the suppression effects of the upstream ORF (uORF) at glass 5’UTR to maintain glass translation. We are investigating the detailed mechanisms of how Nocte regulates glass translation and its global effects on mRNA translation by Ribo-seq.

526V Dynamic time warping on sn- and sc-RNA-seq trajectories of Drosophila adult and larvae testis enables contrasting the different germline developmental stages Soumitra Pati, Amelie Raz, Sharvari Mahadevraju 1) National Center for Biotechnology Information, National Library of Medicine, National Institute of Health, Bethesda, MD, 20894, USA; 2) Laboratory of Cellular and Developmental Biology, National Institute of Diabetes and Kidney and Digestive Diseases, National Institutes of Health, Bethesda, MD, 20892, USA; 3) Whitehead Institute for Biomedical Research, Cambridge, MA, USA

Capturing the single nuclear and single cellular transcriptomics using snRNA-seq and scRNA-seq have advantages of their own. In snRNA-seq the message made by a cell at a certain time is captured whereas in scRNA-seq, the message stored for a longer period is captured as well. Similarly, RNA-seq profiles (sn or sc) captured from different stages of the life-cycle can reveal differences in the transcriptome across life stages. However, co-analyzing such diverse datasets together to gain biological insights poses significant challenges as the datasets could have batch effects eclipsing the biological differences, or even have significantly different RNA-features. Here we use an alternate approach where we first analyze the datasets separately utilizing then individual nuances and then contrast the analyses by a post-analysis alignment procedure. Specifically, we align the trajectory of germ cells from different data sets and set out to identify if there are any differences in the germ cell developmental stages revealed by the nuclear and whole cell transcriptomics at either life-stages.

We analyzed three datasets from different labs: 1) snRNA-seq on adult testis from Fly Single Cell Atlas, 2) scRNA-seq on adult testis and 3) scRNA-seq on larval testis. The snRNA-seq germline has 21,061 nuclei whereas the adult and larval scRNA-seq have 6,438 and 10,652 cells respectively. The trajectory analyses using monocle3 on these datasets separately arrange the germ cell that progresses from spermatogonia to spermatids via different developmental stages. However, to contrast these individual trajectories for uncovering differences in biology, we adapted Dynamic Time Warping (DTW). We were able to align the three different trajectories on a common pseudotime scale. The alignment of germline adult and larval scRNA-seq pseudotime revealed cell states in adults (elongating spermatids) that are absent in the larvae confirming the legitimate alignment. Buoyed by the accuracy of the alignment by DTW, we next focussed on the difference between snRNA-seq and scRNA-seq of adult germ cells and identified a set of genes for which the cell stops producing the RNAs in the nucleus while differentiating from spermatocyte to spermatid but shows purdurance of transcript in the cytoplasm. While some of these genes are known in literature, some are our novel findings that we set out to validate experimentally.

527V Identification of candidate regulators of transposable element (TE) expression from host gene/TE coexpression Matthew Lawlor, Weihuan Cao, Christopher Ellison Rutgers University

The small RNA pathways involved in Drosophila transposable element (TE) control are widely studied, but the role of
host transcription factors in regulating TE expression is not well understood. Prior studies show that related TE families show similar expression patterns and tissue specificity. Furthermore, many TEs show expression patterns biased to specific cell types, likely due to regulation by host-encoded trans-acting factors. We reasoned that TE expression should be associated with natural variation in expression of such host-encoded factors across genetic backgrounds in wild populations of Drosophila. To test this hypothesis, we quantified host gene and TE expression in a previously published RNA-seq study of gene expression in 200 Drosophila Genetic Reference Panel strains. We employed linear models to identify significant associations between TE expression and host genes while accounting for confounding factors such as TE copy number and intronic insertions. We additionally correct for a systematic bias affecting highly expressed genes. Our approach identifies 34 previously reported piRNA pathway genes which show negative associations with three or more TEs. We additionally identify 32 transcription factors and cofactors (including seven unnamed genes) showing exclusively positive associations with three or more TEs in both sexes. Another 31 transcription factors and cofactors (including three unnamed genes) are negatively associated with TE expression. For 11 of these 63 candidates, we were able to confirm their effect on TE expression using publicly available RNA-seq after RNAi knockdown in S2R+ cells. We are in the process of performing in-vivo, tissue-specific RNAi knockdowns of additional candidates. These novel TE regulators will shed light on many aspects of host/TE co-evolution including mechanisms by which TEs co-opt host transcription factors to mobilize in the germline as well as both antagonistic and synergistic interactions between host gene regulatory networks and TE expression.

528V  Integration of BMP, JAK/STAT and EGFR signaling during anterior-posterior patterning of the follicular epithilum. Kelvin Ip, Kaitlin MacDonald, Scott De Vito, Mariana Fregoso Lomas, Laura Nilson McGill University

During development, interpretation of positional cues by individual cells generates stereotyped patterns of gene expression and cell fate. We study the Drosophila follicular epithelium, where cells integrate input from multiple extracellular signals to choose between discrete anterior and posterior cell fates. Central to the patterning of this epithelium is the localized activation of the epidermal growth factor receptor (EGFR) by a ligand, Gurken (Grk), secreted by the underlying oocyte. Early in oogenesis, Grk is localized at the posterior and induces expression of the transcription factor Midline (Mid). Later, Grk moves to the dorsal anterior, where it instead induces the transcription factor Mirror (Mirr). The choice between these alternative outcomes depends on the presence of additional localized secreted ligands. Posterior cells experience Grk in the presence of the JAK/STAT pathway ligand Unpaired (Upd), which cooperates with Grk to induce mid while also independently repressing mirr. At the anterior, the BMP signaling ligand Decapentaplegic (Dpp) cooperates with Grk to induce mirr, while independently repressing mid. In addition, there is mutual repression between mid and mirr. We propose that this joint activation and derepression generates a bistable choice between these two EGFR targets, but how a cell interprets and integrates these signals remains unknown.

We address this question by characterizing reporters bearing putative cis-regulatory regions (CRRs) from the mid and mirr loci. We discovered that, for each gene, the response elements for the known regulatory inputs are distributed across two reporters bearing non-overlapping CRRs. Dpp response elements are located in a different mid CRR than those for Upd or Mirr, and Upd response elements are located in a different mirr CRR than those for Dpp or Mid. Reporters with these CRRs differ in expression boundary and dynamics, providing a tool for assessing the contribution of different regulatory inputs to mid and mirr expression. Further analysis of these CRRs suggests that repression of mirr and mid by JAK/STAT signaling and BMP signaling, respectively, requires binding of the signal mediators to cis-elements. We will be investigating the interplay between the different regulatory inputs to mid and mirr and their role in patterning by manipulating these cis-elements in different sequence contexts.

529V  Myc-regulated miRNAs modulate p53 expression in Drosophila. Gervé Maria Paula, Sanchez Juan, Ingaramo Clara, Dekanty Andres IAL (instituto de agrobiotecnologia del Litoral)

Myc, a conserved transcription factor involved in the regulation of growth and metabolism, has been shown to regulate the biogenesis of miRNAs in cultured mammalian cells, however, the exact mechanisms by which Myc affects miRNA function remain unclear. Here we provide evidences that Myc directly regulates the expression of a high number of miRNAs in Drosophila. ChIP-seq analysis revealed that dMyc is highly enriched in the promoter region of 113 (50%) miRNA genes, and dMyc depletion showed reduced expression of most miRNA genes analysed. Along with reduced pri-miRNA expression, we observed decreased levels of pre- and mature miRNAs and increased expression of miRNA activity sensors (mir-GFP) in myc-depleted cells. Conversely, Myc overexpression increased miRNA levels and reduced miR-GFP expression, strongly suggesting that dMyc modulates the expression and processing of miRNAs. We also show that Myc-dependent regulation of miRNA biogenesis plays a critical role in the response to nutrient stress. Dmp53, the single Drosophila ortholog of mammalian p53, is negatively regulated by miR-305 in the fat body in a nutrient-dependent manner, and its activation is required for maintaining metabolic homeostasis and promoting survival under nutrient deprivation. Our results revealed that dMyc directly binds miR-305 locus and promotes its expression, thus maintaining low Dmp53 levels in the fat body of well-fed animals. Under starvation conditions, however, dMyc protein levels and miR-305 expression are reduced which result in increased Dmp53 levels. These findings demonstrate an essential role for Myc in regulating miRNA expression and highlight the importance of Myc-dependent regulation of miRNA biogenesis in
metabolic homeostasis and organismal survival upon nutrient stress.

530V  The NXF gene family in Drosophila: Evolutionary History and Cell-Type Specific Gene Expression  
Martin Calvino, Jae Hak Son, Christopher Ellison  
The Human Genetics Institute of New Jersey, Rutgers University

In eukaryotes, mRNAs are transported from the nucleus to the cytoplasm by a family of nuclear RNA export factors (NXF). Members of this protein family are characterized by an amino-terminal region (NTR), an RNA-binding domain (RBD), leucine-rich repeats (LRRs), the NFT2-like domain (NTF2), and a Ubiquitin-associated (UBA) domain. In Drosophila melanogaster, the NXF family is composed of four members that have undergone functional diversification. While Nxf1 is involved in nucleocytoplasmic mRNA export, Nxf2 and Nxf3 were co-opted for piRNA-guided transcriptional silencing of transposable elements (TE). The role of Nxf4 remains to be elucidated. While Nxf1 is ubiquitously expressed, Nxf2 and Nxf3 are predominantly expressed in gonads; and Nxf4 displays testis-specific expression only. In this study, we describe the evolutionary diversification of the NXF gene family across more that 100 species of Drosophila and study branch-specific rates of evolution and test for evidence of positive selection among members of this family. We additionally analyzed publicly available single-cell RNA-seq data to identify cell-type specific expression profiles for NXF family members. Our results provide important insight in the evolution of the NXF gene family and its role in transposon suppression.

531V  Social experience and pheromone receptor activity reprogram behavioral switch gene splicing and 
neuromodulatory gene expression in sensory neurons  
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1 Department of Neurobiology, Duke University, Durham; 2 Department of Biology, Duke University, Durham; 3 Department of Biology, University of Chapel Hill, Chapel Hill

Social experience and pheromone signaling in pheromone sensing olfactory neurons affect pheromone responses and male courtship behaviors in Drosophila. We previously showed that social experience and pheromone signaling modulates chromatin around behavioral switch gene fruitless, which encodes a transcription factor necessary and sufficient for male courtship behaviors. Fruitless drives social context dependent modulation of courtship behaviors and pheromone responses in sensory neurons, however, the molecular mechanisms underlying this circuit-level neuromodulation remain less clear. To identify the molecular mechanisms driving social experience-dependent neuromodulation, we performed RNA-seq from antennal samples of mutants in pheromone receptors and fruitless, as well as grouped or socially isolated wild-type males. We found that loss of pheromone detection differentially altered the usage levels of fruitless exons, suggesting changes in splicing patterns. In addition, many Fruitless target neuromodulatory genes, such as neurotransmitter receptors, ion channels, and ion transporters, were differentially regulated by social context and pheromone signaling. Recent studies showed that social experience and juvenile hormone signaling coregulated fru chromatin to modify pheromone responses in olfactory neurons. Interestingly, genes involved in juvenile hormone metabolism were also misregulated in different social contexts and mutant backgrounds. Our results suggest that modulation of circuit activity and behaviors in response to social experience and pheromone signaling arise due to large-scale changes in transcriptional programs for neuromodulators downstream of behavioral switch gene function.

532C  A tale of two functions: Epigenetic programming and RNA splicing by Tip60 histone acetyltransferase  
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Reduced histone acetylation causes chromatin packaging alterations in neurons with concomitant transcriptional dysregulation that is a key initial step in Alzheimer’s disease (AD) etiology. In this regard, we have previously established a central role for Tip60 histone acetyltransferase (HAT) mediated chromatin dynamics in neuronal function, cognition, and AD. However, it remains to be elucidated if Tip60 has alternative cellular functions other than histone acetylation that could contribute towards AD pathogenesis. Altered RNA splicing has recently been highlighted as a widespread hallmark in AD transcriptome that is implicated in the disease. Strikingly, we discovered a novel RNA binding function for Tip60 in the Drosophila brain that is conserved in the human hippocampus and impaired in brains from both, AD fly models and AD patients. Our transcriptomic analysis of the RNA molecules specifically bound to Tip60 by Tip60-RNA immunoprecipitation (RIP) revealed an RNA binding function for Tip60 that was highly specific, selective, and reproducible, with Tip60 RNA targets enriched for critical neuronal processes that are implicated in AD. Remarkably, 79% of Tip60’s RNA targets overlap with its chromatin gene targets, supporting a model by which Tip60 orchestrates bi-level transcriptional regulation at both the chromatin and RNA level, a function unprecedented for any HAT to date. Since RNA splicing typically occurs co-transcriptionally and defects are a recent hallmark of AD, we hypothesized that Tip60 regulates splicing of its RNA targets and that this process is disrupted in AD. Transcriptome analysis from Drosophila brains that model AD showed reduced Tip60 levels and intriguingly, revealed that the residual Tip60 targets a different set of RNA when compared to wild-type flies. Additionally, we observed significant Tip60-associated RNA splicing defects in the Drosophila AD brain, some of which were prevented by increasing Tip60 levels in the brain. Thus, we are the first to uncover distinct histone and RNA binding capabilities for Tip60 that mediate its function in
neural gene control and RNA splicing, respectively, and may underly the chromatin packaging and splicing defects that are now characterized as hallmarks of AD.

533A  Temporal regulation of neuronal maturation by a chromatin anti-looping factor Dahong Chen, Catherine McManus, Behram Radmanesh, Leah Matzat, Elissa Lei NIH

The nervous system undergoes dramatic post-mitotic reorganization to become fully mature, but how chromatin 3D structure of neurons is regulated to assure proper transcriptomic dynamics during this essential process remains undefined in any organism. The tissue-specific chromatin insulator antagonist Alan Shepard (Shep) promotes Drosophila post-mitotic maturation by repressing expression of master regulator genes specifically in maturing neurons. To understand the mechanism of Shep repression of a key target gene, brain tumor (brat), we performed 4C-seq on CNS-derived BG3 cells and found that Shep depletion leads to increased brat promoter looping to proximal regions. Subsequent luciferase reporter assays verified enhancer activity of one of these candidate regions. Notably, depletion of Shep does not affect either enhancer or promoter activities in this artificial juxtaposed context, suggesting that Shep acts as a dedicated anti-looping factor to inhibit brat transcription. Interestingly, in vivo 3C also detected Shep inhibition of looping between the brat promoter and a proximal genomic region in pupal but not larval brains. Subsequent in vivo GFP reporter assays detected enhancer activity of this region specifically in pupal brains. This enhancer activity was not affected by Shep depletion, consistent with an anti-looping mechanism underlying Shep repression of brat transcription in pupal maturing neurons. Moreover, ATAC-seq and CUT&Tag for H3K4me1 on FACS-sorted neurons indicated that Shep is required to restrain chromatin accessibility of the brat enhancer as well as 1,126 other enhancers in pupal but not larval neurons, suggesting extensive Shep-mediated enhancer closure during neuronal maturation genome-wide. These enhancers are highly enriched for Shep-bound loci that are also transcriptionally inhibited by Shep, suggesting temporal Shep inhibition of enhancer accessibility to repress gene expression. Taken together, our results provide the first evidence for a chromatin anti-looping factor that regulates temporal enhancer accessibility and gene expression during organismal development.

534B  Sex-specific variation in R-loop formation in Drosophila melanogaster Timothy Stanek1,2, Rohan Mehra3, Weihuan Cao1, Christopher Ellison1 1) Department of Genetics, Rutgers University, Piscataway, NJ; 2) Department of Pathology, Robert Wood Johnson Medical School, Piscataway, NJ

R-loops are three-stranded nucleotide structures consisting of a DNA:RNA hybrid and a displaced ssDNA non-template strand. Originally viewed as byproducts of transcription, R-loops are now recognized as important regulators of gene expression and genomic stability. Persistent dysregulation of R-loop maintenance can result in replication stress, DNA double-strand breaks, and chromosomal rearrangements that contribute to diseases such as neurological disorders and cancer. Although R-loops are conserved across cell types in mammals, little is known about natural variation in R-loop formation between individuals.

Using DNA:RNA immunoprecipitation followed by high-throughput sequencing (DRIP-seq), we have mapped the R-loop profiles of two D. melanogaster individuals from the Drosophila Genetic Reference Panel (DGRP) in both males and females. R-loops are largely conserved across individuals and between sexes. Consistent with previous studies, R-loops are found at 5'UTRs of protein-coding genes and across many classes of noncoding RNA. R-loops are also enriched at Polycomb response elements and topologically associating domain boundaries. More broadly, R-loops are enriched in both the active RED and Polycomb Group BLUE chromatin states and depleted from GREEN heterochromatin and BLACK repressive states. Differential enrichment analysis reveals a small number of sex-specific R-loops: while non-differentially enriched and male-enriched R-loops form at similar genetic features and chromatin states and contain similar sequence motifs, female-enriched R-loops are enriched at unique genetic features, chromatin states, and sequence motifs. Male-enriched R-loops are most abundant on the dosage-compensated X chromosome, where R-loops appear stronger compared to autosomal R-loops. However, where most R-loop-containing genes exhibit higher levels of expression compared to R-loop-absent genes, male-enriched R-loops on the X chromosome do not correspond to increased gene expression. These male-enriched R-loops also show lower MOF binding and reduced H4K16ac, suggesting that canonical marks of hypertranscription associated with dosage compensation are antagonistic to R-loop formation. Collectively, these results reveal a series of sex-specific characteristics of R-loop formation in vivo and suggest that these hybrid structures can act to the benefit or detriment of specific cellular processes.

535C  HDAC-inhibitory Microbial Volatiles Effect on Slowing Huntington’s Disease in a Drosophila model Rogelio Nunez Flores, Sachiko Haga-Yamanaka, Christi Ann Scott, Anandasankar Ray University of California, Riverside

Eukaryotes coevolve with microbiomes and respond to their secreted metabolites. However, the response to volatile compounds emitted by microbes is poorly understood. We show that a microbial volatile, diacetyl, and other structurally related odorants, can alter gene expression in eukaryotes at a distance from their emission source. These compounds were found to inhibit human histone-deacetylases (HDACs) leading to increased histone-H3K9 acetylation. Most notable, these compounds inhibit HDAC6, a promising up-regulated target in several neurodegenerative diseases. These
epigenetic alterations explain the significant changes in gene expression/physiological changes seen in Drosophila melagonaster models. In a Drosophila melanogaster transgenic model for Huntington’s disease in the fly eye, exposure to the volatiles slowed progression of neurodegeneration similarly to known orally administered HDAC-inhibitor drugs. When we repeated the exposure study in HDAC6[KO] flies bearing the Huntington’s Disease transgene, we also observed similar neuroprotective effects. Our findings suggest that these changes are caused by a conserved atypical signaling pathway that modulates gene expression via changes in chromatin from a distant chemical source.

536A  Unique chromatin characteristics allow a genome-eliminating B chromosome to avoid self-elimination Salina Teklay¹, Haena Lee¹,³, Pooreum Seo¹, Emily Yuguchi², Elena Dalla Benetta¹,³, John Werren², Patrick Ferree¹ 1) Claremont Colleges, Claremont, CA; 2) University of Rochester, Rochester, NY; 3) UC San Diego, San Diego, CA

Insects harbor a wide array of reproductive parasites, including selfish B chromosomes. One of the most dramatic examples is a B chromosome known as PSR (for Paternal Sex Ratio), which resides in natural populations of the jewel wasp, Nasonia vitripennis. PSR is paternally transmitted via the sperm’s nucleus to progeny, and it causes complete destruction of the sperm’s hereditary material, but not itself, during the first embryonic mitotic division. This action converts female-destined embryos into haploid males, which can then transmit PSR. A compelling question is how PSR escapes its own genome-eliminating activity. We previously found that while PSR causes heightened H3K27me1 and H4K20me1 across the paternal genome just before it is eliminated, PSR itself does not incur these marks. Here we have found that the segregation of PSR is only partially affected by the CI activity of Wollbachia. While Wollbachia like PSR causes heightened H3K27me1 and H4K20me1 across the paternal genome, PSR remains unmarked regarding these histone modifications in the presence of this chromatin-altering bacterium. These findings suggest that PSR’s chromatin exhibits unique properties compared to the essential genome complement. We hypothesized that PSR may not participate in the histone-to-protamine transition as a way of evading chromatin disruption. However, this idea was not supported; when the histone chaperone HIRA was knocked down by systemic RNAi, PSR completely failed to segregate. Our current efforts are aimed at examining other chromatin-related characteristics to explain PSR’s intriguing ability to escape its own genome eliminating activity.

537B  Determining how H4K20 methylation contributes to L(3)mbt recruitment to chromatin Megan B. Butler, Aaron T. Crain, Robert J. Duronio University of North Carolina at Chapel Hill, Chapel Hill, NC

Generation and maintenance of specific chromatin domains are essential for proper genome regulation, cell cycle progression, and organisal development. Chromatin is a complex of DNA wound around histone proteins which have N-terminal tails that can be chemically modified post-translationally (PTMs). Proteins termed “readers” bind histone PTMs and either directly or as part of multi-protein complexes alter chromatin structure to control critical genome functions. One such reader discovered in Drosophila melanogaster and called Lethal (3) malignant brain tumor (L(3)mbt) functions as a tumor-suppressor in larval brains. L(3)mbt has been implicated in chromatin condensation and transcriptional regulation. Previous in vitro studies showed that L(3)mbt and its human homolog preferentially bind histone H4 that is methylated on lysine 20 (H4K20me). However, in vivo studies have shown that L(3)mbt does not exclusively co-localize with H4K20me throughout the genome; thus, how L(3)mbt interacts with chromatin and exerts its functions remains incompletely understood. Determining whether L(3)mbt depends on H4K20me for binding the genome in vivo has never been addressed in the most direct way – by mutating H4K20. With our histone gene replacement platform in Drosophila, we generated H4K20A and H4K20R mutants to assess the consequence of eliminating H4K20me on L(3)mbt interaction with the genome. In order to perform this experiment, we built GFP- and FLAG-tagged alleles of l(3)mbt at the endogenous locus using CRISPR-Cas9 and the Scarless Gene Editing system. These alleles complement a null mutation of l(3)mbt, indicating that the epitope-tagged proteins are fully functional. We assessed L(3)mbt’s cytological localization in wildtype and H4K20-mutant animals via confocal microscopy using anti-GFP and anti-FLAG antibodies. In wildtype larval brains, imaginal discs, and adult ovaries, L(3)mbt is predominantly nuclear during interphase with much of the protein residing in the nucleoplasmic compartment. During mitosis, L(3)mbt disperses throughout the cell after nuclear envelope breakdown and does not localize to condensed mitotic chromosomes. In H4K20-mutant animals, we detected a small decrease in total L(3)mbt signal without any change in L(3)mbt localization. These results indicate that we are unable to use cytology alone to make conclusions about whether H4K20me is necessary for L(3)mbt chromatin binding. Therefore, we will next use approaches like CUT&RUN to determine the association of these epitope-tagged L(3)mbt proteins with the genome in wildtype and H4K20-mutant animals at a high resolution.

538C  Interrogating the roles of canonical versus variant histone H3 in genome function and aging Jeanne-Marie McPherson, Robert Duronio, Daniel McKay University of North Carolina at Chapel Hill

Histones are essential for packaging and organizing DNA into chromatin, which regulates all DNA-dependent processes. Regulation of histone abundance is critical, as too many or too few histones is toxic to cells and disrupts development. Cells contain two types of histones: canonical histones expressed during S phase of the cell cycle and variant histones expressed throughout the cell cycle and in post-mitotic cells. Canonical histone H3.2 and variant histone H3.3 are highly

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conserved across eukaryotes, making *Drosophila melanogaster* an ideal model to study histone biology. By manipulating canonical H3.2 copy number in *Drosophila* we discovered that H3.3 is essential for development when canonical histone gene copy number is reduced, uncovering a previously unknown requirement for coordination between canonical H3.2 and variant H3.3 expression. We hypothesize that cells possess a homeostatic mechanism to maintain the correct relative expression of canonical H3.2 and variant H3.3. Histone homeostatic mechanisms could affect histone expression at many levels, including transcription, translation, deposition, or protein turnover. Thus far we have discovered that reducing canonical H3.2 copy number does not result in increased variant H3.3 transcript or protein levels. We will explore the mechanisms of histone homeostasis by conducting a forward genetic screen to identify factors involved in canonical and variant histone coordination. This project expands our fundamental understanding of how canonical and variant histones cooperate to regulate genome function.

539A  A novel mosaic system for performing forward genetics in a sensitized histone mutant background Aaron T. Crain, Robert J. Duronio University of North Carolina at Chapel Hill, Chapel Hill, NC

The mono-methylation of lysine 20 of histone H4 (H4K20me1) is involved in numerous processes necessary for cell proliferation, including DNA replication, gene expression, and chromosome condensation during mitosis. These functions of H4K20me1 have been largely derived from interpreting the phenotypes resulting from manipulation of the activity of Set8, the lysine methyltransferase responsible for depositing H4K20me1. However, such interpretations are complicated by studies demonstrating that Set8 has several non-histone substrates with essential roles in cell proliferation, such as p53 and PCNA, as well as non-catalytic functions. To directly assess the contribution of H4K20me in cell proliferation, we engineered a novel system for generating mosaic tissues containing clones of histone mutant cells. We first created a new, precise deletion of the replication dependent histone locus on chromosome 2 (HisC) that is marked with Act5c-dsRed. This allele was then combined with an established transgenic histone replacement platform in which a transgene containing a synthetic histone gene array capable of complementing homozygous HisC deletions is integrated on chromosome 3. These transgenes are engineered to express either wild-type (WT) or mutant (e.g. H4K20A and H4K20R) histones. We then use FLP/FRT-mediated mitotic recombination to produce dual-color mosaic tissue in which a transgene containing a synthetic histone gene array capable of complementing homozygous HisC deletions is integrated on chromosome 3. These transgenes are engineered to express either wild-type (WT) or mutant (e.g. H4K20A or H4K20R) histones. We then use FLP/FRT-mediated mitotic recombination to produce dual-color mosaic tissue in the *Drosophila* eye, where one color (green) labels cells expressing wild-type histones and the other color (red) labels cells expressing only mutant histones. Thus, we can easily visualize and quantify competitive proliferation of histone mutant cells. Strikingly, we found that both H4K20A and H4K20R mutant cells can proliferate adjacent to wild-type cells. Strikingly, we found that both H4K20A and H4K20R mutant cells can proliferate adjacent to wild-type cells, achieving 34% and 22% of the adult eye, respectively, compared to 42% for H4WT. In contrast, we found that Set8null mutant cells cannot proliferate adjacent to wild-type cells and die. We interpret the lack of proliferation and death of Set8null cells as likely due to H4K20me-independent mechanisms, and thus conclude that the role of H4K20me in cell proliferation remains unknown. To discover the functions of H4K20me in cell proliferation we will utilize our newly established mosaic system to perform a forward genetic screen for modifiers of proliferation of H4K20 mutant cells. This work establishes the first platform for performing forward genetics using histone mutant cells in animals.

540B  Identification of factors involved in rDNA magnification in the male germline Alyssa Slicko1,2, Jonathan Nelson¹, Yukiko Yamashita1,2,3 1) Whitehead Institute, Cambridge, MA; 2) Howard Hughes Medical Institute; 3) Department of Biology, MIT, Cambridge, MA

An important role of the germline is to maintain unstable, but essential genomic elements to ensure the transmission of functional genomes across generations. Among these elements are ribosomal DNA (rDNA), which consists of hundreds of tandem repeats of the genes that are needed for sufficient ribosomal activity. rDNA is known to be one of the most unstable regions in the genome because this repetitive structure is susceptible to deleterious recombination. This instability is countered in the germline to prevent the progressive multi-generational loss of rDNA, but the mechanisms that underlie this feature have remained unknown. Our previous work suggests that rDNA is maintained by copy number expansion in the germline that recovers copies lost in the previous generation, though the factors that achieve this expansion remain largely unknown. Here we describe our efforts to identify these factors through RNAi-mediated knockdown of candidate genes.

We used a well-described system of robust rDNA copy number expansion in the male germline called “rDNA magnification” to identify factors required for rDNA expansion. Previous work revealed that animals with very little rDNA, meaning no rDNA on Y and partial rDNA on X, have the bobbed phenotype, which displays disrupted dorsal cuticle patterning. rDNA magnification is the phenomenon that a small amount of offspring from bobbed males have wild-type cuticles due to the rDNA on the partially deleted X chromosome having been expanded. We tested the ability for RNAi knockdown of candidate genes to prevent rDNA magnification in bobbed males.

We selected candidate factors of known rDNA-binding proteins, and homologs of factors that aid in rDNA maintenance in yeast, to test with the RNAi. We preliminarily identified that an uncharacterized gene on the Y chromosome, PRY (Polycystine-related Y), contributes to rDNA magnification. PRY was tested as a candidate factor since it resides in...
a portion of the Y chromosome previously identified to be necessary for rDNA magnification. We discovered that the transgenic expression of PRY can induce rDNA magnification in both the male and female germline. We are now investigating the activity of PRY in the germline to understand how it contributes to rDNA magnification.

541C Genomic insertion of repetitive DNA can trigger conversion of euchromatin to heterochromatin Safiyo Aden, Derrick carper, Heidi J.J Pipkin, Andrew M Arsham Bemidji State University

Heterochromatin is a key genomic defense against invasive genetic elements, mitigating damage by inhibiting transposition, silencing gene expression, and reducing recombination at insertion sites. How genomes recognize and silence novel threats prior to establishing sequence-specific adaptive defenses like piRNAs is poorly understood. To investigate genome defense against novel repetitive DNA we carried out a transposition mutagenesis screen with a reporter construct expressing the white gene adjacent to a 256-copy tandem array of a 36 nt lac operator sequence from E. coli. Reporter gene expression was robust in the vast majority of transposition mutants recovered, while approximately 1% had variegated eye color suggesting epigenetic silencing by heterochromatin. Mapping of insertion sites by inverse PCR revealed that the lacO reporter construct can trigger silencing of actively transcribed euchromatin. Excision of the lacO array by FLP-FRT recombination suppressed variegation, restoring full reporter gene expression and demonstrating repeat-dependent silencing. Ectopic heterochromatinization a lacO tandem array may model disease processes like the silencing of triplet nucleotide expansions and evolutionary processes like centromere formation.

542A Role of Ulp1, a SUMO E3 Protease, in enabling ‘safe’ Homologous Recombination progression at the nuclear periphery. Nadejda Butova, Irene Chiolo, Chiara Merigliano, Taehyun Ryu, Sydney Prange University of Southern California

Pericentromeric Heterochromatin occupies ~30% of fly and human genomes and is mostly composed of highly repeated DNA sequences prone to aberrant recombination. And yet, repair mechanisms in heterochromatin remain poorly understood. Previous studies from the Chiolo lab in Drosophila cells showed that ‘safe’ homologous recombination (HR) repair of heterochromatic double-strand breaks (DSBs) requires the movement of repair sites away from the heterochromatin ‘domain’ (a distinct nuclear structure in fly cells) to the nuclear periphery, where the strand invasion protein Rad51 is recruited to continue repair. This is believed to prevent aberrant recombination by isolating broken sites and their homologous templates away from ectopic (non-allelic) repair templates before strand invasion. The post-translational modification SUMOylation is essential for this pathway, as it blocks Rad51 recruitment and HR progression inside the heterochromatin domain thus preventing aberrant recombination. Furthermore, work in the Chiolo lab has shown that Ulp1 overexpression leads to abnormal Rad51 recruitment to repair sites inside the HC domain. What restarts HR at the nuclear periphery, the SUMOylation targets, and the role of SUMOylation in repair are major open questions in the field. By identifying these targets and their role in repair of ionizing radiation (IR)-induced DSBs, our studies will unravel new molecular mechanisms preventing chromosome rearrangements and genome instability in heterochromatin, which are still obscure, as well as establish the role of Ulp1 in the spatial and temporal regulation of heterochromatin repair. Through the use of Mtor/Tpr depletions and Ulp1AN mutants that lose the nuclear-pore association domain, thus releasing Ulp1 in the nucleoplasm, my results show that Mtor/Tpr is required to recruit Ulp1 to the nuclear periphery where it is able to complete repair. By testing the role of Ulp1 at the nuclear periphery using Ulp1 depletions by RNAi or degrons, the loss of Rad51 foci at heterochromatic DSBs (marked by γH2Av foci associated with heterochromatin marks), at the nuclear periphery in the absence of Ulp1 was observed. This supported the role for Ulp1 in HR progression once DSBs have relocalized. Overall, these results support the hypothesis that Ulp1 compartmentalization to the nuclear periphery is necessary for preventing abnormal HR progression inside the heterochromatin domain, while enabling ‘safe’ HR progression at the nuclear periphery.

543B Investigating the origin and evolution of CG17359, rapid-evolving, essential ZAD-ZNF gene in multiple Drosophila species Madeline Gruys1,2, Zainab Abdulrahman1,2, Ashlyn Anderson1,2, Safiyo Aden1,2, Solomon Aviles1,2, Derrick Carper1,2, Nathan Dupre1,2, Jack Jurmu1,2, Tetiana Khotko1,2, Zahraa Lami1,2, Hunter Lindsay1,2, Tea Merkl1,2, Heidi J. J. Pipkin1,2, Dennis Quach1,2, Anthony Ruiz1,2, Melissa K. Sawyer1,2, Noah Stone1,2, Bailey Taylor1,2, Barbara Whitlock1,2, Christina Yang1,2, Emily Yang1,2, Andrew M Arsham1) 1) Bemidji State University; 2) North Hennepin Community College

Ancient essential genes are highly conserved due to strict functional constraints that limit their evolutionary potential. It has become clear that young and rapidly evolving genes can also acquire new functions and become essential. The evolution of some of these genes may be driven by the dynamic nature of heterochromatin and to maintain silencing of rapidly-changing non-coding heterochromatic sequences. The evolutionarily dynamic ZAD-ZNF gene family encodes the most abundant class of insect transcription factors, many of which appear to be the product of relatively recent segmental duplication or retrotransposition. CG17359 and CG17361 are adjacent paralogous ZAD-ZNF genes that appear to have formed by retrotransposition between 30 and 40 million years ago. CG17359 is essential in D. melanogaster and shows evidence of positive selection, while CG17361 is not essential and does not show evidence of strong selective
pressure. We have identified a third nearby ZAD-ZNF gene, CG8474/Meics, as a candidate ancestor for both. Using the gene annotation framework of the Genomics Education Partnership, we annotated sequence conservation and synteny for all 3 genes across Drosophila species spanning 40 million years of evolution. The quickly-evolving yet essential gene annotation framework of the Genomics Education Partnership, we annotated sequence conservation and synteny. We have identified a third nearby ZAD-ZNF gene, CG8474/Meics, as a candidate ancestor for both. Using the

544C Towards telomere-to-telomere genome assemblies of Drosophila melanogaster

The goal of genome sequencing and assembly is to capture all sequence features that play critical roles in organisms and to join them together as parts of complete, gapless chromosome reference assemblies. However, in Drosophila melanogaster, the most complete assemblies are at least 15-20% smaller than the known genome size. The segments missing from assemblies are almost entirely composed of repetitive sequences, primarily transposable elements and satellite repeats concentrated in pericentric heterochromatin. These missing regions not only contain essential genes, they also harbor other elements that play crucial roles in genome stability, chromosome segregation, protein translation, and TE repression. To recover these important regions, we are building telomere-to-telomere assemblies of multiple D. melanogaster strains, including the genome reference strain iso-1. Our goal is to produce complete, extremely accurate (fewer than 1 error per megabase) assemblies for all three autosomes and the X and Y chromosomes. Efforts by the human Telomere-to-Telomere (T2T) Consortium have pioneered the completion of virtually gapless chromosome assemblies that span large repetitive arrays, including satellites, scrambled transposable elements, and ribosomal DNAs. These approaches leverage ultra-long sequencing reads to untangle assembly graphs derived from highly accurate long sequence reads. By applying this approach to additional strains, we can study variation in chromosome structures that have previously resisted scrutiny, like centromeres. This open, collaborative initiative aims to produce a gapless assembly of D. melanogaster, outline best practices for extending this approach to other strains and species, and support public accessibility of data releases and methodologies.

545A Silencing and position-effect variegation in a dual-reporter transposition mutagenesis screen

Highly repetitive DNA sequences are often associated with invasive or pathogenic DNA, and can also be found in gene-poor regions of eukaryotic genomes like centromeres and telomeres. Genomes often defensively package repeats in dense, transcriptionally refractory, stably heritable heterochromatin which plays a crucial role in the regulation of gene expression through transcriptional silencing. How these sequences trigger the establishment and maintenance of heterochromatin remains poorly understood, but repeats appear to serve as a signal of foreignness to host genomes and can activate the siRNA, miRNA, or piRNA silencing pathways. Silencing of repetitive DNA is implicated in the formation and function of centromeres and telomeres, in defense against transposons and viruses, and in human trinucleotide repeat expansion diseases like Friedreichs Ataxia and Fragile X syndrome. To investigate the mechanism and regulation of repeat-triggered silencing, undergraduate students participating in a classroom research experience (CURE) carried out a transposition mutagenesis to investigate the effects of large tracts of repetitive DNA on gene expression and chromatin state. Students mobilized a p-element containing a 256-copy tandem array of the E. coli lac operator (LacO), flanked by white and yellow reporter genes. After setting up mobilization crosses, students screened F2 progeny for changes in eye or body color indicating expression of either or both reporter genes (287/14137 or 2% of all male F2 screened). Of transgene-expressing flies, wild type expression of yellow and white comprised 82% of all mutants and indicate presumed euchromatic insertions resulting in high reporter gene expression. The white gene alone was silenced in 11% of identified mutants; the yellow gene alone was silenced in 4%, demonstrating a surprising decoupling of reporter gene silencing. 3% of mutants showed variegation of the white gene and full expression of the yellow gene, but the converse was not observed, nor was variegation of body color. Molecular mapping of insertion sites with inverse PCR suggest that LacO repeat insertion can trigger silencing even in gene-rich euchromatic locations.

546B The Phosphorylated Histone Variant H2Av Associates With Gypsy Insulator Proteins Through Liquid-Liquid Phase Separation

Highly repetitive DNA sequences are often associated with invasive or pathogenic DNA, and can also be found in gene-poor regions of eukaryotic genomes like centromeres and telomeres. Genomes often defensively package repeats in dense, transcriptionally refractory, stably heritable heterochromatin which plays a crucial role in the regulation of gene expression through transcriptional silencing. How these sequences trigger the establishment and maintenance of heterochromatin remains poorly understood, but repeats appear to serve as a signal of foreignness to host genomes and can activate the siRNA, miRNA, or piRNA silencing pathways. Silencing of repetitive DNA is implicated in the formation and function of centromeres and telomeres, in defense against transposons and viruses, and in human trinucleotide repeat expansion diseases like Friedreichs Ataxia and Fragile X syndrome. To investigate the mechanism and regulation of repeat-triggered silencing, undergraduate students participating in a classroom research experience (CURE) carried out a transposition mutagenesis to investigate the effects of large tracts of repetitive DNA on gene expression and chromatin state. Students mobilized a p-element containing a 256-copy tandem array of the E. coli lac operator (LacO), flanked by white and yellow reporter genes. After setting up mobilization crosses, students screened F2 progeny for changes in eye or body color indicating expression of either or both reporter genes (287/14137 or 2% of all male F2 screened). Of transgene-expressing flies, wild type expression of yellow and white comprised 82% of all mutants and indicate presumed euchromatic insertions resulting in high reporter gene expression. The white gene alone was silenced in 11% of identified mutants; the yellow gene alone was silenced in 4%, demonstrating a surprising decoupling of reporter gene silencing. 3% of mutants showed variegation of the white gene and full expression of the yellow gene, but the converse was not observed, nor was variegation of body color. Molecular mapping of insertion sites with inverse PCR suggest that LacO repeat insertion can trigger silencing even in gene-rich euchromatic locations.
Eukaryotic genomes are characterized by a highly orchestrated 3D organization. Binding of insulator proteins to specific DNA sequences through the genome contribute to genome structure by demarcating the boundaries between genome domains. The functional properties of insulator proteins, along with those of other architectural proteins, are under strong scrutiny given that the understanding of genome structure is essential to explain fundamental aspects of genome expression and genome maintenance. Loop extrusion mediated by Cohesin and chromatin-mediated liquid-liquid phase separation (LLPS) are thought to be the major driving forces contributing to domain formation and compartmentalization of the genome. The *Drosophila melanogaster* genome expresses an array of insulator proteins that constitute diverse insulator complexes targeted to different sequences across the genome, but how these insulator complexes specifically contribute to genome structure is currently unknown. Although insulator proteins function in the context of chromatin and nucleosomes, specific histone proteins have never been directly implicated in insulator function. Here, we demonstrate an interaction between gypsy insulator proteins and the phosphorylated form of the histone variant H2Av (γH2Av). We show that components of the gypsy insulator complex, Su(Hw), Mod(mdg4)67.2 and CP190 colocalize with γH2Av throughout the genome and mutation of insulator components prevents stable association of phosphorylated H2Av with chromosomes. Inhibition of the PP2A phosphatase results in a stronger chromosomal association of γH2Av and insulator proteins. Interestingly, γH2Av but not unphosphorylated H2Av associates to insulator bodies after osmotic stress, and phosphatase activity is required for insulator body dissolution after recovery from stress. Our evidence suggests a model in which phosphorylation of H2Av regulates insulator activity by modulating the LLPS properties of insulators and associated architectural proteins.

547C  Cardiac aging prevention through H3K27me3 modulation  Clara Guida, Georg Vogler, Peter Adams, Rolf Bodmer  Sanford Burnham Prebys Medical Discovery Institute

The incidence of heart failure approximately doubles through each decade of life. The detrimental effects of age on heart function are likely due in part to epigenetic dysregulation and associated gene program changes. Indeed, evidence already suggests that the maintenance of the epigenome becomes more error-prone with age, leading to so-called “epigenetic drift”, or accumulation of epigenetic alterations. Interestingly, manipulation of certain epigenetic modifiers was found to expand lifespan in different animal models. However, little is known about the epigenetic mechanism underlying cardiac aging. The challenge of studying cardiac aging relies in part on the low abundance of human samples and long and expensive experimental times for mammalian animal models. Here we used the *Drosophila* heart model, which has several advantages including short lifespan, less genetic redundancy, conserved biological pathways, and in vivo heart analysis protocols, to identify epigenetic mechanisms involved in cardiac aging. In a targeted genetic screen, we found that partial depletion of components of the Polycomb Repressive Complex 2 (PRC2) prevented cardiac aging. This is in line with previous findings reporting increased lifespan in PRC2-/+ flies. Indeed, we found that H3K27me3 content is increased in cardiomyocytes with age in control flies. Moreover, age-related cardioprotection was also achieved when we treated flies with an inhibitor of the H3K27me3 methyltransferase, EZH2. These findings suggest that H3K27me3 marks can mediate age-related heart function deterioration and establish the *Drosophila* model as a useful tool to investigate the basic biology of age-related cardiac epigenetic drift.

548A  Investigating the role of Polycomb repression in *Drosophila* eye specification  Haley Brown, Justin Kumar  Indiana University, Bloomington, IN

During metazoan development, gene regulatory networks (GRNs) are activated in undifferentiated tissues to induce a specific fate. However, when GRNs are disrupted, the tissue can *transdetermine* – losing the programmed fate to adopt another. Epigenetic factors, such as the Polycomb Group (PcG) proteins, ensure proper spatiotemporal control of GRNs. PcG proteins function as a set of complexes to add a repressive histone mark (H3K27me3) and condense chromatin. In turn, the accessibility of chromatin – or lack thereof – regulates differential transcription of genes in certain tissues. While the correlation between GRNs and chromatin modifications in development is widely established, the underlying mechanisms linking the two during transdetermination has yet to be discovered. The overarching goal of this project is to determine how epigenetic modifications affect tissue fate specification. An excellent model to study the mechanisms underlying fate plasticity is the eye-to-wing transformation of *Drosophila* eye-antennal discs (EADs). Previous work from our lab discovered that the EAD-specific removal of one PcG protein, Polycomb (Pc), transforms the eye imaginal tissue to wing – indicating that the loss of epigenetic repression is sufficient to allow cellular reprogramming. To investigate the molecular mechanism underlying this transformation, I have performed RNA-seq on wild-type (WT) wing discs (WDs) as well as WT and *Pc* mutant EADs throughout third instar development. This analysis identified 55 candidate genes that could be responsible for promoting reprogramming of the EAD. My preliminary data suggest the most promising of these candidates is *vestigial* (*vg*), as this locus is directly regulated by *Pc*. Furthermore, overexpression of *vg* in the EAD grants an eye-to-wing transformation, and ectopic *vg* expression is detected in the developing wing pouch of the transformed disc. I will further investigate how repressive and active histone modifications are changing in this system to allow for reprogramming by using a novel epigenome profiling technique, CUT&RUN. I hypothesize that the inability of *Pc* mutants to read H3K27me3 marks allows the epigenome to become malleable, activates wing determination genes, and transforms the eye into a wing. Outcomes of this study will elucidate the mechanistic role
549B  The H3.3K27M oncohistone antagonizes reprogramming in Drosophila Kami Ahmad1, James Anderson1, Steven Henikoff2,3 1) Fred Hutchinson Cancer Research Center; 2) Howard Hughes Medical Institute

Development proceeds by the activation of genes by transcription factors and the inactivation of others by chromatin-mediated gene silencing. In certain cases development can be reversed or redirected by mis-expression of master regulator transcription factors; this reprogramming must involve the activation of new genes and the chromatin-mediated silencing of others. Here, we express the wing-specific Vestigial master regulator to reprogram the developing eye, and test the role of silencing in reprogramming using an H3.3K27M oncohistone mutation that dominantly inhibits histone H3K27 trimethylation. We find that production of the oncohistone blocks eye-to-wing reprogramming. CUT&Tag chromatin profiling of mutant tissues shows that H3K27me3 of domains is generally reduced upon oncohistone production, suggesting that a previous developmental program must be silenced for effective transformation. Strikingly, Vg and H3.3K27M synergize to stimulate overgrowth of eye tissue, a phenotype that resembles that of the oncohistone in human cancers. Transcriptome profiling of elongating RNA Polymerase II implicates the mis-regulation of signaling factors in overgrowth. Our results demonstrate that growth dysregulation can result from the simple combination of crippled silencing and transcription factor mis-expression, an effect that may explain the origins of oncohistone-bearing cancers. We have continued this work to search for additional factors that can synergize with the H3.3K27M oncohistone, and to probe the earliest epigenetic changes in cells as they begin to overproliferate.

550C  X marks the spot: Specifically targeting active chromatin to the X chromosome Joseph Aguilera, Mukulika Ray, Ashley Conard, Erica Larschan Brown University

Precise and coordinated regulation of gene expression during growth and development is essential for the viability of all organisms and to prevent diverse diseases ranging from cancer to neurodegeneration. Chromatin domains coordinate gene expression by concentrating key factors at specific genomic locations in a context-specific way to concordantly activate or repress groups of genes. Before zygotic genome activation (ZGA), chromatin domains have not yet formed and only a few key transcription factors (TFs) are bound which are called pioneer TFs. Pioneer TFs have the ability to bind to closed chromatin, recruit chromatin remodelers to open chromatin, and target additional transcription complexes. Furthermore, pioneer TFs are critical for cellular reprogramming and are often reactivated in cancer. Pioneer TFs can regulate the formation of active and repressive chromatin domains which form after ZGA. However, the fundamental mechanisms by which chromatin domains are formed at the correct genomic locations across time and space remain unknown. One example of a conserved active chromatin domain is the hyperactivated single male X-chromosome in Drosophila, which coordinately upregulates thousands of genes two-fold to equalize transcript levels with the two autosomes. My preliminary data suggest that the interaction between an essential pioneer TF and large transcription complex is important to target this active chromatin domain specifically to the X-chromosome. Therefore, I will use genomic, optogenetic, and machine learning approaches to define new mechanisms by which a pioneer TF nucleates the formation of an active chromatin domain that upregulates the male X-chromosome over time and space.

551A  Intercalary heterochromatin prevents local somatic pairing loss in interspecies Drosophila hybrids James Baldwin-Brown, Nitin Phadnis University of Utah

Homologous chromosome pairing is essential to all eukaryotes. Even so, our understanding of the molecular mechanisms underlying pairing is limited. We used changes in the pattern of pairing in Drosophila hybrids to find new mechanisms connected to pairing, and find that BLACK (intercalary) heterochromatin correlates with successful pairing. This discovery will help us understand pairing generally.

Although pairing is often associated with meiosis, it also occurs in somatic cells. Complete somatic pairing is the wild type state in Drosophila and other dipterans, but reproducible patterns of unpaired regions exist across the genome in interspecific Drosophila hybrids. Because chromosome pairing machinery exists in all eukaryotes, finding the genomic elements that drive this non-pairing will help us understand the drivers of pairing generally.

We crossed Drosophila melanogaster and Drosophila simulans, then used Hi-C to measure the rate of chromosome pairing with high resolution across the genome. Compared to within-species crosses, this hybrid Hi-C shows dramatic regions of high and low pairing. Repeating experiments in multiple tissues showed that hybrid pairing loss uniquely affects polytenic tissues. In keeping with results from mosquitoes, an inversion between D. melanogaster and D. simulans showed reduced pairing around the inversion breakpoints, serving as a positive control.

While past microscopy experiments showed the existence of non-pairing regions, our high-resolution methods show a complex tapestry of high- and low-pairing regions of varying breadths and intensities. The clearly resolved peaks are consistent with the “button” model of pairing (multiple, discrete pairing sites) rather than the “zipper” model (continuous pairing along the chromosomes). We showed that local pairing maxima and minima are apparently uncorrelated with expected pairing drivers such as sequence similarity and insulator binding site density. High pairing is, however, highly correlated with BLACK chromatin. This chromatin is underreplicated in tissues with multiply replicated
Paramutation is the phenomenon by which a silent allele can turn off a normal allele in trans in an epigenetic manner. The silenced state of the wildtype allele can persist through generations even in the absence of the original paramutagenic allele. The mechanisms underlying paramutation are poorly understood. Further, little is known about how paramutation shapes gene expression under natural conditions. In this study, we investigate a system of genomic paramutation in Drosophila virilis. Previous studies have shown that maternally transmitted piRNAs that target the center divider (cdi) gene in D. virilis have the capacity to silence expression of cdi in the next generation. In addition, it has been shown that piRNAs that target cdi can be maintained in subsequent generations in the absence of the original silencing allele. However, it is not known whether this pattern of piRNA biogenesis and maternal transmission coincides with epigenetic repression of cdi expression across multiple generations. To determine if cdi piRNA biogenesis mediates paramutation, we measured the expression of cdi in the ovaries of females heterozygous for the silencing allele, as well as their daughters that lack the silencing allele.

In two independent experiments, cdi expression was quantified in 20 F1 heterozygous mothers and 20 first-generation backcross daughters, lacking the original silent allele, using RT-qPCR. Confirming previous studies, we found that heterozygous females that maternally inherited the piRNA producing allele had low expression of cdi in ovaries. We further found that the first-generation backcross daughters - lacking the paramutagenic allele – had significantly lower cdi expression in the ovaries compared to a genotypically identical strain with no piRNAs mapping to cdi. This study thus describes a new system of paramutation in gene expression of Drosophila virilis, which can serve as a baseline for future studies that seek to understand how paramutation is regulated.

Paramutation with implications for genome architecture. Studies show that transvection regulates gene expression via distinct DNA elements at specific timepoints in stochastic expression of naturally occurring gene activation and repression between chromosomes. We found a biological role for transvection in controlling the

A telomere associated system of paramutation in Drosophila virilis mediated by maternally provisioned piRNAs

DNA elements act across long genomic distances to regulate gene expression in processes including enhancer-promoter interactions and genetic imprinting. During the gene-regulatory phenomenon of transvection in Drosophila, DNA elements on one allele of a gene act between chromosomes to increase or decrease expression of another allele of the gene. Despite the discovery of transvection over 60 years ago, little is known about its biological role. Furthermore, how separable DNA elements contribute to activating or repressing transvection at distinct times during development is unclear. Here, we study the expression of spineless (ss) in the developing fly eye as a paradigm to understand gene activation and repression between chromosomes. We found a biological role for transvection in controlling the stochastic expression of naturally occurring ss alleles. We characterized CRISPR engineered deletions of sequences across the ss locus and identified DNA elements required for activating and repressing transvection. We found that enhancers participate in transvection at distinct times in development to promote gene expression. Finally, bringing a silencer element on a different chromosome into proximity with the ss locus “reconstitutes” the gene, leading to repression. Our studies show that transvection regulates gene expression via distinct DNA elements at specific timepoints in development with implications for genome architecture.

Activating and repressing stochastic gene expression between chromosomes

Posttranslational modifications of histones impact chromatin structure and function, which underlie everything from genome stability to cell differentiation and disease. Understanding how histone modifications are established and maintained is critical to understanding these processes. BRWD3 (Bromodomain and WD repeat-containing protein 3) is a specificity factor for a CUL4 ubiquitin ligase complex. BRWD3 has been implicated in DNA replication, transcription, and regulation of histone H3 methylation and acetylation. The underlying mechanisms of BRWD3 action, however, are largely unknown. To uncover how BRWD3 functions, we performed both dBRWD3 IP-MS and BRWD3-dependent ubiquitination IP-MS. As expected, we identified the BRWD3-Cul4-DDB1 E3 ubiquitination complex and multiple histones as significantly enriched in our dBRWD3-IP. Interestingly, we identified the Origin Recognition Complex (ORC) as both associated with BRWD3 and ubiquitinated in a BRWD3-dependent manner. Bioinformatic analysis revealed that ~35% of ORC2 binding sites overlap with dBRWD3 binding sites throughout the genome. Furthermore, depletion of dBRWD3 significantly reduces DNA proliferation in the S phase. In addition to ORC, we have also identified several key chromatin-related proteins that could provide mechanistic insight into BRWD3 function. In conclusion, our approach has successfully identified likely direct targets of the BRWD3-Cul4-DDB1 E3 ubiquitin ligase complex, which could help explain how
BRWD3 functions in diverse chromatin-related processes.

**555B Y2H screening reveals potential interactors of a B chromosome-expressed toxin in the jewel wasp Isabella Draper, Sammy Lee, Max Richmond, Patrick Ferree Claremont Colleges, Claremont, CA**

Thousands of plants and animals are known to contain parasitic B chromosomes, many which drive, or segregate at super-Mendelian frequencies. An intriguing question is how B chromosome drive occurs at the mechanistic level. Recently we discovered a toxin gene, named *haploidizer*, which is expressed by the Paternal Sex Ratio (PSR) chromosome in the jewel wasp, *Nasonia vitripennis*. The presence of PSR causes complete elimination of the paternally inherited half of the wasp’s genome, an action that converts female-destined embryos into males, the PSR-transmitting sex. Reduction of *haploidizer* transcripts in the testis by RNAi caused strong suppression of PSR’s genome eliminating activity, demonstrating that this gene is essential in this function. To begin to understand the role of *haploidizer*, we performed yeast 2-hybrid (Y2H) screening with the putatively encoded *HAPLOIDIZER* protein. In the relatively small group of potential interactors, we found several proteins that may perform chromatin-associated functions. These findings are consistent with the hypothesis that PSR disrupts some aspect of paternal chromatin remodeling following sperm entry into the egg. Our group is currently using RNAi to test these candidate *HAPLOIDIZER* interactors.

**556C DNA methylation machinery is required for transcriptome regulation and early development in the wasp Nasonia Jeremy Lynch1, Deanna Arsala1,2 1) University of Illinois at Chicago; 2) University of Chicago**

While DNA methylation is present in most insects, the toolkit has been lost in the lineage leading to Drosophila. Previous experiments in the wasp *Nasonia vitripennis* showed that the methylation toolkit component DNA methyltransferase 1 (Nv-dnmt1a) plays an important role in early embryogenesis, and we are using this model to understand the ancestral role of DNA methylation in insect embryos, and how it was replaced in species where the toolkit has been lost. We found that embryonic lethality of Nv-dnmt1a knockdown is preceded by scattered failures of blastoderm cellularization, and subsequent failures of morphogenetic movements. Such phenotypes are typical of defects in the maternal-zygotic transition (MZT), indicating that DNA methylation may play a role in regulating this process in the wasp. Using whole genome bisulfite sequencing, we show that knockdown of Nv-dnmt1a leads to strong reduction of gene body methylation throughout the genome. Using RNAseq, we show that ~90% of genes downregulated after Nv-dnmt1a RNAi are methylated (in wild-type embryos). This is not unexpected, as it has been previously shown that insect DNA methylation is associated with efficient transcription (and not repression or imprinting as found in vertebrate models). As development proceeds, more and more non-methylated loci are affected, showing a likely indirect effect of DNA methylation on the regulation of transcripts during the MZT. We propose that moderate disruption of the large number of methylated genes has cascading effects on the MZT that lead to failure of proper maternal clearance and/or zygotic activation of genes in early embryogenesis of *Nasonia*.

**557A The histone chaperone NASP has multiple functions during development. Reyhaneh Tirgar, Shannon Leahy, Kendal Broadie, Jared Nordman Vanderbilt University, Nashville, TN**

Histone chaperones aid in the disassembly and reassembly of nucleosomes during the processes of replication and transcription. While histone chaperones have been studied extensively at the molecular level, there remain more open questions about understanding the broader role histone chaperones play during development and disease conditions. Our focus is on the Nuclear Autoantigenic Sperm Protein (NASP) chaperones for histones H1 and H3/H4. These chaperones are required for proper DNA replication, cell cycle progression and cell proliferation. In mammals, NASP is homozygous lethal, but the contributions NASP makes during development are largely unknown. To understand the consequence of NASP loss on development, we used CRISPR/Cas9 to generate a NASP null mutant in *Drosophila*. Zygotic NASP is not essential for viability, however, loss of NASP function results in female sterility. This suggests that maternally deposited NASP has a critical function during embryogenesis. Further, NASP null mutants display severe movement impairments. This striking lack of mobility suggests neurological or neuromuscular defects, which we are currently investigating.

**558B Environmental Effects on the Epigenetic Silencing of Transposable Elements Jennifer McIntyre University of California Irvine**

Rapidly changing dietary and environmental conditions are of concern as to how they influence our genome. Transposable elements (TEs) are widespread DNA sequences that are able to move throughout the genome. They have large potentials to impact our genome, including perturbing the expression of neighboring genes via epigenetic mechanisms. Alzheimer’s, muscular dystrophy, predisposition to cancer, and other diseases are often caused by TEs. Accordingly, it is important to determine how our environment affects this jumping DNA. To mitigate the spreading of TEs, repressive markers are enriched at the TE location. While this epigenetic marker is successful at limiting the movement of TEs, it can also repress neighboring genes. In this experiment, we used a reporter assay that consists of a reporter gene next to a TE. Altered expression level of the reporter gene would inform the extent of TE-mediated...
silencing of neighboring reporter gene. We focus on two major directions: varied climate variables and altered dietary conditions. Our preliminary results suggest that, while changing climate conditions have inconclusive effects on repressive markers, altered diet is likely to have significant influences. We have already seen that a high sugar diet causes increased expression of the reporter gene. This indicates reduced TE-mediated silencing of neighboring genes. We will continue testing other dietary conditions to identify factors that altered TE-mediated silencing. Identification of these prevalent factors could help mitigate TE-mediated abnormal expression of important genes that cause detrimental diseases.

559V The detachment of lamin Dm0 from the nuclear envelope increases variability in 3D positioning of LADs within Drosophila melanogaster nuclei Simon Bondarenko1,2, Igor Sharakhov1,2 1) Department of Entomology and the Fralin Life Sciences Institute, Virginia Polytechnic Institute and State University, Blacksburg, VA 24061, USA; 2) Department of Genetics and Cell Biology, Tomsk State University, Tomsk, Russian Federation

The attachment of chromatin to the nuclear envelope correlates with gene repression, which may play an important regulatory role in the interphase nucleus. Lamins interact with the internal nuclear membrane (INM) and together with other proteins form nuclear lamina. Lamina-associated domains (LADs) of chromatin are shown to interact with the nuclear lamina, however the role of lamins in the 3D position of LADs inside the nucleus is not well understood. In this study, we developed LAD- and non-LAD-specific fluorescent oligo-probes for the chromosome 3 of Drosophila melanogaster and hybridized them with the wild type (wt) Canton S and Lam[A25] mutant nuclei. This mutant lamin lacks the hydrophobic CaaX box responsible for tethering the lamin Dm0 to the INM. The mutant Dm0 protein is not confined to the nuclear perimeter but is distributed throughout the nuclear interior, colocalizing with chromosomes. We performed the confocal microscopy of highly-polytenized nuclei of salivary glands (SG) and low-polytenized nuclei of proventriculus (PV) labeled with LAD- and non-LAD-specific probes using Zeiss LSM 880 with Airyscan module. The radial distribution of DAPI intensity was measured by the radial profile plugin for Fiji software and the radial distribution of LADs and non-LADs was analyzed with the TANGO plugin for Fiji. The radial distribution of chromatin in Lam[A25] nuclei was significantly shifted toward the center of the nucleus in comparison with the wt, which suggests that the lamin Dm0 is necessary for the attachment of chromatin to the NE. Although, we did not detect a significant difference in the radial distribution of LADs and non-LADs between the mutants and wt (PCA analysis), the variability in LADs in the PV nuclei was 29% higher in Lam[A25] than in wt (significant, t-test, p<0.05), whereas the variability in the radial position of LADs in SG was only 10.5% higher in Lam[A25] than in wt (significant, t-test, p<0.05). At the same time, the variability of the radial position of non-LADs was not significantly different among all groups. The results suggest that the Dm0 protein plays an important role in the 3D position of LADs inside the nucleus.

560V ORC associates with the Nup107-160 subcomplex, coupling nucleoporins to replication initiation Logan Richards1, Christopher Lord1, Mary-Lauren Benton2, John Capra1,3, Jared Nordman1 1) Department of Biological Sciences, Vanderbilt University, Nashville, TN; 2) Department of Computer Science, Baylor University, Waco, TX; 3) Bakar Computational Health Sciences Institute and Department of Epidemiology and Biostatistics, UCSF, San Francisco, CA

The regulation of DNA replication is critical to ensure the accurate duplication of genetic information and maintain genome stability. Replication initiates at thousands of replication start sites throughout the genome. The origin recognition complex (ORC) binds throughout the genome to initiate DNA replication. In metazoans, however, it remains largely unknown how ORC is targeted to replication origins to facilitate helicase loading and replication initiation. We hypothesized that ORC’s genomic binding was driven through protein-protein interactions. To address this, we performed immunoprecipitations coupled with mass spectrometry for ORC2, a subunit of ORC, in Drosophila embryonic extract. Surprisingly, we found that ORC2 associates with several subunits of the Nup107-160 subcomplex of the nuclear pore. We determined that this interaction is developmentally regulated, occurring most strongly in the first six hours of embryogenesis. Bioinformatic analysis revealed that, relative to all modENCODE factors, nucleoporins are the most enriched factors at ORC2 binding sites. Critically, ORC2 binding to chromatin is dependent on ELYS, a member of the Nup107-160 complex. Consistent with a function in ORC2 loading, knock down of ELYS delays S phase entry and impairs DNA synthesis. Our work reveals a new connection between ORC, replication initiation, and the nuclear pore. We propose that specific nucleoporins have a previously unrecognized function in metazoan replication initiation.

561V Hinf1 is a guardian of the somatic genome by repressing transposable elements Niraj Nirala, Y Tony Ip University of Massachusetts Chan Medical School, Worcester, MA

Large portions of eukaryotic genomes contain repetitive sequences that include transposable elements, which can be mobilized to generate new integrations in the host genome, therefore affecting genome function, stability and evolution. Germ cells possess the Piwi-interacting RNA (piRNA) pathway to repress transposable elements to maintain genome stability across generations. Whereas, uncontrolled transposable element expression in somatic cells causes mutations that do not get passed on to future generations but may lead to pathological consequences. We have uncovered that loss of function of a single zygotic gene, Hinf1, which encodes a conserved zinc finger transcription factor, is sufficient to cause de-repression of most transposable elements, resulting in a substantial DNA damage in somatic tissues. Deep
sequencing, mutant clonal analyses and cell type specific RNAi experiments reveal that the key cell-autonomous target of Hinfp in this process is the linker histone Histone1, which has important roles in heterochromatin chromatin formation and transposable element repression. Pharmacological inhibition of reverse transcriptase by a combination of 3TC and AZT resulted in rescue of DNA damage in gut enterocytes. Moreover, transgenic expression of Hinfp or Histone1, but not Histone4 of core nucleosome, was sufficient to rescue the defects in repressing transposable elements and host genes. We also demonstrated that down regulation of Hinfp caused neurological defects in aging flies and enhanced the Ras oncprotein-induced lethality and tissue growth. Overall, our findings suggest that Hinfp acts as a pivotal physiological regulator of Histone1-dependent silencing of most transposable elements, as well as many Drosophila genes, and serves as a new venue for studying genome stability, cancer progression, neurodegeneration and aging.

562V  Dual loss of HP1B and HP1C impacts chromatin structure  
Sarah Sims, Nicole Riddle University of Alabama at Birmingham

Heterochromatin Protein 1 (HP1) proteins are non-histone chromosomal proteins that are highly conserved in eukaryotes. HP1 proteins can form homo- and heterodimers, which bind to other chromatin elements such as histones, DNA, and a variety of protein partners. Due to their diverse nuclear functions and maintenance of chromatin states, HP1 proteins are essential for ensuring the safety and functions of the genome. The genome of Drosophila melanogaster contains three somatically expressed HP1 genes: Su(var)205 encoding HP1a, HP1b, and HP1c. Loss of each of the three proteins has important functions, with mutations leading to the misexpression of hundreds of genes and decreased viability and/or fertility. When mutations in HP1 proteins are studied, the impact on the other HP1 family members typically is not assayed, despite them occurring together in protein complexes. Here, we investigate how HP1 proteins interact by examining double-mutant fly strains lacking HP1B and HP1C. We find that the viability of these animals is strongly dependent on genetic background, with one background producing a homozygous viable, fertile, and healthy stock despite lacking two HP1 proteins, while another background allows less than 10% of animals to survive. Examination of polytene chromosomes from these HP1b/HP1c mutants suggests that their morphology is affected, while HP1a continues to localize to the centromeres and telomeres. Our data demonstrate that complete loss of HP1B and HP1C is survivable in D. melanogaster and demonstrate the importance of genetic background. Ongoing studies focus on how pairwise HP1 loss affects gene expression and high-resolution chromatin analysis through CUT&RUN. Our study highlights possible crosstalk and cooperative functions of HP1 proteins and has the potential to provide further insights into the functions of the sole remaining HP1 protein, HP1a.

563V  Repair of double-strand breaks in Drosophila polycomb bodies  
Marieke Wensveen, Aditya Dixit, Aniek Janssen University Medical Center Utrecht

Eukaryotic cells are continuously exposed to DNA damaging insults that can break or chemically modify their DNA. Double-strand breaks (DSBs) are particularly dangerous because their improper repair can directly lead to insertions, deletions, or major structural chromosomal rearrangements. DNA is packaged into a variety of chromatin domains, which each have specific molecular and biophysical properties that can influence the DSB response. Whereas euchromatin contains open, actively transcribed regions, heterochromatic regions mainly consist of compact, silent genomic regions. One major type of heterochromatin is facultative heterochromatin, which is essential to silence developmental genes throughout organismal development. Facultative heterochromatin is enriched for trimethylation of histone 3 lysine 27 (H3K27me3) as well as polycomb proteins, and accumulates in sub-nuclear foci, called polycomb bodies. Although the DNA repair response in euchromatic regions has been extensively studied, the repair response in polycomb-enriched chromatin remains largely unknown. We hypothesize that this heterochromatin subtype requires distinct chromatin- or DNA repair- responses to ensure safe DSB repair. To study the DSB response in polycomb chromatin, we here integrate a previously established inducible single DSB system (A. Janssen et al. G&D 2016) in Drosophila tissue in either euchromatic or polycomb chromatin. Using this system, we find that DSBs in polycomb chromatin employ the canonical repair pathways Homologous Recombination and Non-Homologous End-Joining. Moreover, chromatin analysis reveals local changes in the f-Het chromatin landscape specifically at the break site. We hypothesize that these DSB-induced chromatin changes are necessary to detect, access or process the DNA break. Together, our data indicate that DSBs in polycomb chromatin use canonical DSB repair pathways and require specific local chromatin changes for their faithful repair.

564V  Ectopic heterochromatin triggered by insertion of repetitive DNA is temperature-sensitive  
Melissa Sawyer1,2, Safiyo Aden1,2, Zayid Dakane1,2, Nathan Dupre1,2, Jack Jurmu1,2, Hunter Lindsay1,2, Heidi J.J. Pipkin1,2, Anthony Ruiz1,2, Jessica Xiong1,2, Luke Ziegler1,2, Andrew M. Arsham1 1) Bemidji State University, Bemidji, MN; 2) North Hennepin Community College, Brooklyn Park, MN

Heterochromatin is a key genomic defense against invasive genetic elements, mitigating damage by inhibiting transposition, silencing gene expression, and reducing recombination at insertion sites. How genomes recognize and silence novel threats prior to establishing sequence-specific adaptive defenses like piRNAs is poorly understood. To investigate genome defense against novel repetitive DNA we carried out a transposition mutagenesis screen, mobilizing a
Chromatin insulators are DNA-protein complexes that promote chromatin 3D organization by mediating interactions between distant genomic sites. Insulators can prevent the spread of repressive chromatin and block communication between enhancers and promoters to regulate gene expression. Altering insulator function can lead to defects in cellular differentiation and organismal development. In *Drosophila*, the well-studied *gypsy* insulator consists of three core proteins: Su(Hw), Mod(mdg4)67.2, and CP190. Multimerization of insulator proteins forms insulator bodies, of which normal localization is correlated with proper insulator function. To identify factors required for insulator body formation, we used a cell line expressing a functional Mod(mdg4)67.2-GFP fusion protein and performed a high-throughput visual RNAi screen. This strategy identified Nurf301, a nucleosome remodeling factor, as a potential novel regulator of *gypsy* insulator body formation.

Previous work showed how nucleosomes are shifted at Su(Hw) sites in Nurf301 null hemocytes, but it remained unclear how Nurf301 mechanistically affects *gypsy* insulator function. First, we found that Nurf301 promotes *gypsy*-dependent insulator barrier activity in a ubiquitous manner using an in vivo luciferase assay. Co-IP and IP-mass spec results indicated that Nurf301 physically interacts with CP190, Su(Hw), and Mod(mdg4)67.2. Furthermore, Nurf301 co-localizes with insulator proteins, and depletion of Nurf301 extensively alters the distribution of *gypsy* insulator proteins on chromatin. ChIP-seq and NeuRNA-seq profiles in cells lacking Nurf301 and Su(Hw) revealed that Nurf301 and Su(Hw) mutually affect the binding of one another on chromatin and regulate expression of a similar set of genes. Correlation between altered nucleosome positioning and reduced Su(Hw) binding in Nurf301 null cells suggested a functional relationship. Analysis of chromatin accessibility using ATAC-seq supported the hypothesis that Su(Hw) serves as a docking site for recruitment of CP190, Su(Hw), and Mod(mdg4)67.2-GFP fusion protein and performed a high-throughput visual RNAi screen. This strategy identified Nurf301, a nucleosome remodeling factor, as a potential novel regulator of *gypsy* insulator body formation.

Animal genomes fold into contact domains defined by enhanced internal contact frequencies with debatable functions in establishing independent gene regulatory domains. A large fraction of mammalian contact domain boundaries form by stalling of chromosomal loop-extruding cohesin by CTCF, but most *Drosophila* boundaries form CTCF-independently. However, how CTCF-independent boundaries form and impact organismal development remains largely unexplored. Here, we assess genome folding and transcriptional regulation defects in fly embryos completely lacking the ubiquitous boundary associated factor Cp190. We find that sequence-specific DNA-binding proteins like CTCF and Su(Hw) directly interact with Cp190-containing core complexes and recruit Cp190 to form most promoter-distal boundaries. Cp190 is essential for early development and prevents regulatory crosstalk between gene loci that pattern the embryo. Cp190 is thus currently the major player in fly boundary formation and function, revealing that diverse mechanisms evolved to partition genomes into independent regulatory domains.

Interphase genomes are organized into an intricate three-dimensional (3D) structure that facilitates accurate gene expression and maintains genome stability. This organization is largely conserved, yet how karyotype evolution influences 3D genome organization remains largely unexplored. Here, we use DNA Oligopaints to visualize autosomes and the Z sex chromosome (chZ) in *B. mori* (N=28) and compare our findings to *D. melanogaster* (N=4). We show that *B. mori* nuclei are highly compact, with nuclear volumes similar to *D. melanogaster* despite the *B. mori* genome being over three times larger. This discrepancy in nuclear size cannot be explained by differences in inter-chromosomal interactions or chromosome territory (CT) formation: both species harbor spatially separated CTs with minimal intermixing.
Furthermore, CTs in both species are non-randomly organized, with gene-poor chromosomes being most peripheral. However, homologous chromosome copies share a single CT in D. melanogaster due to somatic homolog pairing, which does not occur in B. mori. Moreover, we find significant differences in intra-chromosomal interactions. Unlike loose chromosome folding observed in D. melanogaster, B. mori chromosomes are tightly folded within CTs, with nearly all tested chromosomes showing interactions between both telomere domains and the center of the chromosome.

Our first-time visual analysis of individual chromosomes further allowed us to address the outstanding question of how dosage compensation (DC) is achieved in B. mori, a ZW species. In B. mori males, both chZs are similar in size and shape and are more compact than autosomes or the female chZ, suggesting that both male chZs are partially and equally downregulated during DC. This mechanism contrasts with DC in D. melanogaster, where the single male chX is upregulated. Thus, B. mori DC is instead more similar to DC in the nematode C. elegans. Using both FISH and ATAC-seq, we find that the female chZ repositions toward the nuclear center and the chromatin becomes more open concomitant with increased Z-linked gene expression at the onset of DC. Together, these studies represent the first non-sequencing-based support for Ohno’s hypothesis for the evolution of dosage compensation. We uncover significant differences in chromosome folding between D. melanogaster and B. mori and intriguing similarities between DC in B. mori and C. elegans, despite these lineages harboring evolutionarily distinct sex chromosomes (ZW/XY). We propose a model where holocentricity may be more influential for interphase genome organization and DC than evolutionary relatedness.

568V  Details of transgene construction determine effective siRNA production

Sudeshna Biswas, Victoria Meller
Wayne State University

Flies correct for imbalance of X chromosome dosage between the sexes by increasing X-linked gene expression in males. The Male Specific Lethal (MSL) complex, composed of proteins and roX RNA, localizes to the male X and participates in gene up-regulation. How the X is identifed remains unclear. Loss of roX RNA causes mislocalization of MSL proteins and male lethality. Small interfering RNAs (siRNA) and satellite repeats enriched on the X contribute to X recognition. Our studies showed that ectopic expression of siRNA from 1.688X (cytological position 3F) partially rescues roX1 roX2 males and localization of the MSL complex, but siRNA from other repeats with similar sequence, including 1.6883A, did not. We hypothesize that details of transgene construction could determine function. To test this, I engineered an siRNA-producing transgene with 1.6881A sequence but with size, phasing and orientation identical to the biologically active 1.6883A transgene. In accord with my hypothesis, the new 1.6881A transgene rescued roX1 roX2 males to the same level as a 1.6883A transgene. This suggests that details of transgene construction play a crucial role in production or processing of double stranded RNA. In the future, I will take advantage of siRNA-producing transgenes that differ in their ability to promote X recognition to determine if epigenetic marks at 1.688 repeats are enhanced only when transgenes that promote X recognition are present. My studies will serve to disentangle the role of siRNA and chromatin modification in dosage compensation.

569V  Aid from repeat-binding and architectural maintenance proteins important in D. melanogaster dosage compensation

Maggie Sneideman, Victoria Meller
Wayne State University

Drosophila melanogaster males carry one X and one Y chromosome, but females have two X chromosomes. To equalize the amount of expression of the X-linked genes between the sexes, males increase the expression of X-linked genes approximately two-fold. This is mediated by the Male-Specific Lethal (MSL) complex, which modifies chromatin to elevate expression. The MSL complex first binds at Chromatin Entry Sites (CES) on the X, and then spreads into nearby active genes. CES contain a short motif that is bound by the adapter protein CLAMP. CLAMP is necessary to attract the MSL complex to the CES. However, these motifs are also found on the autosomes. These autosomal motifs also bind CLAMP, but fail to recruit the MSL complex. CLAMP with X-specific or male-specific effects, proteins with AT hook motifs, and heterochromatin factors. I will use knock down lines of my candidate genes to test for a male specific phenotype. I will mate these lines to a dosage compensation compromised line to determine if it genetically interacts with the CES. To determine if these proteins localize to the 1.688X repeats, ChiP for several candidates suspected to bind in this repeat region will be performed. Lastly, I will use Reem Makki’s dual luciferase reporter assay to identify candidate genes necessary for recruitment of compensation by the 1.688 repeats or CES. Thus far, ISWI, D1, and SAF-A appear to have a genetic interaction with the CES.

570V  Investigating the function of Stonewall in the maintenance of Drosophila female germline stem cells

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The balance between stem cell self-renewal and differentiation is essential to maintain tissue homeostasis. Previous
studies have highlighted that the heterochromatin-associated protein, Stonewall (Stwl), is important for the maintenance of germline stem cells (GSC) in the Drosophila ovary; Stwl loss-of-function results in agametic ovaries while its overexpression results in an increase of undifferentiated cells. However, the mechanism by which Stwl maintains the GSC fate in the female germline remains incompletely understood. We hypothesized that Stwl regulates the balance between self-renewal and differentiation by modulating gene expression in GSCs. To test this, we measured gene expression changes upon overexpression of Stwl in a GSC-enriched population. Our data showed that Stwl overexpression was associated with the downregulation of genes encoding components of extracellular matrix, germline-enriched ribosomal genes as well as germline-associated cytoskeletal genes. Strikingly, we also observe that germline-specific knockdown of Stwl is associated with a specific loss of nuclear lamina in GSCs. Interestingly, tethering genomic loci to the repressive environment of the nuclear lamina has been shown to regulate cell-type-specific gene expression. As such, loss of the nuclear lamina may allow improper gene expression in Stwl-depleted GSCs, leading to their loss. Together, we propose that Stwl tethers differentiation genes to the nuclear lamina in GSCs and orchestrates a gene expression program that promotes self-renewal.

571V  Investigating the consequences of histone overexpression in Drosophila  Risa Takenaka1,2, Harmit Malik1,3  1) Division of Basic Sciences, Fred Hutchinson Cancer Research Center, Seattle, WA; 2) Graduate Program in Molecular and Cellular Biology, University of Washington, Seattle, WA; 3) Howard Hughes Medical Institute, Fred Hutchinson Cancer Research Center, Seattle, WA

Eukaryotic genomes are packaged into chromatin, which is composed of histone-containing nucleosomes. Insufficient histone levels result in poor packaging and promiscuous transcription, whereas excess histone levels impede transcription and other essential cellular processes. Therefore, an optimal histone-to-genome ratio is critical to maintain cellular functions and organismal fitness. In Drosophila, the abnormal oocyte (abo) gene encodes a repressor of core histones. abo is a maternal-effect lethal gene first isolated by Larry Sandler. abo homozygous-mutant females produce fewer offspring relative to their heterozygous sisters. This defect can be attributed to an overproduction of histones, which disrupt the maternal-to-zygotic transition during embryogenesis. Despite abo’s important function in histone regulation, abo-mutant adults show no other morphological phenotypes. Therefore, the Drosophila abo is an intriguing model for studying novel aspects of tissue-specific vulnerabilities to histone overexpression. To this end, I have generated a precise, CRISPR/Cas9-mediated knockout of abo flies to accurately interpret the in-vivo consequences of histone overexpression. Using this abo-knockout fly, I will measure transcriptomic consequences of histone overexpression to potentially explain why different tissues and developmental stages might vary in their responses to histone overexpression.

572A  Spargel/dPGC-1 is required in eggshell patterning and proper cytoskeleton organization during oogenesis and embryogenesis  Mohammed Shah Jalal, Sabarish Nagarajan , Atanu Duttaroy Howard University, Washington, DC

PGC-1 (Peroxisome proliferator-activated receptor-gamma coactivator-1) is a robust activator of mitochondrial biogenesis in the liver and skeletal muscles. On the other hand, Drosophila PGC-1, Spargel (Srl), is predominantly expressed in the Drosophila ovary, and germline-specific knockdown of srl arrests late-stage egg chamber development, resulting in arrested oogenesis and female sterility. Unlike srl RNAi, srl hypomorphic mutants (srl1/srl1) have a marked decrease in growth rate, body size, life span, and fecundity. A CRISPR-mediated srl deletion (srl(del)) is an amorphic allele that is embryonic lethal. But a trans-heterozygous mutant mother (srl(del)/srl1) is fertile and used to investigate further the effects of low dosage of Spargel on oogenesis. We determined the maternal requirement of Spargel during embryogenesis because trans-heterozygous (srl(del)/srl1) females produced deformed embryos, most of which remained unhatched. A range of embryonic deformities was noticed in eggs produced by srl(del)/srl1 mother, which includes ventralized eggshells with phenotypes including single dorsal appendage (DA), 2 short DAs, or 2 DAs close together compared to the control. Broad (Br), a zinc-finger transcription factor that specifies the dorsal appendages forming cells, was not repatterned to clear the dorsal-anterior region in 10 of the 12 stages 10 egg-chambers investigated in srl(del)/srl1 ovaries. Roughly, a third of the post-vitellogenic oocytes are found with Gurken mislocalization. Interestingly, more than 90% of the srl(del)/srl1 egg chambers above stage 9 were found to have actin cytoskeletal defects, which might contribute towards dumping defects leading to small eggs. Follicle cell-specific knockdown of srl has minimum effect on dorsal appendage formation, implying that germ cell Spargel may regulate eggshell patterning via signaling interaction with follicle cells. We also found gaps in the distribution of the syncytial nuclei (nuclear fallout) in a majority of the embryos from srl(del)/srl1 mother, which also show cytoskeletal defects in pseudocleavage furrow formation. Together, our findings suggest that maternal spargel is required for eggshell patterning and cytoskeleton organization during oogenesis and early embryonic development.

573B  Identification and characterization of novel genes in Drosophila’s retinal development utilizing a transcriptomics approach  Sequioa Smith, Mardelle Atkins Sam Houston State University

All cells in a multicellular eukaryotic organism contain a complete genome. However, these organisms possess different cell types with diverse morphologies and functions; thus, highlighting the importance of the regulation of gene expression. Transcriptome data is essential for understanding differential gene expression during development. An
increasing stream of larger and more complex datasets has poured into public databases in the last decade. The storage and maintenance of this shared data are federally funded. Our research seeks to repurpose existing datasets to increase their public value while generating discoveries. Traditionally, genetic screens and isolation of spontaneous mutants have been used to identify factors that affect developmental processes. Mutant isolation and identifications are labor-intensive and time-consuming. We established a rapid data analysis strategy to identify candidate genes required for eye development, beginning with publicly available transcriptomes. Our transcriptome data analysis process produced a candidate list of 47 genes. I will present our strategy for candidate gene identification and initial characterization of several candidate genes.

574C Structure-function analysis of Defective proventriculus (Dve) in Drosophila melanogaster eye development Anuradha Chimatla1, Madhuri Kango-Singh1,2,3,4, Amit Singh1,2,3,4,5 1) Department of Biology, University of Dayton, Dayton, OH; 2) Premedical Program, University of Dayton, Dayton, OH; 3) Center for Tissue Regeneration & Engineering (TREND), University of Dayton, Dayton, OH; 4) Integrative Science and Engineering (ISE), University of Dayton, Dayton, OH; 5) Center for Genomic Advocacy (TCGA), Indiana State University, Terre Haute, IN

Organ development is a highly regulated process to transform a monolayer organ primordium into a fully developed organ. Formation of morphogen gradient acts as critical cue to determine cell fate in developing tissues. In Drosophila melanogaster (Fruit fly), wg morphogen acts as a negative regulator of eye development and wg gradient determines the eye vs. head fate. Previously, we have identified defective proventriculus (dve), an ortholog of SATB1 as a novel dorsal patterning gene that regulates transcription of wingless (wg) morphogen. Axial patterning is required to establish Antero-posterior (AP), Dorso-Ventral (DV), and Proximo-Distal (PD) axes. Of the three axes, DV axis is the first lineage restriction event during eye development and any deviation results in developmental birth defects. In the dorsal gene hierarchy, dve acts downstream of GATA-1 transcription factor pannier (pnr) and upstream of wg. Loss-of-function of dve results in dorsal eye enlargement and increase in wg while gain-of-function results in eye suppression and reduced wg expression. In humans, SATB1 functions as a transcriptional regulator and chromatin organizer and requires tetramerization by the ULD domain. Furthermore, SATB1, the human ortholog of dve is highly upregulated in cancers. Here we have used the Drosophila eye model to understand the role of different Dve protein domains in regulating wg during eye development. We hypothesized that different domains of Dve protein might be critical for regulating downstream target, wg. Understanding this can increase our knowledge of which domains might be pathogenic in human development or disease. We performed structure-function analysis of Dve protein to elucidate the role of various domains in regulating wg and eye development. We have developed several transgenic lines, which will allow us to induce expression of the specific domains of Dve protein and assay their effect in the eye. Dve has a ULD domain for tetramerization, HOX domains for DNA binding and PPP4R2 domain for H2AFX dephosphorylation. Here we present our results on ectopic induction of these domains and their effect on eye phenotype and wg expression in the developing eye. We demonstrate the requirement of tetramerization by ULD domain for Hox domain mediated eye suppression function of Dve and wg regulation during eye development.

575A Hh signaling coordinates stereotyped and stochastic patterns in the Drosophila eye Alison Ordway, Caitlin Anderson, Lukas Voortman, Elizabeth Urban, Robert Johnston Johns Hopkins University

Development of an organism requires both stereotyped and stochastic patterning. Stereotyped patterning robustly generates nearly identical structures across individuals. In contrast, stochastic cell fate specification produces randomized patterns that are unique to each individual. Stereotyped fate decisions are required for the development of many sensory organs, including visual and olfactory systems. Despite their importance, how the molecular mechanisms controlling stereotyped and random patterns intersect within the same tissue has not been addressed. This project aims to determine how gene regulatory mechanisms are tuned to generate both highly regular and stochastic patterns within the same tissue using the Drosophila eye as a model.

The Drosophila eye is composed of ~800 ommatidia in a near perfect array. Each ommatidium comprises eight photoreceptors (R1-8) which develop in a predictable fashion. As photoreceptors are differentiating during larval eye development, a wave of morphogenesis driven by Hedgehog (Hh) signaling promotes the highly reproducible structure of the eye.

Underlying the uniform morphology of the fly eye is a random pattern of photoreceptor subtypes. Two R7 photoreceptor subtypes are defined by expression of light-detecting Rhodopsin proteins. Random patterning of these two R7 subtypes is controlled by stochastic ON/OFF expression of the transcription factor, Spineless (Ss). SsON R7s express Rhodopsin 4 (Rh4), whereas SsOFF R7s express Rhodopsin 3 (Rh3). ss is regulated by an interplay of transcription and chromatin regulation during larval eye development.

I found that Hh signaling plays a second role in eye development to regulate stochastic patterning. hh mutants display a reduction in the percentage of SsON R7s. Cubitus Interruptus (Ci), an effector of Hh signaling, binds at an eye specific
enhancer in ss. This site overlaps with a binding site for Klumpfuss (Klu), a repressor of ss, suggesting competitive binding and regulation.

These results indicate that Hh signaling is finely tuned to drive both stereotyped eye morphology and generate stochastic R7 subtype patterning during development. My ongoing work aims to determine how Hh regulates transcription and chromatin compaction at the ss locus. I am also working to determine how Ci and Klu compete for binding and regulation of stochastic ss expression. My studies will describe how regulation of stochastic and stereotyped patterning is coordinated in a single developing tissue.

576B   Decapentaplegic Regulates the Boundary Expression of Midline and Groucho in the Developing Eye Imaginal Disc of Drosophila Alani Perkin, Parag Bhatt, Britney Roberts, Sandra Leal Harris-Stowe State University

The development of an organism relies on interactions between genes within specific signaling networks across diverse tissues. The regulatory genes of these networks are responsible for programming the proliferation, cell fate specification, and differentiation of unique cell populations that create a tissue pattern central to organogenesis. A well-characterized family of developmental regulatory genes is represented by the T-box (Tbx) gene family that encodes transcription factor (TF) proteins sharing an evolutionarily conserved T-box DNA-binding domain. The Drosophila melanogaster family of developmental regulatory genes is represented by the T-box (Tbx) gene family that encodes transcription factors and co-factors responsible for retinal specification and patterning anterior of the MF. Using the UAS-Gal4 binary expression system, we found that Dpp maintains the proper boundaries of mid and Groucho (Gro) expression within the eye imaginal disc. We are now investigating whether the Notch-Delta signaling pathway plays a role in modulating dpp and mid activity to pattern the eye disc.

577C   Insights into the evolution and development of stochastic Drosophila retinal patterning through cross-species comparison with yellow-fever mosquito, Aedes aegypti Zachary Goldberg, Julia Ainsworth, Yunchong Zhao, Crystal Diei, Michael Perry University of California San Diego

How do adaptations arise within the constraints of existing genetic programs? The vast diversity of insect eyes and downstream neural circuits present an ideal system for studying how evolution modifies developmental programs to create morphological and neural novelty. Because of the interconnectedness and physiological precision necessary for vision, any modification must be accommodated at multiple levels, including visual processing circuits in the brain. Insect eyes are made up of repeating “simple eyes” called ommatidia, and contain a surprising diversity of arrangements and modifications that provide specialized functions. In the Diptera (true flies), each ommatidium contains eight photoreceptors (PRs): six used for motion vision and two used to make color comparisons. Different subtypes of ommatidia have color vision PRs which express different combinations of Rhodopsins sensitive to different wavelengths of light. In Drosophila, stochastic distributions of PR subtypes ensure that different color detectors cover the entire visual field. In contrast, Aedes aegypti mosquito retinas sacrifice stochastic patterning to instead place a subset of color-sensitive Rhodopsins in a local, regionalized ventral stripe of unknown function. What genetic regulatory changes are needed to repattern the eye? How do downstream neural circuits adapt to changes in upstream retinal patterning? What constraint exists on evolution of a system as complex as vision? By leveraging the vast array of tools and knowledge in the stochastic Drosophila retina with the non-stochastic patterning of the Aedes retina, we seek to understand how insects have evolved to pattern their retinas at the level of gene regulation. Previous work has shown that the transcription factor Spineless (Ss) controls stochastic PR patterning in Drosophila. We are investigating the role of Ss in the Aedes retina to understand if gene regulatory changes at the Ss locus alone are sufficient to repattern the eye and rewire the brain. Studying this novel neural feature and how the rearranged receptors in the eye connect to the proper circuits in the brain will greatly enhance our understanding of how complex interconnected systems evolve.

578A   The Goldilocks effect: proper dosage of PAX6 levels is required for proper retinal differentiation and patterning in Drosophila. Claude Jean-Guillaume, Justin Kumar Indiana University

The transcription factor PAX6 is the master regulator of eye development in all seeing animals. In Drosophila melanogaster, its orthologs Eyeless (Ey) and Twin of Eyeless (Toy) sit atop the retinal determination network (Rdn), which represents a group of around fourteen transcription factors and co-factors responsible for retinal specification and patterning. Many of PAX6’s roles were first elucidated in Drosophila through studies of ey, and it’s been believed that the specific loss of ey would result in the collapse of the Rdn and the loss of eye development; while the early embryonic loss of toy results in a headless fly (both Ey and Toy are required for overall growth of the eye-antennal imaginal disc), toy knockdown during larval development does not affect the compound eye. However, eye development can proceed in a complete loss-of-function ey mutant background. Our big question is to understand how the fly’s PAX6
molecules regulate eye development. More specifically, we seek to demonstrate how these molecules cooperate to ensure proper expression of the general morphogen decapentaplegic which is required for proper patterning of the eye. Moreover, we wish to show how a proper balance between the expression levels of eya and toy is required for proper development of the two fly’s visual systems, the ocelli and compound eyes, and dorsal-ventral boundary establishment within the compound eye. Using a set of genetic and molecular tools, we have shown that eye development can proceed in the absence of Ey due to Toy, which binds to and activates Ey targets. We have also shown that Ey plays a previously unknown role early in establishing the anterior dorsal head vertex field, which is the region that later gives rise to the ocelli. Early retraction of Ey from that region is required for proper ocellar and dorsal head vertex development. Finally, we have shown that loss of Ey can lead to a mis-regulation of dorsal-ventral patterning genes in the compound eye.

579B The timing of cell fate decisions is critical for initiating pattern formation in the Drosophila eye Justin Kumar, Bonnie Weasner Indiana University

Patterning of the compound eye begins at the beginning of the third larval instar when a wave of morphogenesis initiates from the posterior margin of the eye-antennal disc and sweeps across the eye primordium. The leading edge of this differentiating wave is visualized by a groove in the epithelium called the morphogenetic furrow. A number of signaling cascades including the Hedgehog, Decapentaplegic, JAK/STAT, EGF Receptor, and Notch pathways are required for the initiation of the furrow. If these pathways are disrupted early in development, then the furrow is prevented from initiating from the posterior margin. As a result, the developing eye field fails to be properly patterned and adult flies lack the compound eyes. These phenotypes are reminiscent of eyeless (eya), eyes absent (eya), and sine oculis (so) loss-of-function mutants. These genes belong to the retinal determination (RD) gene regulatory network and are responsible for specifying the fate of the compound eye. Interestingly, expression of hedgehog and decapentaplegic are lost when the RD network is disrupted. Based on these observations, it has been proposed that the RD network is reiteratively used to first specify the fate of the compound eye and then later to initiate pattern formation. Here we describe an alternate mechanism that explains the role that the RD network plays in pattern formation. We have made the unexpected discovery that the RD network functions within the margin of the disc to control the timing of cell fate decisions. In normal development, cells of the margin which surround the compound eye are transformed into head epidermis during mid to late pupal development. This is well after the eye has been completely patterned. These cells then generate a gradient of Wingless (Wg) signaling that, in turn, establishes distinct cellular fates along the periphery of the retina. The result is a clear and smooth transition from ommatidia to bristle laden head epidermis. We show that in RD mutant discs this transformation occurs prematurely during larval development. This precocious change in cellular fate appears to be the underlying reason for the collapse of hedgehog and decapentaplegic expression and for the failure of the morphogenetic furrow to initiate from the posterior margin. We propose that a cardinal role for the RD network is to control the timing of cell and tissue fate decisions.

580C Elucidating the role of the Drosophila melanogaster TENT5 homolog in eye development Abdulgater Al-nouman, Jennifer Curtiss New Mexico State University

Eyeless is a paired-homeodomain transcription factor that sits at or near the top of a network of transcriptional factors that initiate eye development in D. melanogaster. Eyeless is conserved across metazoans, with the human ortholog being Pax6. Transcriptional networks such as this are required for cell fate determination in all tissues especially during development when a single egg cell differentiates into the astonishing variety of cells seen in an organism. Previous transcriptomic data and chromatin immunoprecipitation sequencing (ChIP-seq) analysis suggest that the D. melanogaster gene CG46385 is a direct transcriptional target of Eyeless. CG46385 is the D. melanogaster ortholog of human TENT5, which has been shown to function as a non-canonical poly-A polymerase (PAP). PAPs are involved in modification of mRNA by adding long strings of adenine bases to the 3’ end, which can affect mRNA stability and translational efficiency. Because CG46385 is a direct transcriptional target of Eyeless, we hypothesize that it has a role in regulating gene expression during eye initiation by regulating mRNA stability. We are performing fluorescent in-situ hybridization on larval eye-antennal imaginal disks and embryos to determine at what stage of development and in what cells CG46385 may function. In addition, we are generating CRISPR-Cas9 mediated CG46385 mutants and plan to observe the eye phenotypes of these mutants to determine CG46385’s role in eye development. Our studies will provide insight into the function of the human ortholog TENT5, which has been linked to congenital diseases including retinitis pigmentosa and osteogenesis imperfecta.

581A Extradenticle expression in the Drosophila Melanogaster eye regulates ectopic patterning on the ventral margin of the eye-antennal imaginal disc Jasmine Warren, Bonnie Weasner, Justin Kumar Indiana University Bloomington

Many kinds of eye diseases, including those that result in blindness originate from genetic mutations of key genes involved in eye morphogenesis. Extradenticle (exd), is a TALE homeobox family transcription factor that is used in a variety of molecular processes such as embryogenesis and development of the central nervous system. However, its role in patterning the Drosophila eye has not been completely understood. The compound eye of Drosophila is
composed of 800-unit eyes, called ommatidia, which are arranged in a stereotyped hexagonal array. This organization, down to the directional angle of the ommatidia, is key to proper vison in adult flies. This specific cell organization is achieved by a wave of cell differentiation called a morphogenic furrow. This furrow arises from a single point on the most posterior end of the eye-antennal imaginal disc and migrates anteriorly in a single wave of differentiation. This patterning event involves several signaling pathways to properly pattern the undifferentiated cells that composed the eye field. My preliminary findings have shown a role for \textit{exd} in retina patterning, which revolves around the regulation of cell differentiation in the ventral side of the \textit{Drosophila} eye. During a genetic screen I found that reductions in levels of Exd leads to the inappropriate release of a second wave of differentiation from the ventral margin of the eye field. This was accomplished by utilizing RNAi and a unique driver, c311-GAL4, that drives expression in a specific tissue layer of the imaginal disc known as the peripodial epithelium. This ectopic patterning results in a completely disorganized retina. A possible cause for the disorganized retina is that \textit{wingless (wg)} expression is lost on the ventral margin when \textit{exd} is knocked down. It has been well documented that \textit{wg} signaling in the eye acts as a repressor to other key patterning genes. I hypothesize that the loss of \textit{exd} leads to the loss of the ventral repression pathway of the eye field. Additionally, I conducted a genetic screen to identify potential binding partners of \textit{exd} that functional in patterning the \textit{Drosophila} eye. The results of my screen showed that the knockdown of \textit{hth} in the peripodial epithelium phenocopies the loss of \textit{exd}. This data shows that \textit{hth} and \textit{exd} may functioning as co-factors to properly spatially restrict the initiation and progression of cell differentiation in the \textit{Drosophila} eye disc.

**582B  Heterodimerization-dependent secretion of BMP5/7 is required for wing patterning in \textit{Drosophila}**

*Milena Bauer1, Gustavo Aguilar1, Kristi Wharton2, Shinya Matsuda3, Markus Affolter1  1) University of Basel, Basel, Switzerland; 2) Brown University, Providence, USA*

One of the key processes of embryonic development is the establishment of differentiation patterns. These, in turn, tightly depend on cell-cell communication, often mediated by secreted signalling molecules. Morphogens are a paradigmatic example: upon secretion from a localized source, morphogens disperse forming concentration gradients that will activate distinct gene targets at different concentration thresholds. Bone Morphogenetic Proteins (BMPs) act as indispensable morphogens in multitude of scenarios. Different BMPs are often expressed in the same tissue acting, potentially, in a combinatorial manner, forming both homo- and heterodimers. While the biochemical properties of the different ligand combinations have been extensively studied, little is known about the impact that they have on morphogen gradient formation.

In the \textit{Drosophila} developing wing there are two BMP-type ligands, Decapentaplegic (Dpp) and Glass bottom boat (Gbb). It has been proposed that they can form both hetero and homodimers. Although is assumed that wing development mainly requires Dpp, Gbb has also been shown to be involved in patterning and growth, despite relatively weak phenotypes. Based on genetic analysis it has been suggested that Dpp and Gbb exhibit different effective signalling ranges. However, dissection of these process exceeds the power of current genetic tools. We have generated \textit{ad hoc} CRISPR reagents and protein binder-based tools to visualize and manipulate the dispersal of Gbb and Dpp.

**583C  Evolutionarily young genes \textit{flf1} and \textit{flf2} are required for Wingless signaling in the wing development of \textit{Drosophila}**

*Yusuke Kurihara, Yuka Doi, Tomoko Nakamura, Yoko Keira, Hiroyuki Ishikawa  Graduate School of Science, Chiba University, Japan*

Conserved and ancient genes often play essential roles in various biological processes, whereas young genes, which exist in only one or a few species, had been considered to perform relatively minor functions. Recent studies in \textit{Drosophila} have demonstrated that young genes play important roles in morphogenesis and behavioral decision. However, little is known about the involvement of young genes in the development of the \textit{Drosophila} wing. Here we show two young genes \textit{four-jointed localization factor (flf)} 1 and \textit{flf} 2 function redundantly to regulate Wingless (Wg) signaling in the \textit{Drosophila} wing development.

In a previous study, we identify \textit{flf1} as a modulator of the cellular localization of the Golgi kinase Four-jointed. Moreover, we identified \textit{flf2}, a paralogous gene of \textit{flf1}, by using nucleotide BLAST against the \textit{Drosophila} reference genome. \textit{flf1} and \textit{flf2} are tandemly arrayed genes, and the amino acid sequences of their translation products are highly similar. For further analysis of functions of \textit{flf1} and \textit{flf2}, we produced mutations of these two genes by using CRISPR/Cas9 system. Single mutant flies that lack \textit{flf1} or \textit{flf2} developed and reproduced without an obvious phenotype. By contrast, the number of chemosensory bristles was decreased in the adult wings of \textit{flf1} \textit{flf2} double mutant. An abnormal number of chemosensory bristles in the adult wings is found in \textit{wg} mutants. Therefore, we decided to investigate relationship between \textit{flf1(s)} and Wg signaling. Wg puncta diffusing away from the dorso-ventral boundary cells was decreased in the wing imaginal discs of \textit{flf1} \textit{flf2} double mutant. This result indicates that \textit{flf1} and \textit{flf2} are required for normal expression pattern of Wg in the wing imaginal discs. To test whether Flf(s) affect Wg diffusion in a dose dependence manner, we overexpressed \textit{flf1} or \textit{flf2} in the wing imaginal discs. Overexpression of Flf1 or Flf2 enhanced Wg diffusion in the wing imaginal discs. Our results suggest requirement of \textit{flf1} and \textit{flf2} for proper expression pattern and function of Wg in the \textit{Drosophila} wing development.

**584A  Defining the Role of \textit{CG11617} in the Transcriptional Control of Muscle Development in \textit{Drosophila}**

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The mammalian Mohawk transcription factor is known to be expressed in embryonic precursors of skeletal muscle and functions by regulating the transcription of slow-twitch myosin-heavy-chain isoform expression in fast-twitch muscle fibers, through Sox6 repression, during muscle development. However, we do not fully understand its mechanistic role in invertebrate skeletal muscle development. By using the Drosophila melanogaster Mohawk ortholog, termed CG11617, we first analyzed CG11617 localization. Thoracic flight muscles reveal nuclear CG11617 localization whereas the tergal depressor of the trochanter muscles (TDT, or jump muscles) reveal nuclear and cytoplasmic localization between the myofibrils. Analysis of CG11617 knockdown animals revealed that these flies are lethal in the pharate adult stage. Upon examination, these mutant flies showed fibrillar disorganization of indirect flight muscles, the absence of jump muscles, and a significant decrease in imaginal wing disc myoblasts compared to the wild-type. Together, these findings may account for the overall skeletal muscle impairment and pupal lethality observed in the knockdowns. Furthermore, utilizing Gal4 driver lines containing fiber-specific enhancer-lacZ constructs, to follow muscle fiber fate, we observed a fibrillar muscle to tubular muscle identity switch in these mutant flies. Overall, these findings suggest that CG11617 may skew skeletal muscle precursor differentiation towards one fiber-type versus another, and CG11617 is a determinant of skeletal muscle fiber-type specification and differentiation.

585B  Single-cell sequencing of Drosophila embryonic heart and muscle cells during differentiation and maturation Georg Vogler1, Bill Hum2, Marco Tamayo1, Yoav Altman2, Rolf Bodmer1 1) Sanford Burnham Prebys Medical Discovery Institute, Development, Aging and Regeneration, La Jolla, CA; 2) Sanford Burnham Prebys Medical Discovery Institute, NCI-designated Cancer Center, La Jolla, CA

The developing Drosophila heart consists of cardial cells that differentiate into different types of cardiomyocytes and pericardial cells. A large body of work has identified numerous genes and pathways involved in heart specification and differentiation, downstream of cardiac transcription factors, such as Tinman (NKX2-5) and Dorsocross1/2/3 (TBX5). The advent of single-cell RNA sequencing (scRNAseq) technology allowed us for the first time to describe the transcriptome of different cardiac cell types in the Drosophila model at high resolution. Here, we applied scRNAseq on sorted cells of late-stage Drosophila embryos expressing a cardiac GFP reporter. We find distinct expression profiles of cardioblasts as they mature to cardiomyocytes, as well as discretely clustering pericardial cells, including a set expressing Tinman that potentially assist in heart morphogenesis. In addition, we describe other cell types that were sequenced as by-catch due to low but distinct extracardiac expression of the GFP reporter. Our studies on wildtype cardioblasts will be the foundation for investigating developmental profiles in mutant backgrounds and for generating gene regulatory networks at single-cell resolution during cardiogenesis.

586C  Discs Large is a novel regulator of the Enteroblast Mesenchymal-to-Epithelial Transition in the adult Drosophila midgut Fionna Zhu, Michael Murray, Georgia Malloy University of Melbourne

Maintenance of epithelial barrier function is essential for human health. The adult Drosophila midgut provides a powerful model system for understanding mechanisms of epithelial homeostasis. It consists of a monolayered epithelium of absorptive enterocytes (ECs) and secretory enteroendocrine cells, underpinned by stem cells and intermediate progenitor enteroblasts (EBs). When EBs differentiate into ECs, they undergo a Mesenchymal-Epithelial Transition (MET) and intercalate into the epithelium.

While much is known about the transcription factors and signalling pathways that regulate EB identity and differentiation, the mechanical aspects of MET are less well studied. To identify novel EB MET regulators we conducted an RNAi screen using the lineage tracing tool, ReDDM. We screened genes likely to be involved in that process and identified three components of septate junctions (SJs): Discs Large (dlg), Mesh and Snakeskin.

Dlg is a member of the MAGUK family of scaffolding proteins and is known to regulate SJ formation, cell polarity and proliferation. EB-specific knockdown of dlg using Su(H)GBE-GAL4, resulted in a multilayered epithelium, a more rounded EB morphology, defective SJs and premature expression of the EC-marker Pdm1. dlg knockdown also led to a shortened lifespan and increased mortality due to chemical damage by DSS, likely as a result of impaired barrier function.

MARCM clones deficient for dlg, recapitulated the multilayering and SJ protein mislocalisation RNAi phenotypes. Interestingly, the extent of multilayering appears more severe in the anterior midgut compared to the posterior. Patches of Mesh and Ssk were mislocalised to the apical membranes of clones and were complementary to enriched F-actin, suggesting disruption of microvilli structure. We also observed intracellular actin-rich vacuoles (likely autophagosomes) and apical membrane disruption, features previously associated with delamination and cell death upon Ecc15 infection, coordinated by the EGFR pathway. Isolated clones were occasionally found detached from the basement membrane, supporting this hypothesis. Gaps in the epithelium were also found nearby clones, with enrichment of SJ proteins at their boundaries.
We are currently examining whether these phenotypes are caused by disruption to key signalling pathways. Together, our results highlight the role of Dlg in midgut homeostasis and MET and the utility of the Drosophila model to expand our understanding of epithelial homeostasis.

587A A role for the apterous gene in adult survival of Drosophila melanogaster Cindy Reinger, Michèle Sickmann, Markus Affolter, Martin Müller University of Basel

The apterous (ap) gene codes for a transcription factor belonging to the LIM-homeodomain family. ap is best known for its role in the patterning of the adult wing where it determines the identity of cells in the dorsal compartment. We have previously shown that the expression of ap in the wing relies on two tissue-specific enhancers, apE and apDV. The phenotypic consequences of an apE deletion are viable and fertile flies devoid of wings, which can be maintained as a homozygous stock. True ap null alleles also produce wingless adults. However, and in contrast to ΔapE, such flies also show the precocious adult death syndrome. This condition had attracted some interest ~40-60 years ago from investigators like King, Butterworth and Wilson. They reported that the syndrome correlates with female sterility, abnormal adipose tissue and adult death within 72 to 96 hrs after eclosion. Among the three, the latter phenotype is the most obvious and striking: ap null flies readily hatch in good numbers and are active. But after about one day, they become lethargic and die within the next 72 hrs. A model for the comprehensive explanation of the syndrome and the connection between the individual disorders is still missing.

In the course of our studies on the ap locus, we have recently started to address the genetic and cellular basis of the precocious adult death syndrome. We will present an overview of our initial results.

588B Patterning and Morphogenesis of the Posterior Midgut Daniel S. Alber1,2, Maria Avdeeva3, Liu Yang2, Shannon E. Keenan1,2, Eric F. Wieschaus1,2, Stanislav Y. Shvartsman3,4 1) Department of Chemical and Biological Engineering, Princeton University, Princeton, NJ, USA; 2) The Lewis-Sigler Institute for Integrative Genomics, Princeton University, Princeton, NJ, USA; 3) Center for Computational Biology, Flatiron Institute, Simons Foundation, New York, NY, USA; 4) Department of Molecular Biology, Princeton University, Princeton, NJ, USA

The organogenesis of the Drosophila posterior midgut (PMG) offers a genetic and morphological model system to study the fundamentals of tissue patterning and morphogenesis in a highly conserved and broadly relevant tissue. ERK-mediated signaling acts primarily through two key regulators, Tailless (Tll) and Huckebein (Hkb), to pattern the posterior region of the embryo prior to gastrulation. We use immunostaining to follow the expression patterns of 16 potential targets of these regulators, including overlapping domains of Brachyenteron (Byn), a highly conserved T-box transcription factor, and Wingless (Wg). We elaborate on the conventional model of specification to include effects of Tll and Hkb mutants on the expression patterns of these 16 targets and establish a hierarchy of expression. We reveal the dynamics of this signaling network via transcriptional live reporters created via CRISPR and live imaging using light-sheet microscopy. Using a novel computational method to fuse imaging datasets of the individual components of posterior patterning into a common multivariable trajectory, we show a low intrinsic dimensionality of posterior patterning. Additionally, we classify the morphological changes as the primordial mid- and hind-gut invaginates and involutes into distinct geometric transformation events and connect it to spatiotemporal expression patterns of key PMG-specifying markers. We focus on the morphological effects of Byn, which plays a key role in the cell shape changes and rearrangements in zebrafish and frog embryogenesis, by studying PMG morphogenesis in various mutant backgrounds. Our analysis supplements the existing understanding of epistatic relationships within the PMG-specifying gene regulatory network and describes PMG morphogenesis in the context of gene expression patterns.

589C Identifying split-GAL4 drivers for targeting and manipulating enteroendocrine cells in the Drosophila midgut Jessica Holsopple1,2, Ellen Popodi1,2, Kevin Cook1,2 1) Indiana University, Bloomington, IN; 2) Bloomington Drosophila Stock Center, Bloomington, IN

The Drosophila midgut is a valuable model tissue in many ways, one of which is its utility for characterization and manipulation of cells involved in intestinal endocrine signaling. As in other organisms, the Drosophila intestine contains a group of secretory cells called enteroendocrine cells. This unique group of cells releases peptide hormones to induce systemic effects that allow the whole organism to respond to changing conditions in the gut. Enteroendocrine cells are often divided into subcategories based on the profile of peptides a cell secretes. The diversity of these cells requires methods of identifying, targeting, and manipulating subpopulations of them to clearly understand the functions of discrete subgroups. The split-GAL4 system provides a promising avenue to address this issue and previous studies have identified many combinations of split-GAL4 drivers that target small groups of enteroendocrine cells. Our current work investigates select pairs of split-GAL4 drivers and characterizes the cells reported by these pairs using the expression profiles of peptide-encoding genes reported by lexA knock-in drivers known to target common, midgut-secreted peptides. Using this technique, we are able to characterize the subset(s) of enteroendocrine cells that a split-GAL4 pair reports. We also test various double reporters containing both UAS and lexAop components and analyze the strengths and weakness of these reporters when compared to each other. Overall, this work provides evaluation of lexA drivers for select peptides, detailed characterization of split-GAL4 driver pairs that allow for manipulation of known subpopulations
of enteroendocrine cells, and a framework for future, straightforward characterization of such pairs not characterized in this study.

590A Characterization of novel Drosophila Egf receptor signaling targets with roles in eggshell structure and morphology Molly Yuschock, Jessica White, Taylor Reiff, Lisa Kadlec Wilkes University

Drosophila epidermal growth factor receptor (Egfr) signaling plays a critical role in many aspects of development including oogenesis, embryogenesis, and proper development of wing and eye tissues. For example, during wing development, Egfr signaling helps specify vein tissues, and in the ovary Egfr signaling is known to establish the body axes during oogenesis. Microarray screens by our lab and others have identified potential downstream transcriptional targets of the Egf receptor using the Drosophila ovary as a model system. Our initial work compared gene expression in fly ovaries where the activity of the Egfr pathway was reduced (gurken mutant), wild-type (OreR), or constitutively active (CY2/ATop). We have employed a number of approaches to further investigate the expression, biological function, and mechanism of action of a subset of putative genes of interest, focusing primarily on genes of previously unknown function. A small-scale functional screen using available collections of UAS-RNAi transgenic flies and P-element insertion lines was used to investigate the possible functions of a group of these novel EGFR-responsive genes. A number of these genes were observed to play roles in normal eggshell structure and morphogenesis. Gene mutant/knockdown phenotypes include decreased chorionic integrity, shortened eggs, and various dorsal appendage malformations, as well as decreased fertility. We have used the CRISPR-Cas9 system to create mutations in some of these “morphogenesis genes.” These mutants have so far recapitulated the previously observed phenotypes, and in at least one case resulted in the observation of an additional phenotype in our null mutant, not seen in the original P-element insertion. We are currently using these CRISPR mutants for further study and characterization of the genes.

591B Characterizing the Role of Doublesex in Creating Sexual Dimorphism in the Somatic Gonad Natalie Murphy, Ellen Baxter, Mark Van Doren Department of Biology, Johns Hopkins University, Baltimore, MD

The Doublesex (Dsx) and Mab-3 Related Transcription factor (DMRT) family proteins are paramount for sex determination in most animals, from planaria to flies, birds, mice, and humans. In Drosophila, the embryonic gonad is formed when a bipotent cluster of somatic gonadal precursors (SGPs) coalesces with the germ cells. Sex determination is regulated by X chromosome dosage, which activates an alternative splicing cascade that yields Dsx in females and Dsx in males. Both Dsx isoforms have the same DNA binding domain, but they regulate their targets differently to yield sexual dimorphism. Dsx, like mammalian Dmrt1, is first expressed in the somatic gonad during embryogenesis and is required for male vs. female gonad development.

It is known that dsx is expressed in early SGPs during development, but the exact timing and role of Dsx in sex-specific cell fate specification during gonad development is unknown. We are taking a multi-pronged approach to investigate the cell fate decisions that are regulated by Dsx, as well as the target genes it controls to make these decisions. First, we used an endogenously tagged GFP-Dsx to characterize Dsx expression in the somatic gonad. We found that throughout development, males express Dsx in the early somatic support cells (i.e., the hub and early cyst cells), as well as the terminal epithelium. Female Dsx expression is dynamic throughout development. By adulthood, Dsx is limited to the early somatic support cells (i.e., the cap cells and escort cells) — this is a similar pattern as seen in the male. However, at pupal and larval stages, Dsx expression is more promiscuous, and is seen in a wider range of somatic cell types.

This points towards an early “establishment” need for Dsx in some cell types, and a more limited need for “maintenance” as the gonad ages. We will use small dsx LOF clones to further investigate this hypothesis, which cells of the gonad require autonomous sex information, and which cells may be regulated through cell-cell signaling. Ultimately, we aim to elucidate the mechanisms of Dsx in regulating the sex determination of Drosophila gonad somatic stem cell development.

592C The role of the extracellular protease AdamTS-B and BMP signaling in wing vein formation Olivia De Grace, Afshan Ismat University of St. Thomas

The Drosophila wing is composed of longitudinal veins (L1-L6) and crossveins, the posterior crossvein (PCV) and anterior crossvein (ACV). AdamTS-B is one of three ADAMTS proteins expressed in Drosophila, specifically in the wing imaginal disc. It was previously shown that loss of AdamTS-B in the wing resulted in extra PCVs or deltas. Moreover, over-expression of this protease resulted in a complete absence of the PCV and distal end of the L5 longitudinal vein. The BMP pathway is known to play an essential role in wing vein formation, especially the PCV. The question we are asking is whether AdamTS-B functions through the BMP signaling pathway to inhibit wing vein formation. In order to explore this question, we are performing genetic interaction studies between AdamTS-B and components of the BMP signaling pathway to examine whether these two proteins function in the same signaling pathway or not. Using the GAL4/UAS system of over-expression, we are using a wing-specific GAL4 line (MS1096-GAL4) to over-express combinations of AdamTS-B and each of the following BMP signaling pathway components: dpp, gbb, and tkv. I predict that over-
expression of these BMP pathway components may counteract the defects seen in AdamTS-B over-expression and result in a more “normal” wing vein pattern. The anticipated impact of this research will be more knowledge of how this important BMP signaling pathway is regulated. We will also learn more about potential functions of the AdamTS-B gene in other tissues as it is also expressed in the embryonic trachea.

593A  A single cell atlas of Drosophila embryonic epidermal and salivary gland cells highlights spatiotemporal gene expression during tube morphogenesis Annabel May, Katja Röper MRC LMB

Many internal organs are composed of epithelial tubes, the morphogenetic effectors responsible for controlling morphogenesis in these tubes and their order of action is yet to be comprehensively assessed. Within our tubulogenesis model, the Drosophila embryonic salivary gland, morphogenetic effectors have previously been identified either individually, as a result of mutant phenotypes and whole embryo microarrays or systematically, though gain of function screens. Currently there has been no attempt to assess the entire transcriptional landscape of the developing Drosophila melanogaster embryonic salivary gland.

Using a combinatorial method of strictly staged embryos, fluorophore driven embryo cell sorting and 10x chromium single cell sequencing we describe two new stage 11 embryonic single cell sequencing data sets. A comprehensive cell lineage of the embryonic salivary gland, central nervous system and muscular cells driven by the transcription factor forkhead covering developmental time points of salivary gland specification, invagination and migration and a second dataset at an identical time point isolating epithelial cells expressing the Drosophila homologue of Beta-Catenin, Armadillo.

Here we present both data sets as a spatiotemporal specific bioinformatic tool to identify and investigating genes involved in salivary gland development. Following bioinformatic labelling of the two cell populations we employed differential expression analysis. This technique alone yielded well-known salivary gland markers and a number of novel genes not previously implicated in salivary gland development. These genes include salivary gland specific and early expressed transcription factors, highly up-regulated genes within the developed salivary gland, genes specifically excluded from the salivary gland compared to surrounding epithelia, and a number of as-yet uncategorised genes. Further investigation into the forkhead driven cell population using pseudotime and marker based identification provided a useful tool for prediction of temporal based expression across salivary gland development. This technique again successfully identified both well-known genes and genes previously not described to be temporally expressed within the salivary gland. This specific dataset has lead to the identification of novel gene expression patterning within the early salivary gland, identification of temporally controlled expression of adhesion molecules in time with morphogenetic movement of the tissue and highlighted upregulation of secretory machinery far earlier than previously reported.

594B  Using NaNuTrap method to provide insight into synchronized remodeling of adjacent tissues ectoderm and mesoderm at gastrulation Zsuzsa Akos, Angelike Stathopoulos California Institute of Technology

Early large scale tissue remodeling often happens concomitantly in neighboring tissues in embryos. In Drosophila melanogaster, the fast phase of germ band extension (GBE) that drives stretching in the ectoderm happens at the same time as cell division in the neighboring mesoderm. The concurrence of these processes suggests that this synchronization relates to (i) a molecular mechanism that induces division at the same time as GBE happens or, alternatively, (ii) cell division in mesoderm is a response to tension created by the ectoderm. The latter mechanism would ensure coordination between these processes and help release the tension when it reaches a level too high to tolerate. We recently developed a new method, NaNuTrap that utilizes maternally deposited fluorescent proteins (FPs) and zygotically expressed nanobodies to label tissues early in development, which previously was challenging in Drosophila due to its fast development compared to the maturation time of the FPs. We use NaNuTrap to label the mesoderm and separate it from the adjacent ectoderm in order to track cell movement and division in these tissues. In this way, we can investigate whether the more detailed spatial pattern of cell division in the mesoderm also follows the stretching pattern of the adjacent ectoderm tissue in wildtype embryos. We also seek to determine whether cell division in the mesoderm would be affected if tension created by the ectoderm is altered. In summary, we use the NaNuTrap method to investigate if these two processes, stretching of ectoderm and division of mesoderm cells, are coordinated through physical attachment or by synchronized molecular pathways that induce them at the same time.

595C  The small GTPase Rap1 promotes polar cell survival and morphogenesis to form the migratory border cell cluster Luke Messer, Jocelyn McDonald Kansas State University

So-called ‘organizer cells’ help other cells develop through directing cell fate specification, stem cell maintenance, and even cell migration. Often these various organizer cell functions are due to production of secreted, diffusible signals that initiate specific responses in neighboring cells. One such organizer is the polar cells that form at the anterior and posterior poles of egg chambers, the functional units of the developing ovary. Polar cells are initially produced in excess. Regulated apoptosis eliminates these ‘supernumerary’ polar cells to produce the ‘mature’ polar cell pairs. Later in
oogenesis, the anterior polar cell pair secretes the cytokine Unpaired to activate JAK/STAT signaling in adjacent follicle cells, which become migratory border cells. If excess polar cells survive, too many border cells are recruited, whereas if one or both mature polar cells die, too few border cells form, either of which inhibits successful migration. Thus, survival and maintenance of the mature polar cells must be tightly regulated so only two polar cells survive, yet the mechanisms regulating this are unclear. Here we show that the small GTPase Rap1 controls border cell cluster size by promoting survival of the mature polar cells. We found that egg chambers deficient for Rap1 form border cell clusters with fewer cells than normal. At least 20% of border cell clusters were missing at least one polar cell in addition to fewer border cells. Loss of Rap1 did not alter polar cell specification or the timely elimination of supernumerary polar cells during early oogenesis. Nor did Rap1 regulate activation of STAT in the follicle cells that will form the border cell cluster. Instead, during mid-oogenesis, Rap1 maintained the viability of anterior follicle cells including the mature polar cells. Without proper Rap1 activity, one or both anterior polar cells underwent apoptosis resulting in smaller border cell clusters. Notably, Rap1-deficient border cell cluster size was rescued by blocking cell death using the apoptosis inhibitor p35. These data together show that Rap1 regulates cell survival through blocking apoptosis, resulting in clusters with the optimal number of border cells. Remarkably, Rap1-deficient polar cell shape was drastically altered, potentially contributing to polar cell death. Thus, Rap1 is a new regulator of organizer cell viability that promotes polar cell survival and morphogenesis to build the migratory border cell collective.

596A Phosphoinositide PI3,4,5P3 turnover modulates cytoskeletal forces controlling Drosophila eye morphogenesis Jacob Malin, Christian Rosa-Birriel, Victor Hatini Tufts University School of Medicine, Boston, MA

Epithelial remodeling relies on modulation of mechanical forces that alter the shape and the relative arrangement of cells. In the Drosophila pupal retina, remodeling is dependent on the repeated pulsed contraction and expansion of cell-cell contacts at the level of adherens junctions (AJs). While actomyosin network assembly controls contraction, branched F-actin network assembly controls expansion. Our goal is to uncover mechanisms that control the pulsatile dynamics of the two networks and coordinate their activities. We previously found that levels of the phosphoinositide PI3,4,5P3 (PIP3) increase during contact expansion and decrease during contraction. This result suggests that PIP3 activates the WAVE regulatory complex (WRAC) to control F-actin branching and protrusive membrane dynamics that drive expansion. To examine the system further, we investigated the role and localization of the phosphatase and tensin homolog (Pten) that hydrolyzes PIP3 to PI(4,5)P2 (PIP2), and separately phosphoinositide 3-kinase (PI3K), the enzyme that phosphorlates PIP2 to PIP3. We find that Pten localizes to AJs and accumulates dynamically during both expansion and contraction. The dynamic localization of PI3K is still being investigated. In pten mutant eyes, the dynamics of PIP3 and of protrusive and contractile effectors are disrupted and cell-cell contacts are shortened or lost altogether. Loss of pten and constitutive activation of PI3K produce similar phenotypes, and PI3K depletion induces related phenotypes. Overall, our findings suggest that the cyclical production and hydrolysis of PIP3 is central to the control and coordination of contractile and protrusive dynamics. PIP3 has been previously implicated in controlling actomyosin contractility and endocytosis in epithelial remodeling. Our findings suggest that PIP3 turnover, rather than overall PIP3 levels, is required to fine-tune contractile and protrusive dynamics that shape this epithelium during development.

597B Customization of tissue growth coordinates organ form and function in the embryo Rajprasad Loganathan1, Daniel Levings2, Ji Hoon Kim1, Michael Wells3, Hannah Chiu1, Yifan Wu1, Matthew Slattery2, Deborah Andrew1 1) Johns Hopkins University; 2) University of Minnesota Medical School; 3) Idaho College of Osteopathic Medicine

Developmental tissue growth in the Drosophila embryo has hitherto been ascribed exclusively to the contributions from early mitotic cycles. Post mitotic non-proliferative growth of embryonic tissues has, therefore, been considered negligible for organogenesis. In this work, we describe a significant contribution of non-proliferative cell growth to the formation of two embryonic tubular organs—the salivary gland (SG) and the trachea. The BTB-domain transcription factor Ribbon (Rib) plays a critical role in promoting growth of both organs. In the SG, loss of Rib results in abnormal cell shape and a significant reduction in cell size, attributable to a loss of cytoplasmic volume gain during embryogenesis. In the trachea, Rib loss also results in a significant reduction in cell volume and a failure of segmental branch connectivity. Tissue-specific ChiP-Seq revealed that, in the SG, Rib binds almost all SG-expressed ribosomal protein genes (RPGs). Follow-up transcription assays (RT-qPCR and FISH) revealed that Rib is required for high levels of expression of all ten of the RPGs we have so far tested. Although Rib can directly bind SG RPG enhancers in vitro, the binding is both weak and not sequence selective. We have demonstrated, however, that Rib may attain specificity for RPG enhancer binding and transcriptional activation through direct cooperative interactions with three previously known activators of RPG expression—Trf2, M1BP, and Dref. Rib, in addition to binding SG RPGs, also binds translation factors and chaperones (other components of the “translatome”) implicated in protein synthesis. The mechanism by which Rib-dependent SG cell growth occurs—boosting expression of RPGs and other components of translation—suggests functional consequences beyond cell growth, i.e., priming this organ for its secretory function by enriching the translational machinery to coordinate organ form with function. Surprisingly, Rib binds non-ribosomal genes to promote cell growth in the embryonic trachea where tissue growth is geared primarily to meet the demands of branch connectivity rather than secretion. The differential deployment of cell growth mediators in these two tissues by a single transcription factor
suggests a model in which the growth of individual tissues is customized to coordinate organ form with function.

598C  Investigating the role of Uif and Gprk2 in tissue-specific growth of the larval trachea Zihao Yu, Robert Ward Case Western Reserve University

Most animal species show allometric growth, which means that different organs and tissues grow at different rates relative to each other. Understanding the mechanisms of how different tissues grow is important, since it can provide insight about unique signals and pathways required for development of different organs. The Drosophila larval trachea is an excellent model for tissue-specific growth, since larvae are transparent and the trachea can be easily visualized. In addition, we and other labs have identified a small number of genes that are required for growth specifically in the larval trachea. Two of these genes are uninflatable (uif) and G-protein coupled receptor kinase 2 (Gprk2). uif encodes a single pass transmembrane protein that is expressed on the apical surface in epithelial cells, and is strongly expressed in the trachea. Loss of Uif results in larval lethality with trachea that are roughly have the relative size of that found in wild type larvae. Gprk2 encodes the only non-neuronal G-protein receptor kinase encoded by Drosophila. Loss of Gprk2 also results in larval lethality, but the mutant larvae have long and highly convoluted trachea. In other developmental contexts, Gprk2 is required to phosphorylate G-protein coupled receptors to mark them for endocytosis and functional attenuation. Preliminary results indicate that Uif levels are higher in Gprk2 mutant larval trachea, raising the possibility that Gprk2 may promote the endocytosis of Uif. We are looking at the genetic and biochemical interactions between uif and Gprk2, and their effect on growth through the insulin signaling pathway.

599A  Regulated actomyosin turnover is essential for eye epithelial morphogenesis Christian Rosa, Jacob Malin, Victor Hatini Tufts University

Epithelial morphogenesis is fundamental to animal development and the understanding of congenital disorders. In the fly retina, the repeated assembly and disassembly of contractile actomyosin and branched F-actin networks drive epithelial morphogenesis. However, the roles and the mechanisms that control the pulsatile dynamics of each network are poorly understood. The Rho1 RhoGTPase regulates the assembly of the contractile actomyosin network by activating both the formin Diaphanous (Dia) to promote the assembly of linear actin filaments and non-muscle Myosin II (MyoII) to bind and contract these filaments. We hypothesized that actomyosin turnover is essential for rebalancing forces in the epithelium for proper remodeling and for preventing tissue damage. To test this idea, we overexpressed a constitutively active myosin light chain kinase (MLCK.ct) and separately a constitutively active Dia (Dia.ca) to respectively activate either MyoII or the assembly of linear F-actin. Both manipulations led to defects in epithelial remodeling. MLCK.ct inhibited pulsatile actomyosin assembly and induced ruptures in the epithelium. Likewise, Dia.ca disrupted epithelial remodeling. To determine the mechanisms that regulate actomyosin turnover, we searched for RhoGEFs and GAPs affecting epithelial remodeling. We identified RhoGEF2 and RhoGAP71E as regulators of actomyosin turnover in this process. RhoGEF2 localized mediologically, while RhoGAP71E localized both mediologically and at the cell surface. Genetic analysis combined with live imaging revealed that RhoGAP71E inhibits pulsatile actomyosin dynamics and decreases MyoII levels, while RhoGEF2 has opposite effects. Our data suggest that RhoGEF2 and RhoGAP71E activate and inhibit Rho1, respectively, to sustain actomyosin turnover, promote proper remodeling and rebalance forces in the tissue to inhibit tissue rupture.

600B  Anisotropic Myosin Recruitment Responds To A Static Source During Drosophila Body Axis Elongation Matthew Lefebvre, Nikolas Claussen, Noah Mitchell, Sebastian Streichan University Of California Santa Barbara

Morphogenesis is an inherently dynamic process which bridges cellular and tissue scales. Across metazoans, the actomyosin cytoskeleton plays a central role in sculpting morphogenetic geometry. Yet the mechanisms by which it is controlled remain poorly understood. Recent evidence suggests two regulatory principles: instructive genetic programs, and a dynamic response to mechanical stimuli. The two are not mutually exclusive and have proven difficult to disentangle. In D. melanogaster body axis elongation, a canonical example of convergent extension, both the required genetic patterning inputs, and the force-generating cytoskeletal actors have been identified. Non-muscle myosin II (MyoII) is recruited to dorso-ventral cell-cell junctions where it generates forces. In anterior-posterior patterning mutants axis elongation is impaired, but the way in which genetic information orients the cytoskeleton remains unclear. We study this process using live (in-toto) imaging and dynamic, quantitative analysis. A systematic comparison of the genetic patterning system as well as the magnitude and orientation of junctional MyoII reveals a series of discrepancies. Crucially, during the course of axis elongation, gene patterns permanently deform with the flowing tissue, while the MyoII orientation remains aligned to the dorso-ventral axis, with only a temporary deflection. Therefore, we propose that MyoII is recruited by a geometrically defined static source rather than cell-intrinsic genetic patterning. Under this hypothesis, we can explain and quantitatively predict MyoII dynamics. Our results suggest that genetic patterning does not directly instruct cytoskeletal regulation during axis elongation. Mechanical cues, in particular embryo-scale tension, are a natural candidate for the static source whose signatures we uncover, since they need not behave in the same way as genetic cues in deforming tissue. Our results add a novel perspective on developmental biology: morphogenesis may be organized by tissue geometry without constant input of...
genetic patterning, allowing for robust, modular and self-organized processes.

601C ArfGAP1 regulates collective cell migration in vivo. Alison Boutet1,2, Carlos Zeledon1,2, Xiaojuan Sun1, Gregory Emery1,2 1) Institut de recherche en immunologie et en cancérologie, IRIC, Montréal, Canada; 2) Université de Montréal, Montréal, Canada; 3) School of Basic Medical Sciences, Henan University, Kaifeng, China

Introduction: Collective cell migration plays important roles in morphogenesis and embryonic development and is a main feature of the formation of metastases in several cancers. Unlike single cell migration, collective cell migration is characterized by cell-cell adhesion and cell-cell communication. We have previously demonstrated that vesicular trafficking plays a critical role in cell guidance and cell-cell communication during collective cell migration. A recent screen aimed at identifying new regulators of vesicular trafficking involved in collective migration identified ArfGAP1 as a regulator of border cell migration in the Drosophila ovary.

Methods and Results: Between stages 9 and 10 of the Drosophila egg chamber development, the so-called border cells form a cluster that is attracted by the oocyte through the secretion of ligands to receptor tyrosine kinases (RTKs). We found that the depletion of ArfGAP1 specifically in border cells induces migration defects. Indeed, clusters devoid of ArfGAP1 are able to initiate their migration, but lose directionality. Investigating the cause of this phenotype by analyzing various determinants of border cell migration, we found that the depletion of ArfGAP1 reduces the level of active RTKs at the plasma membrane as they accumulate in late endosomal compartments. Looking further in the degradative pathway, our results indicate an increase of active RTKs inside late endosomes, combined with an increase in Rab7 signal and lysosomes in border cells depleted of ArfGAP1. Our results suggest that ArfGAP1 is necessary for the proper sorting of active RTKs in endosomes to maintain RTKs at the plasma membrane and allow directionality. Moreover, rescue experiments and genetic interactions revealed that the role of ArfGAP1 in border cell migration is dependant of its GAP activity and could act through Hrs and Llrk, two proteins involved in multivesicular body formation and RTKs degradation.

Conclusion and Relevance: We identified ArfGAP1 as a new regulator of border cell migration that might acting through vesicular trafficking to maintain of active receptor tyrosine kinases at the plasma membrane. This study could potentially reveal a new important mechanism in collective cell migration, and by extent in cancer dissemination.

602A Uncovering the mechanism of hematopoietic niche formation Kara Nelson, Stephen DiNardo University of Pennsylvania Philadelphia, PA

Niches have been well-established as a source for important extrinsic components that regulate the balance between progenitor cell maintenance and differentiation in tissues. Most well-studied niches are positioned reproducibly within tissues, and are comprised of an organized collective of cells. This implies that formation of niche structure is regulated, and suggests that the tissue-specific structure of niches is pertinent to the underlying biology of the tissue that the niche supports. However, mechanisms that determine the initial organization of cells constituting a niche, and the location of the niche within a tissue remain less-understood. To investigate mechanisms of niche formation, we use the posterior signaling center (PSC)—the niche of the larval lymph gland—as a model niche. The PSC niche maintains stem cell-like hematopoietic progenitors that eventually differentiate into mature hemocytes that respond to immune challenges and infliction of wounds, and replenish hemolymph during homeostasis. Initial formation of the PSC occurs during embryogenesis: PSC cells are specified laterally during mid-embryogenesis, and in late embryos, the PSC resides at the dorsal surface of the embryo, coalesced at the posterior of the lymph gland. Preliminary live-imaging of PSC formation revealed that alary muscle (AM) and visceral mesoderm (vm), two tissues nearby the PSC, move towards the dorsal midline in concordance with the PSC. Thus, we hypothesized that one of these tissues, AM or vm, sends guidance cues that direct PSC migration towards the dorsal midline/lymph gland posterior. Using tissue-specific ablation of the AM, we demonstrate that it is not required for PSC formation. However, analysis of late-stage mutants that lack visceral mesoderm shows that fewer PSC cells are present at the lymph gland posterior, suggesting a requirement for vm in the formation of the PSC. Future experiments will investigate if, in the absence of vm, fewer PSC cells were specified, or some PSC cells underwent apoptosis, or PSC cells were misplaced within the embryo during migration – the latter of which would suggest that vm sends a directional or competency cue important for PSC cell migration.

603B Transcriptome analysis reveals temporally regulated genetic networks during border cell collective migration. Emily Burghardt1, Jessica Rakijas1, Antariksh Tyagi2, Pralay Majumder3, Bradley J.S.C. Olson1, Jocelyn A. McDonald1 1) Kansas State University; 2) University of North Dakota; 3) Presidency University, Kolkata, India

Collective cell migration underlies many processes essential to the life of an organism, including sculpting organs during development, wound healing in the adult, and cancer metastasis. Drosophila border cells, which undergo collective cell migration during normal development, are a genetically tractable model in which to study molecular drivers of collective cell migration. In the ovary, a group of 6-10 follicle cells are specified as border cells. At mid-oogenesis, border cells round up as a cluster, detach from the underlying epithelium and begin their migration. The cluster first extends directed
actin-rich protrusions to help them move rapidly through the surrounding tissue. Later, as migration slows down, the cluster rotates several times, then stops at the oocyte border. Successful border cell migration relies on cell signaling, polarization of the cluster, including directed protrusions, remodeling of the actin cytoskeleton, and maintenance of adhesion between border cells and with the nurse cell migratory substrate. Signals from ecdysone, JAK/STAT, EGFR/PVR, and other pathways initiate and direct the migration of border cells. Downstream targets of these signaling pathways, however, are poorly characterized. Nor is it known which genes, if any, are differentially expressed during distinct migration stages. Thus we wanted to identify genes whose expression changed during border cell migration. We performed RNA-sequencing on border cells specifically isolated at pre-, mid-, and late-migration stages. Transcriptome analyses of these cells showed that 1,794 transcripts (1,394 unique genes) were significantly differentially expressed during border cell migration. Downstream analyses, including clustering of genes by expression patterns and testing for gene ontology enrichment and genetic networks, identified nine groups of differentially expressed genes. Many of these genes have known roles in cell junction assembly, epithelial differentiation, cellular morphogenesis, the actin cytoskeleton, and epithelial-to-mesenchymal transitions, but also metabolic processes. To validate this RNA-sequencing approach, we confirmed the expression and/or function of a subset of these genes in border cells, including CG11147, CHES-1-like, and Arf51F. Thus, our transcriptome identified differentially expressed genes in migrating border cells and highlighted multiple genetic networks, as well as individual genes, that may function in border cell migration.

605A Investigating the role of Ecdysone signaling during mid embryogenesis using Halloween genes Jae Ho Lee, Riti Mital, Robert Ward Case Western Reserve University

20-hydroxyecdysone (20E) is a well-characterized steroid hormone required for major development changes in Drosophila. 20E surges before each larval molt, before pupariation, and during terminal differentiation of the adult structures. During molting and metamorphosis, 20E binds to its receptor to directly activate a group of early genes such as Broad-Complex (BR-C), E74 and E75, which activate late genes that are performing cell death of obsolete larval tissues, cell proliferation, etc. There is also a less well characterized pulse of 20E during mid embryogenesis. Previous studies indicate a role for 20E signaling in germband retraction, head involution, dorsal closure, and cuticle secretion. To gain a mechanistic understanding of the function of the 20E of the 20E during embryogenesis, we are characterizing two Halloween mutants, disembodied (dib) and shroud (sro). These genes encode biosynthetic enzymes required for 20E synthesis. Using confocal microscopy of fixed and live wild-type and mutant embryos, we are examining the signaling and cytoskeletal dynamics required for dorsal closure. In addition, we conducted RNA sequencing of wild-type and mutant 9- and 13-hour embryos and found approximately 2000 genes that are differentially expressed between wild-type and mutant embryos. Gene ontology analysis identified genes involved nucleic acid binding and transferase activity to be significantly over represented in this collection. Interestingly, among the top 50 20E-induced genes, 11 encode Inc RNA or Antisense RNA. To extend this study, we are using RNA interference and loss-of-function mutants of 20E induced genes to investigate their role in dorsal closure.

605B Regulation of epithelial tissue sealing during Drosophila dorsal closure by the PI4P phosphatase Sac1 Kimberley Gauthier*, Julie Brill1,2 1) Cell Biology Program, The Hospital For Sick Hospital, Toronto, Canada; 2) Department of Molecular Genetics, University of Toronto, Toronto, Canada

A recurring theme throughout development and epithelial tissue morphogenesis is the sealing or fusion of epithelial tissues, forming new cell junctions and a single seamless layer of cells. This process is central in wound healing, and dysregulation can lead to developmental anomalies such as cleft palates and neural tube defects. Epithelial sealing is essential in dorsal closure during Drosophila embryogenesis in which the epidermis wraps around the developing embryo, covering the underlying epithelium (the amnioserosa) and rejoining at the dorsal midline where the tissue seamlessly zippers shut. Dorsal closure is driven by actomyosin contractility in the amnioserosa, Jun kinase (JNK)
signaling in the leading edge cells of the epidermis, and formation of a supracellular actin cable around the dorsal hole. Previously, embryos with Sac1 mutations were found to die during embryogenesis, leaving behind gaping dorsal anterior holes. Sac1 is a lipid phosphatase that dephosphorylates phosphatidylinositol-4-phosphate (PI4P), a lipid needed for membrane trafficking from the Golgi. Sac1 mutant embryos also have ectopic JNK activation and excessive cellular constriction in the amnioserosa. How Sac1 regulates JNK signaling and actomyosin contractility during dorsal closure remains unknown. Using time-lapse confocal microscopy to image a fluorescent myosin regulatory light chain marker, I found that dorsal closure proceeds at a slower pace in Sac1 mutants, and that some embryos develop tears in the amnioserosa and epidermal puckering towards the final stages of dorsal closure. In ongoing studies, I will further characterize the role of Sac1 in regulating epithelial sealing, actomyosin contractility, and JNK signaling activation. My results will uncover how Sac1 and phospholipid homeostasis orchestrate contractile forces and cellular signaling to promote tissue morphogenesis.

607C  Smog GPCR regulates distinct myosin pools and cortical actin organization during Drosophila SG invagination Vishakha Vishwakarma, Thao Phuong Le, SeYeon Chung Louisiana State University

Epithelial tube formation requires Rho-dependent actomyosin contractility that generates cellular forces to drive cell shape changes and rearrangement. Rho signaling is activated by G protein-coupled receptor (GPCR) signaling at the cell surface in both invertebrates and vertebrates. During Drosophila embryonic salivary gland (SG) invagination, the GPCR ligand Folded gastrulation (Fog) activates Rho signaling to drive apical constriction. Two GPCRs, Smog (ubiquitous) and Mist (mesoderm-specific), regulate myosin contractility downstream of Fog in the early Drosophila embryo. However, the SG receptor for Fog that translates Fog signal to cytoskeletal reorganization has not yet been identified. Using genetic suppression assay and in vitro cell contraction assay, we revealed that Smog transduces Fog signal to regulate Rho kinase (Rok) and myosin accumulation in the apicomedial region of SG cells to control apical constriction during invagination. We also discovered Fog-independent roles of Smog in maintaining epithelial integrity. smog loss results in reduced/mislocalized junctional myosin, which correlates with reduced and discontinuous signals of the apical determinant protein Crb and the key adherens junction protein E-Cad; this leads to a distorted embryonic morphology with an abnormally elongated SG placode. Moreover, smog null mutants also show enhanced bleb formation in the apical domain of SG cells during invagination, suggesting disorganized cortical actin networks upon smog loss. Genetic interaction tests with actin polymerizing/depolymerizing factors reveal that Smog regulates the cortical actin networks by regulating actin assembly/disassembly. Our study supports a model that Smog regulates apicomedial and junctional myosin pools in a Fog-dependent and -independent manner, respectively, and reveals a new role of Smog in regulating cortical actin organization during epithelial tube formation.

608A  Snail drives epithelium-to-mesenchymal transition by cytoplasmic sequestering of polarity protein Bazooka/Par-3 mo weng1, Rolin Sauceda1,2 1) University of Nevada, Las Vegas; 2) Stanford University School of Medicine

Epithelial-to-mesenchymal transition (EMT) converts cells from highly connected and apicobasally polarized epithelial states into detached and often migratory mesenchymal states. It is an essential process during development and its misregulation underlies tumorigenesis. In the conventional model, a key step in EMT is to eliminate cell-cell junctions through transcriptional repression by EMT transcription factors, such as the conserved EMT driver Snail. However, recent advances suggest most EMTs display reversible intermediate states and do not reach the final mesenchymal state where junctions are completely lost. We show previously that in Drosophila gastrula, Snail drives EMT in mesoderm through a post-transcriptional mechanism: disassembling junctions through downregulation of polarity protein Bazooka/Par-3. Here we show that this downregulation of Bazooka/Par-3 is through a cytoplasmic sequestering of Bazooka/Par-3 proteins. Snail is necessary and sufficient in downregulating cortical Bazooka/Par-3. Despite the loss of cortically localized Bazooka/Par-3, we do not observe significant changes in Bazooka/Par-3 protein levels upon Snail overexpression in embryo extracts. To examine the change in cytoplasmic levels of Bazooka/Par-3 in response to Snail expression, we induced Snail overexpression clones in egg chamber follicle epithelium which normally do not express Snail. We find that the Snail-dependent downregulation of cortical Bazooka/Par-3 is accompanied by an increase in cytoplasmic levels of Bazooka/Par-3 proteins. This suggests Baz is regulated at post-translational levels and preferentially localizes in cytoplasm. In addition to the loss of Bazooka/Par-3 and adherens junctions, Snail expression in these cells also leads to cell flattening and even delamination from the follicle epithelium. Overall, these results suggest a post-translational pathway is downstream of Snail in regulating Bazooka/Par-3 localization and it provides a possible molecular mechanism for EMT intermediate states.

609B  Physical aspects of Drosophila gastrulation Konstantin Doubrovsinski1, Joel Tchoufag2, Mohamad Ibrahim Cheikh1, Kranthi Mandadapu2, Amanda Goldner1 1) UT Southwestern; 2) UC Berkeley

Epithelial morphogenesis is a process through which epithelia develop complex shapes. A common mode of epithelial morphogenesis is folding, where an initially flat sheet of cells bends out of plane to create a fold. A popular system to study folding is ventral furrow formation during gastrulation in Drosophila melanogaster. Despite a long-standing effort to understand the mechanism of tissue folding, the mechanism of ventral furrow formation remains unclear.
We argue that the key missing information required to understand tissue folding is the knowledge of tissue material properties. Indeed, as is known from basic physics considerations, the knowledge of tissue material properties together with the knowledge of tissue deformation dynamics determine the forces driving the dynamics uniquely. Hence, determining material properties of embryonic tissues is absolutely required to arrive at a predictive model of morphogenetic dynamics.

Here, we present a novel technique based on micron-sized bendable cantilevers to directly probe material properties of epithelia in a developing embryo. Our data characterizes how embryonic tissues respond to localized nanonewton range forces. In particular, we show that (1) the tissue restores its shape incompletely after an external pulling force is removed, (2) the tissue restores its shape to a lesser extent if the applied force is spread over a larger area, (3) when constant force is applied, the displacement of the point being pulled increases according to a characteristic 1/2 power law. To interpret our data, we developed a novel 3D computational model of epithelia. Based on a very small set of simple assumptions, our model quantitatively explains all the above as well as several other key features of our measurements.

Our modeling approach can be used further to explain key aspects of tissue dynamics during Drosophila gastrulation. Specifically, we characterize gastrulation dynamics in a genetic background (anillin knockdown) where cells remain completely open to the yolk-sack. Despite cells being completely open on their basal site, ventral furrow formation proceeds (almost) completely unperturbed, with ventral furrow forming to the full extent. Our computational model readily accounts for this unexpected observation. In this way, our in vivo measurements together with computational modeling strongly constrain the relative contributions of the various physical effects involved in ventral furrow dynamics.

610C Investigating a morphogenetic role for septate junction proteins in cell shape changes and polarity during dorsal closure Oindrila De, Robert Ward Case Western Reserve University

Septate junctions (SJs) form an occluding barrier in invertebrate epithelia. More than 30 SJ constituents have been identified and characterized for their canonical barrier function but some studies have highlighted a non-occluding role for SJ genes in Drosophila morphogenesis. In our previous studies, we demonstrated a non-barrier requirement for a subset of core SJ genes during head involution, dorsal closure, and salivary gland development in the fly embryo. However, the mechanistic nature of this requirement is unknown. To address this, we are using dorsal closure (DC) as our model to investigate the morphogenetic function of core SJ genes. DC is a mid-embryogenesis process that seals an epidermal gap, which occurs as a result of germ band retraction. It is driven by actomyosin-based contraction of the extraembryonic amnioserosa cells, tension at the leading-edge, and zipper at the canthi mediated by filopodial dynamics and adhesion. Additionally, cells of the lateral epidermis elongate dorsoventrally and exhibit planar polarized expression of various molecular components including actin, tubulin, and Rab GTPases. Loss-of-function mutations in core SJ genes, including coracle, Macroglobulin complement-related, and Neurexin-IV result in a failure to complete DC and is associated with a dorsal open phenotype in cuticle preparation of embryos. Here we are characterizing cellular defects in SJ mutant embryos that may contribute to their failure in DC. We examined fixed tissue of SJ mutants and observed a lack of a smoothly organized leading-edge during advanced stages of DC. Also, lateral epidermal cells in SJ mutants show significant defects in cell shape, including a lower aspect ratio, reduced area, and increased circularity of their apical surfaces. Additionally, SJ mutants fail to maintain robust organization of microtubules and actomyosin in the epidermis, amnioserosa, and leading-edge towards the end of DC. We are using live confocal imaging to quantify defects in actomyosin dynamics at the leading-edge and amnioserosa in SJ mutants. We are analyzing SJ mutants for subtle alteration in apical-basal polarity of the dorsal epidermal cells that may affect cell shape. We are also examining SJ mutants for defects in localization of Rab GTPases, which may cause reduced polarized trafficking of cytoskeletal and adhesion markers, resulting in defects in cell shape changes and ultimately leading to a defective DC.

611A Molecular players of mis-specified cell elimination during development Menna El Gammal, Olga Klipa, Fisun Hamaratoglu Cardiff University

Elimination of aberrantly specified cells is vital for development and the prevention of tumour formation. While our cells carry the same genetic information, only a subset of genes are expressed in a certain cell type, determining its fate. Normal organ development heavily relies on the maintenance of correct gene expression patterns in the different cell types within it. Genetic or epigenetic mutations may cause mis-expression of fate-determining factors in cells, disrupting proper patterning and compromising organ function. It is therefore important for a mechanism to be in place to detect and remove those harmful cells. Importantly, the same strategy may be accountable for recognizing and eliminating early cancer cells with incorrect fates. However, little is known about the responsible molecular players. To elucidate this, we induced aberrantly specified patches of cells by modifying the levels of the transcription factor Apterous in Drosophila. This was done by mis-expressing its negative regulator dLMO. Apterous is expressed in the dorsal compartment of the imaginal wing disc, where it defines dorsal fate. We have since undergone 3 screens. First, we identified genes which are differentially expressed in the context of mis-specified cell elimination using RNA-seq. Then, we tested their involvement in the process with two RNAi verification screens, where we observed any potential changes in adult fly wings and larval wing discs. We reasoned that if a gene is required for aberrant clone elimination, then its downregulation would rescue
those clones, leading to wing defects. In this way, we identified a set of genes that are likely involved in the process of mis-specified cell elimination.

612B Exploring the function of Canoe's intrinsically disordered region in linking cell junctions to the cytoskeleton during morphogenesis Rachel Szymanski, Noah Gurley, Zuhayr Alam, Mark Peifer University of North Carolina at Chapel Hill

The remarkable ability of cells to change shape and move without disrupting tissue integrity is a hallmark of embryonic development. To do so cells must link the contractile actomyosin cytoskeleton to cell-cell and cell-matrix junctions, and this linkage must be robust and dynamic. We have focused on the multidomain scaffolding protein Canoe, homolog of mammalian Afadin, which links the cadherin-catenin complex to actin. It includes two N-terminal RA domains, which bind the small GTPase Rap1, followed by FHA, Dilute and PDZ domains. These are separated from the C-terminal F-actin binding domain by a long intrinsically disordered region (IDR). IDRs are now recognized as important players in the multivalent interactions that assemble multiprotein complexes, including those that form phase-separated biomolecular condensates. One of our tasks is to define Canoe's mechanistic role, by taking it apart as a machine. We began by exploring the roles of its PDZ and F-actin binding domains, thinking they provided the direct linkage between cadherin and actin. However, deleting each domain had only modest effects on protein function. We now have returned to this analysis, exploring a surprising result from earlier work—proteins null alleles of canoe have a milder zygotic phenotype than the “canonical” canoe alleles, whose defects in dorsal closure gave the gene its name. To explore this, we have sequenced a series of 22 EMS-induced canoe alleles. Most result from premature stop codons, arrayed across the coding sequence. Intriguingly, the two strong “canonical” alleles carry stop codons early in the IDR. We suspect they encode truncated proteins that act in a dominant negative fashion, interfering with maternally-contributed Canoe protein. We are now exploring the function of this diverse set of Canoe proteins, examining both zygotic and maternal/zygotic mutants. We think this will help define the role of Canoe’s IDR in protein function.

613C Defining the roles of the small GTPase Rap1 and its regulator Dizzy in embryonic morphogenesis Kristi Yow, Kia Perez-Vale, Noah Gurley, Mark Peifer University of North Carolina at Chapel Hill

During embryonic development, epithelial cells must change shape and move while maintaining a robust linkage between one another. In Drosophila, multiple molecular mechanisms are involved in maintaining this linkage between adherens junctions and the cytoskeleton. Our lab has focused on the role of the junction-cytoskeletal linker Canoe, which regulates diverse events in embryonic morphogenesis, from apical-basal polarity establishment to apical constriction to convergent elongation and collective cell migration. Canoe is activated by the small GTPase Rap1, which in turn is regulated by diverse Guanine nucleotide exchange factors (GEFs). We and others have defined the roles Rap1 plays during cellurization and mesodermal invagination, but little is known about its role in later embryonic morphogenesis. We are thus exploring the roles of Rap1, and those of one of its GEF regulators, Dizzy, contrasting these with roles of the Rap1 effector Canoe. We used maternally driven shRNAi to deplete both maternal and zygotic Dizzy. Our data reveal Dizzy is required to reinforce junctions under tension during germband extension, thus making Dizzy loss comparable to canoe maternal/zygotic mutants. Dizzy knockdown also altered junctional planar polarity, though not to the same extent as Canoe. Later in development the dizzy mutant epidermis remains relatively intact, with the most severe defects localized to the ventral epidermis, as seen previously in canoe mutants. Together these data suggest that Dizzy is the predominant GEF regulating Canoe via Rap1. We’re now extending this analysis to explore the role of Rap1 in morphogenetic movements. Our preliminary data suggest that Rap1 mutants have substantially more severe defects in epithelial integrity. Cell shapes are altered beginning at the onset of germband extension. The balance of apical contractility between different cells appears to be lost, leading to epithelial folding, and by the end of germband extension epithelial junctions become fragmented in many cells. These data suggest Rap1 must have other effectors in addition to Canoe—intriguingly, the Rap1 phenotypes are more similar to those of embryos lacking both Canoe and Pyd (fly ZO-1) function. These data also suggest additional GEFs must regulate Rap1 during these stages. Together, our data provide new insights into the mechanisms regulating cell shape change and ensuring tissue integrity.

614A Protein biogenesis factors Nascent Polypeptide Associated Complex–alpha and Signal Recognition Particle are required in heart development Analyne Schroeder, Georg Vogler, Alexandre Colas, Rolf Bodmer Sanford Burnham Prebys Medical Discovery Institute, La Jolla, CA

Congenital Heart Disease (CHD) is driven by a strong genetic predisposition, yet only a small subset of patients (~20%) are diagnosed with a precise genetic cause. Therefore, expanding the pool of genes associated with CHD and establishing the functional relationships between genes can assemble a more comprehensive genetic network to better understand cardiac development and pathogenesis. In our studies, we identified protein biogenesis cofactors Nascent polypeptide Associated Complex (NAC) and Signal Recognition Particle (SRP) that bind disparate subsets of emerging nascent polypeptides at the ribosome exit site to direct polypeptide fates, as novel regulators of cell differentiation and cardiac morphogenesis. Knockdown (KD) of the α- (Naca) or β- subunit (bicaudal) of NAC in the developing Drosophila heart led to disruption of cardiac remodeling during pupal stages resulting in an adult fly with no heart. Heart loss was
rescued by combined KD of Naca with the hox gene abd-B. This genetic interaction between hox genes and Naca was recapitulated in differentiation assays using human Multipotent Cardiac Progenitors (MCPs). KD of Naca in MCPs led to decreased cardiomyocytes (CM) and increased fibroblasts (Fib) which were reversed upon co-KD with mammalian hox genes HOXC12 and HOXD12. KD of cardiogenic transcription factors (TFs) Gata4/6 and MyoCD led to different population growth profiles, with decreased CMs but no change in Fibs, indicating that Naca may utilize distinct mechanisms in driving cell fates compared to these TFs. The effect of Naca KD on the fly heart was temporally regulated, in that KD in embryo or in pupae caused only a partial loss of the heart, whereas KD during both stages led to heart loss similar to continuous KD throughout life. This suggests that embryonic Naca KD may in part reprogram cells leading to aberrant cardiac remodeling during pupal stages. Lastly, KD of several SRP subunits individually in the fly heart produced a range of cardiac phenotypes that targeted specific segments and cell types, indicating spatially regulated activities of SRP components in the heart. Together, these data suggest that despite NAC and SRP ubiquitous presence, they displayed spatially and temporally fine-tuned activities for proper cardiac morphogenesis. Naca’s interaction with cardiac-specific hox gene functions builds upon the novel role of this pathway and expands our understanding of the complex genetic networks involved in cardiac development and pathogenesis.

615B  The role of Akirin/NuRD interactions during heart development Mia Jones, Hayley Milner, Scott Nowak Kennesaw State University

Congenital heart defects are the most prevalent birth defect presented in the human population. A number of these cases have been ruled as sporadic, or resulting from different interactions of many independent genetic loci and alleles. Gene combinations and chromosomal changes are known to play a crucial role in the development of congenital heart defects, but the precise genetic and environmental factors involved in this process remain poorly understood. In Drosophila, the process of heart specification and patterning is controlled by a number of transcription factors that work together with the nuclear co-factor, Akirin, to mediate cardiac gene expression. Preliminary work in the Nowak Laboratory found that Akirin likely regulates gene expression by working together with the Nucleosome Remodeling and Deacetylase (NuRD) complex. Embryos bearing mutations in different NuRD subunits produce hearts, but they are often severely misshapen, poorly patterned, and have reduced numbers of cardiomyoblasts. Moreover, quantitative live imaging of cardiac contractions in pre-hatching embryos indicates that these hearts display impaired cardiac function. Together, these data suggest that Akirin/NuRD interactions may be key regulators of proper heart function.

616C  Tissue scale viscoelastic properties influence 3-D organ morphology in the developing fly retina Jacob Decker, Xiao Sun, Ilaria Rebay University of Chicago

Development of functional tissue morphologies requires precise spatiotemporal regulation of cellular processes that generate mechanical forces and modulate tissue material properties. Extensive work has revealed the existence of common cell biological strategies that are employed to shape epithelial sheets; however, the influence that tissue scale material/viscoelastic properties impart on morphogenetic processes is not understood. There is growing evidence that within a single cell, different organization states of the cytoskeleton result in different cellular viscoelastic properties, but how these cellular behaviors—in aggregate—influence tissue-scale material properties, is an open question. The Drosophila retina, which has long been used as a model system to understand the cell biological basis of tissue patterning, presents an intriguing opportunity to explore the cell biological and biophysical processes that coordinate to shape the fly eye at the tissue scale.

The fly retina is a neuroepithelium comprised of ~750 multicellular units called ommatidia, each of which comprises a core cluster of photoreceptor neurons surrounded by a supportive lattice of non-neuronal cells. During development, ommatidial units are positioned in a stereotyped curvature that is critical for the optical resolution of the adult compound eye. During pupal development, the apical surfaces of the photoreceptor cells involute and then elongate along the optical axis of the tissue in a process called apical expansion. We have recently discovered that the retina establishes its curvature prior to apical expansion onset, rather than concomitantly as it was previously assumed. This suggests that retinal curvature establishment is a discrete morphogenetic process driven by a distinct mechanism. Interestingly, coincident with curvature formation, the retinal epithelium reorganizes apical cell contacts and cell adhesion machinery, resulting in precise hexagonal packing of ommatidia. Live-imaging experiments of retinal epithelia during this period suggest that the tissue undergoes a transition from a viscous-fluid like state to an elastic-solid like state. This transition depends on both packing geometry of ommatidia in the epithelium and actomyosin contractility. This newly described morphogenetic process in the pupal retina provides a model to probe the relationship between changes in tissue material properties and final tissue morphology. Quantitative live-imaging coupled with perturbations to cytoskeletal networks in the developing retina can provide mechanistic insight into the cellular processes that tune tissue material properties during tissue morphogenesis.

617A  Characterization of mechanosensitive regulation of cell adhesion by membrane kinase Gish Reina Koran, Mo Weng University of Nevada Las Vegas
Mechanical input, much like a biochemical signal, can be converted into a biological response and trigger downstream effects. Adherens junctions, the major cell-cell junctions that resist physical tension, often mediate mechanosensitivity. Our lab has shown that in the mesoderm precursor epithelium of Drosophila gastrula, adherens junctions are strengthened in response to physical tension generated by apical myosin contraction. Such mechanosensitive junction strengthening is essential for mesoderm morphogenesis, but the molecular mechanism mediating this regulation is unknown. We have identified Gilgamesh (Gish) as a potential novel mechanosensitive junction regulator. Using multiple approaches, we show that Gish appears to be recruited to junctions in mesoderm upon Myosin activation but not in tissues without active Myosin. This recruitment is disrupted in alpha-catenin mutants, suggesting its dependency on adherens junctions. Importantly, loss of function of Gish leads to severe defects in adherens junctions and mesoderm internalization. Consistent with weakened adherens junctions, Gish mutants phenocopy junction mutants in membrane morphology. These data suggest Gish is a critical player in mechanosensitive remodeling of adherens junctions.

**618B  Quantitative Models of Mechanical Feedback in Morphogenesis**

Nikolas Claussen¹, Matthew Lefebvre³, Hannah Gustavson¹, Noah Mitchell¹, Stefano De Renzis², Boris Shraiman³; Sebastian Streichan¹ 1) University of California, Santa Barbara; 2) European Molecular Biology Laboratory, Heidelberg

Morphogenesis is an inherently dynamic process which bridges cellular and tissue scales. Quantitative analysis and mathematical modelling applied to dynamic imaging data can play a crucial role in elucidating developmental processes and can turn biological hypotheses into quantitative, testable predictions. The actomyosin cytoskeleton plays a central role in sculpting the developing embryo across metazoans, yet the mechanisms by which it is controlled are poorly understood. Recent evidence suggests that the cytoskeleton can dynamically respond to mechanical stimuli. Yet distinguishing which behaviors are key signatures of such mechanical feedback has proven difficult. In *D. melanogaster* body axis elongation, a canonical example of convergent extension, non-muscle myosin II (MyoII) is recruited anisotropically to cell-cell junctions where it generates force. But how the cytoskeleton is oriented remains incompletely understood.

We show how simple, physical models with a small number of biologically meaningful parameters allow us to describe the behavior of MyoII expected if the cytoskeleton responds to mechanical signals and test our conjectures using an \emph{(in-toto)} characterization of MyoII. We can quantitatively predict the dynamics of both the magnitude and orientation of junctional MyoII based on mechanical cues, cell deformation rates and embryo geometry. We find excellent agreement with experimental data in both wild-type embryos and when using genetic and optogenetic perturbations. Our results show how mathematical models can rationalize morphogenetic behaviors and uncover mechanisms to be targeted by molecular investigations. We add a novel perspective on developmental biology: morphogenesis may be organized by mechanical and geometric signals without constant input of genetic patterning, allowing for robust, modular and self-organized processes.

**619C  Lztr1 is a conserved regulator of Ras/MAPK activity**

Giovanna Collu, Marek Mlodzik Icahn School of Medicine at Mount Sinai

*LZTR1* was recently identified in a screen of human chronic myeloid leukaemia cells aimed at discovering the genetic basis of drug resistance mechanisms. Specifically, LZTR1 was shown to regulate RAS ubiquitination and RAS/MAPK pathway activation in cell culture models. Here we demonstrate a conserved function for the fly orthologue Lztr1/CG3711 in regulating Ras activity *in vivo*. Knockdown of CG3711 leads to Ras gain-of-function phenotypes in the wing, which can be rescued by loss of one copy of *Ras*. Further, through epistasis experiments we show that CG3711 acts in the Egrf/Ras/MAPK cascade to control wing vein patterning and eye development. Consistently with the involvement of LZTR1 in human disorders affecting the nervous system, such as Schwannomatosis, *Drosophila* CG3711 is also expressed in specific neurons and glia. We are currently investigating the function of CG3711 in these tissues, and genetic interactions with other homologues involved in similar human pathologies.

**620A  Robustness of Early Pattern Formation in the Drosophila Visual Map**

Charlotte Wit¹², Melinda Kehribar¹², Robin Hiesinger² 1) equal contribution; 2) Freie Universitât Berlin

The *Drosophila* visual map is a highly patterned synaptic brain region formed by photoreceptor axon terminals in the lamina. We investigated the spatiotemporal organization and the underlying molecular determinants that ensure early pattern formation of this visual map. Unit eyes (ommatidia) and photoreceptor neurons (R1-R6) differentiate in a temporal wave in the developing eye disc; axon outgrowth follows the same wave pattern through the optic stalk that connects the eye disc and the lamina. Photoreceptors bundles are organized in two ways: first, each ommatidium preserves the rotational organization of R1-2-3-4-5-6 neighboring each other (intra-bundle organization); second, R1-R6 bundles preserve their relative positions to each other through the optic stalk and into the lamina (inter-bundle organization). Both types of bundle organization require the cell adhesion molecule Sidekick to preserve the pattern originating in the eye disc. Following the well-characterized differentiation order in the eye disc (R2/5, followed by R3/4, followed by R1/6), the R2 and R5 axon arrive first in the lamina, where they form an ‘equator-blind’ and initially orthogonal pattern. The adhesion G-protein coupled receptor Flamingo/Starry Night (Fmi) is strongly expressed in the
Mechanisms ensuring robustness in organ size regulation are critical for proper development. In epithelial tissues, mechanosensitive Piezo channels are critical for maintaining homeostasis through regulations of cell division and apoptosis. Stretch activation of Piezo triggers cell proliferation. It has been hypothesized that calcium spiking from Piezo activates ERK signaling and induces a G2-to-M transition during cell division. However, the tissue-level mechanistic and functional roles of Piezo during development remain unknown. Here, we have combined pharmacological and genetic approaches to study Piezo's roles in regulating the development of the Drosophila wing imaginal disc, an excellent organism model for epithelial organ development. We investigated the combined outcomes on tissue size, shape and mechanical properties. We found that Piezo overexpression increases the relative concentration levels of cytoskeletal integrin and non-muscle myosin-II leading to increased tissue curvature. Both genetic overexpression and pharmacological activation of Piezo increased proliferation and apoptosis in the tissue. Surprisingly, calcium signaling activity was blocked when Peizo was either overexpressed or knocked down but was increased with acute pharmacological activation. We hypothesize that this loss of calcium activity is due to desensitization of the Piezo channel. Further, knockdown of Piezo increased the numbers of cell neighbors, increased cell area, and decreased cell anisotropy. Piezo overexpression slightly increased the number of cell neighbors and cell area. This highlights the critical role of Piezo in regulating overcrowding within the epithelia during organ growth. In sum, these results support Piezo as a key regulator of epithelial homeostasis through a feedback mechanism that regulates the balance of forces within the epithelium. The outcomes of this feedback loop contribute to the robustness of organ size regulation.

Piezo ensures robust tissue size regulation by balancing proliferation, cell size, anisotropy and cell death Nilay Kumar, Megan Levis, Mayesha Sahir Mim, Maria Unger, Jeremiah J. Zartman University of Notre Dame

Role of Piezo in regulating cell proliferation and apoptosis. Development of an organism is dependent upon proper regulation of gene expression. Initiation of gene expression often relies on long-range signals referred to as morphogens; these morphogens form concentration gradients that aid in specific activation of genes responsible for proper body patterning. In Drosophila, one such morphogen is Dorsal (DI), a transcription factor that helps with patterning of the dorsal-ventral (DV) axis in the early embryo. The impact of DI is further refined by gene regulatory loops that help to control the dynamics of the DI gradient. One regulatory loop of interest is the negative feedback loop with Cactus (Cact). Cact is initially bound to DI, sequestering it to the cytoplasm, but Toll signaling on the ventral side of the cell degrades Cact and allows DI to enter the nucleus. There, DI can activate target genes, one of which is Cact, suggesting that DI may regulate its own inhibition. Our work currently focuses on establishing a system through which Cact can be examined in live embryos. Protein expression and use during development is very rapid; the turnover of Cact happens too quickly for standard live imaging.

Investigating the Role of Septate Junction Proteins during Border Cell Migration Giovanni Sabatino, Haifa Alhadyanian, Robert Ward 1) Case Western Reserve University; 2) University of Kansas

Border cell migration during oogenesis is an excellent system to study aspects of cell migration including polarity, leading edge dynamics, and migratory adhesion. We previously found that knocking down any of several septate junction genes in the border cells resulted in poorly penetrant defects including delayed border cell migration and fragmentation of the border cell cluster. The low penetrance is likely due to the long lifespan of septate junction proteins, coupled with the short time frame during which border cell migration occurs. The septate junction is an analogous structure to the vertebrate tight junction, allowing for occlusion in the epithelium, but septate junction proteins have also been shown to be required for morphogenetic events independent of their role in forming an occluding junction. To gain a better understanding of the function of septate junction proteins in border cell migration we are using confocal microscopy of fixed and live egg chambers expressing RNAi against coracle and Macroglobulin-complement related in heterozygous mutant backgrounds. We are particularly interested in directionality of migration, migratory cell cohesion, and the dynamics of the cytoskeleton at the leading edge. Live microscopy should allow us to observe the actin-myosin leading edge dynamics in real time, as well as better understand if loss of axial direction causes defective border cell migration. Septate junction proteins play diverse roles in embryonic development, and so, by characterizing their effects on border cell migration, we hope to discover new information about the mechanisms of embryogenesis and collective cell migration.

Negative feedback regulation in Drosophila dorsal-ventral patterning Allison Schloop, Gregory Reeves 1) NC State University, Raleigh, NC; 2) Texas A&M University, College Station, TX

Development of an organism is dependent upon proper regulation of gene expression. Initiation of gene expression often relies on long-range signals referred to as morphogens; these morphogens form concentration gradients that aid in specific activation of genes responsible for proper body patterning. In Drosophila, one such morphogen is Dorsal (DI), a transcription factor that helps with patterning of the dorsal-ventral (DV) axis in the early embryo. The impact of DI is further refined by gene regulatory loops that help to control the dynamics of the DI gradient. One regulatory loop of interest is the negative feedback loop with Cactus (Cact). Cact is initially bound to DI, sequestering it to the cytoplasm, but Toll signaling on the ventral side of the cell degrades Cact and allows DI to enter the nucleus. There, DI can activate target genes, one of which is Cact, suggesting that DI may regulate its own inhibition. Our work currently focuses on establishing a system through which Cact can be examined in live embryos. Protein expression and use during development is very rapid; the turnover of Cact happens too quickly for standard live imaging.
techniques, like fluorescent protein fusions. Fluorescent proteins like GFP do not have enough time to mature and fluoresce before the associated protein is degraded. To work around this, we have utilized LlamaTags (Bothma et al. 2018) to image the dynamics of Cactus. We show that Cactus, while predominantly cytoplasmic, is also in the nucleus and shows a recoverable pattern after photobleaching. This provides initial evidence for an identifiable role of nuclear Cactus.

624V  Gene Regulatory Networks in Development: Genetic Variation and Robustness of Anterior-Posterior (AP) Axis Formation in *Drosophila* Lossie (Elle) Rooney1, Prasad Bandodkar2, Cranos Williams3, Gregory Reeves1 1) Graduate Genetics Program, NC State University, Raleigh, NC; 2) Artie McFerrin Department of Chemical Engineering, Texas A&M University, College Station, TX; 3) Department of Electrical & Computer Engineering, NC State University, Raleigh, NC

Body plan patterning is a critical step in embryonic development that has health and viability consequences across the life of the organism. Anterior-posterior (AP) axis formation is an early event in body plan patterning and establishes the head-to-tail orientation for determining cell and tissue fates. In *Drosophila melanogaster*, Bicoid is a well-studied transcription factor that acts as a morphogen in AP axis patterning by influencing expression of the Gap genes in a concentration-dependent manner to create distinct expression profiles. The Gap genes influence additional target genes that also show distinct expression profiles. Though this system has been studied extensively and many of the relevant genes have been identified, the mechanisms that allow robustness of AP axis formation across genetic backgrounds are not well-characterized. We address this gap using the natural variation of the *Drosophila melanogaster* Genetic Reference Panel (DGRP). By quantifying spatial expression patterns of AP genes across lines of the DGRP, we can identify genetic backgrounds that show significant changes in expression. We expect to identify genomic regions (QTLs) associated with these changes in expression, which we will interrogate for potential causal elements such as enhancers of AP genes. We will discuss current imaging results using ~70 lines and QTLs under investigation.

625V  *trithorax* regulates the expression of multiple *Hox* genes within the embryonic dorsal vessel and is required for heart proper and aorta specification Adam Farmer1,2,3, Shaad Ahmad1,2,3, Kristopher Schwab1,2,3 1) Department of Biology, Indiana State University, Terre Haute, IN; 2) The Rich and Robin Porter Cancer Research Center, Indiana State University, Terre Haute, IN; 3) The Center for Genomic Advocacy, Indiana State University, Terre Haute, IN

The *Drosophila melanogaster* embryonic dorsal vessel is a linear myoepithelial tube comprised of a posterior large-diameter heart proper region that intakes and anteriorly propels hemolymph through a narrow tube designated the aorta. The aorta is divided into an anterior and posterior region consisting of contractile cardiac cells (CC) derived from differ ing lineages. The anterior-most aorta contains Tinman (Tin)-positive CCs, while the posterior aorta and heart proper CCs are arranged in repeated ‘hemisegments’ composed of two anterior Seven-up (Svp)-positive cells and four Tin-positive cells. Several Hox genes have critical roles in the anterior-posterior patterning of the dorsal vessel. Furthermore, the *Antennapedia* (*Antp*) and the *Bithorax Complex* (*Bx-C*) genes are expressed in a spatial colinear manner within the dorsal vessel.

In the developing embryo, *Hox* gene expression is positively regulated by *trithorax group* (*TrG*) genes coordinating the proper segmental and morphological identities. Here, we identify the *trithorax* (*trx*) gene as an essential regulator of *Hox* gene expression within the dorsal vessel. Within *trx* null embryos, the heart proper region of the stage 16 dorsal vessel remains structurally identical to the narrow posterior aorta; the posterior region does not develop the luminal dilation, nor the CCs assume the characteristic cellular morphology of heart CCs. Additionally, the posterior region of the *trx* null dorsal vessel adopts a posterior aortic fate as indicated by the loss of the heart luminal marker Multiplexin, an extracellular ligand necessary for heart luminal dilation. The homeotic transformation of the heart proper into the posterior aorta implies dysregulation of *Hox* gene expression with the *trx* null embryo. Indeed, Abdominal-A (*Abd-A*) expression is completely lost within the *trx* null dorsal vessel resembling the *abd-A* null phenotype. Additionally, other *Hox* genes are dysregulated, notably at the anterior and posterior ends of the dorsal vessel. The anterior-most expression of *Antp* was absent from the posterior aorta, while Abdominal-B (*Abd-B*) expression localized to the posterior-most CCs of A8 segment was also lost. Although the higher *Ubx* levels expressed in the posterior aorta were reduced, low levels of Ultrabithorax (*Ubx*) expression persisted within the dorsal vessel. Overall, *trx* is required for *Antp* and Bx-C protein expression within the dorsal vessel which establishes heart and aorta specification. The inactivation of *trx* results with a striking homeotic transformation of the heart proper region into the posterior aorta similar to the *abd-A* null phenotype.

626V  In vivo analysis of a *Hox* gene enhancer required for segment-specific sense organ patterning Xinyuan Liu, Teresa Orenic University of Illinois at Chicago, UIC, Chicago, IL

*Hox* genes encode conserved transcription factors (TFs) that specify segmental or regional identity along the anterior-posterior (A/P) axis of developing animal embryos and also function during later developmental stages in patterning of limbs and other organs. During embryonic development, the *Hox* TFs function at the top of the hierarchy that controls intra-segmental patterning to generate differences in segmental/regional patterning. On the other hand, recent studies in *Drosophila* and other insects suggest that the *Hox* genes are targets of intra-segmental patterning genes
and function downstream of these genes to generate morphological differences among limbs. We are investigating the regulation of the Hox gene *Sex combs reduced* (*Scr*) in response to the intra-segmental patterning networks that control development of the *Drosophila* adult legs. *Scr* is expressed throughout T1 legs, but its expression is elevated in defined domains of developing legs within the primordia of a group sense organs, the transverse bristle rows (TBRs). We have identified an *Scr* enhancer (*ScrE*) that drives expression in the TBR primordia and is required for TBR development in T1 legs. Furthermore, the proximal/distal (P/D) patterning genes *Distalless* (*Dll*) and *bric-a-brac1/2* (*bab1/2*) regulate *Scr* expression through the enhancer. *Dll*, a homeodomain (HD) TF, activates *Scr* expression through multiple sites dispersed throughout the *ScrE* enhancer. In addition, *Scr* is responsive to repression by Bab1/2 TFs and potential Bab-response sequences have been mapped to a 78bp conserved block within *ScrE*. An in vivo functional analysis of these sequences is in progress to determine the necessity of these sites for *Scr* expression and patterning of T1 leg sensory organs. This investigation will provide insight into how Hox gene integration of intra-segmental patterning information leads to the development of segment-specific limb morphologies.

**627V** Physical mechanisms of tissue compartmentalization in the *Drosophila* embryo Gonca Erdemci-Tandogan, Jessica Yu, Negar Balaghi, Veronica Castle, Rodrigo Fernandez-Gonzalez Institute of Biomedical Engineering, University of Toronto, Toronto, ON

Compartment boundaries prevent cell mixing and are essential for embryonic development. Cables formed by actin and the molecular motor myosin II are often found at compartment boundaries. How boundaries are established and maintained remains unclear. In the *Drosophila* embryo, the mesectoderm separates ectoderm and mesoderm, forming the ventral midline. Eventually, mesoderm cells are internalized becoming part of the central nervous system. We found that ectoderm and mesoderm remained separated as the mesoderm was internalized, suggesting the presence of a boundary between the tissues. Using live microscopy, we found an enrichment of myosin at the mesoderm-ectoderm boundary (MEB), forming a supracellular cable. Myosin levels at the MEB decreased as the mesoderm was internalized. To study the role of myosin cables at the MEB, we simulated mesoderm internalization using a vertex model. Our model predicted that tension at the MEB maintains the linearity of the interface, prevents cell mixing, and controls the timing of mesoderm internalization. Consistent with this, pharmacological inhibition of myosin disrupted the MEB, leading to mesoderm-ectoderm cell mixing and premature mesoderm internalization. Our model also predicted that cell divisions in the ectoderm play a role in maintaining the linearity of the MEB, a hypothesis that we are testing.

**628V** Identifying Proteins that Mediate Increased Proliferation at Higher Intracellular pH Laura Martins, Jenna Hunter, Daniel Orozco, Bree Grillo-Hill San Jose State University

Proliferation is a key cellular process that is tightly regulated in cells and essential for the proper growth of multicellular organisms. Many studies have identified genes and proteins that are essential for regulated proliferation, but much less is understood about environmental factors that control proliferation, such as intracellular pH (pHi). pHi is tightly regulated by cells, and emerging evidence suggests regulated pHi dynamics modulate regulated cell proliferation. To regulate pHi, cells use a wide variety of ion exchangers and acid loaders/extruders to maintain pH near physiological levels. *NHE1* in mammals (*DNhe2* in *Drosophila*) is a ubiquitously expressed sodium proton exchanger that acts as a rheostat to maintain physiological pH. In diseases such as cancer, cells have constitutively increased pHi which in turn alters functions of pH-sensitive proteins leading to altered cell behaviors, like increased proliferation. However, it is unknown which specific pH-sensitive proteins are dysregulated at an increased pHi. To study the role of dysregulated pHi in cancer, our lab generated transgenic flies that inducibly express *DNhe2*, the homolog of *NHE1*, in the *Drosophila* eye. In previous work, our lab demonstrated that overexpression of *DNhe2* in developing *Drosophila* tissues is sufficient to increase pHi and increase cell proliferation in vivo, and results in a rough eye phenotype in adult flies. Here we describe a reverse genetic screen to identify candidate pH-sensitive proteins that promote cell proliferation. We screened a collection of 193 *Drosophila* lines covering 94% of the second chromosome. We visually inspected flies for enhancement or suppression of the *GMR>DNhe2* rough eye phenotype. We identified 35 regions of the second chromosome that show an interaction with *GMR>DNhe2*. We are focusing on two overlapping deficiencies that both enhanced the *GMR>DNhe2* rough eye phenotype, *Df (2L)ED1203* and *Df (2L)ED1315* spanning 38B4-38C6. We obtained smaller deficiencies and single-gene mutations within this region to map the interaction to a single gene, and are currently testing for genetic interactions with *GMR>DNhe2*. Next, we will quantify the effects of candidate genes by quantifying proliferating cells in the third larval instar eye and wing imaginal discs. Understanding the effects of increased pHi and which possible pH-sensitive proteins induce hyperproliferation can help us understand and possibly uncover therapeutic targets.

**629V** Characterization of *kayak* (*kay*) mutant phenotypes in *Drosophila* melanogaster eye development Manuel Alejandro Zúñiga-García1,2, Juan Riesgo-Escovar2 1) Maestría en Ciencias (Neurobiología); 2) Instituto de Neurobiología (INB), Universidad Nacional Autónoma de México, Campus Juriquilla.

During development, *kayak* (*kay*) is a pleiotropic gene transcribed in many places and stages. We characterized the eye phenotypes of three *kay* alleles. We generated mutant clones for *kay1*, *kay3*, and *kay4* using the FLP/FRT system,
and analyzed them by optical and scanning electron microscopy (SEM). Mutant kay\(^2\) and kay\(^4\) clones exhibit bristle defects by SEM. kay\(^4\) mutant clones show ommatidium and bristle defects by SEM, and sport big cuticle indentations in the medial-anterior eye region. Semithin optical sections analysis revealed ommatidium polarity and photoreceptor formation defects (ommatidiums with fewer photoreceptors). Taken together, these results indicate that kay plays an important role in eye development.

630V  **Quantitative input-output mapping of cytoskeleton regulator localization demonstrates linearity in developing epithelia systems** Akanksha Sachan\(^1\), Nilay Kumar\(^2\), Alexander Dowling\(^1\), Jeremiah J. Zartman\(^1\) 1) University of Notre Dame; 2) IIT Bombay

The signalling pathways that operate in Drosophila are conserved with remarkable fidelity in all multicellular organisms and help us understand the emergence of birth defects and cancer in humans. For example, epithelial morphogenesis relies on coordination between multiple conserved cytoskeletal effectors; however, quantitative regulatory relationships governing cell mechanics and the tissue scale remain largely unknown. Here, we utilized immunohistochemistry, a semi-automated Python feature extraction workflow, and regression analysis to identify constitutive relationships between cytoskeletal regulators of cell mechanics in wildtype and mutant Drosophila wing imaginal discs. We evaluated possible correlation model functions between cytoskeletal regulators. In particular, we tested for linear regulation of actin accumulation by the strength of integrin concentration levels, which in turn regulates the basal curvature in the developing wing disc. RNAi-based knockdown of integrin with the Gal4/UAS system reduced basal actin levels, confirming integrin regulates actin. Both actin and integrins localize more in areas with increased local basal curvature and reduced cell height. Non-muscle Myosin II concentration also correlates with Integrin and actin concentration levels at the basal surface. Interestingly, basal actin concentration does not depend strongly on the spatial location along the Anterior-Posterior axis. Multiple polynomial relationships were analysed for predicting actin concentration, and a linear model with a single input of Integrin (or Myosin) as an independent predictor variable was selected based on the Akaike information criterion. Prediction error of the model was observed across different locations of the disc, to check how the model performs spatially. The standardized residual error was observed to be higher across the edges of the pouch region. In summary, concentrations of cytoskeletal effectors show a simple linear relationship, a signature feature of robustness in complex systems. Identification of highly collinear dependencies in biological systems also guides in constraining and calibrating biophysical models of epithelial growth and morphogenesis.

631V  **Investigating if the linker phosphorylation sites in Drosophila Smad2 control its stability and transcriptional activity** Edward Eivers, Pablo Flota, Kenny Castro California State University Los Angeles

Signals from the transforming growth factor beta (TGF-\(\beta\)) superfamily of ligands has been shown to be important for cell fate determination, proliferation and differentiation during embryonic development and tissue homeostasis in the adult. This broad family of signaling molecules is typically divided into the bone morphogenetic protein (BMP) and Activin/TGF-\(\beta\) sub-families. Our labs interest, focuses on the activities of a group of TGF-\(\beta\) proteins known as receptor Smads (R-Smads). These transcription factors transmit intracellular signals for each pathway when phosphorylated in their C-terminal domains by ligand-activated transmembrane receptors. In this study, we identified four potential phosphorylation sites in the central linker domain of Drosophila Smad2 (dSmad2). We present data showing that mutation of these linker sites to non-phosphorylatable alanines resulted in stabilization of C-terminally phosphorylated dSmad2 proteins. We also found that when dSmad2 linker mutants were misexpressed in Drosophila wing imaginal discs using Gal4 drivers, adult wings were, reduced in size, bifurcated and displayed severely disrupted venation when compared to control wings. We are also investigating if mutation of the linker phosphorylation sites in dSmad2 can affect expression of downstream TGF-\(\beta\) target genes. In conclusion, our experiments aim to broaden our understanding of the signaling activities of the TGF-\(\beta\) pathway, in particular dSmad2 signal regulation and duration during Drosophila wing development.

632V  **Frizzled receptor-mediated mechanisms of Wingless signaling in developing Drosophila wing epithelium** Swapnil Hingole, Varun Chaudhary Indian Institute of Science Education and Research (IISER) Bhopal, India

Wnts are evolutionarily conserved lipid-modified glycoproteins. Secreted Wnts can travel up to several cell distances over receiving cells and generate a response in a concentration-dependent manner. Their ability to regulate the growth and differentially pattern the tissue makes them crucial for embryonic development and adult tissue homeostasis. Wnts can act directly at a short-range and long-range, which is best studied in the developing *Drosophila* wing imaginal disc. Wingless (Wg, Wnt1 homolog in *Drosophila*) is secreted from narrow stripes of cells along the dorsoventral boundary of the wing disc and activates both short and long-range signaling. However, the mechanisms of long-range gradient signaling are sparsely understood and highly debated. Replacing endogenous Wg with membrane-tethered Wg was able to maintain long-range target gene expression even in the absence of gradient in wing disc and developed a normally patterned wing. Moreover, studies have also demonstrated that long-range target gene expression can be maintained in a ligand-independent manner.

Our findings show that Frizzled 2 (Fz2), a receptor of Wg, maintains target gene expression beyond the reach of Wg
protein in a ligand-independent manner. Fz2 acts redundantly with Fz1 to activate Wg-dependent canonical signaling. However, we found that the maintenance of signaling is a novel non-redundant function of Fz2. This makes Fz2 highly important for the survival and patterning of cells in the absence of the proper Wg gradient. We also show that the Fz2 mediated maintenance of signaling is not dependent on other Wnt ligands. Together these findings and the elevated Fz2 levels in the absence of Wg protein suggest that receptor oligomerization and internalization could be possible mechanisms behind the maintenance of signaling. Thus the combinatorial effect of direct Wg dependent signaling and Fz2 mediated ligand-independent maintenance provides robustness to the developing wing epithelium experiencing varying concentrations of Wg ligand.

633V  Modulation of integrin levels triggers actomyosin reorganization essential for proper tissue folding  
Andrea Valencia Expósito1, Nargess Khalilgharibi2, Yanlan Mao2, María Dolores Matín Bermudo1 1) Centro Andaluz de Biología del Desarrollo, CSIC/Universidad Pablo de Olavide/JA, Sevilla, Spain; 2) MRC Laboratory for Molecular Cell Biology, University College London, London, UK

Folding of epithelial tissues is essential for the formation of complex three-dimensional organ structures during development. However, while folding of epithelial tissues towards the apical surface has been long studied, little is known about the mechanisms underlying epithelial folding towards the basal side. Both apical and basal epithelial tissue folding are accompanied by changes in actomyosin organization and in cell adhesion. Manipulation of apical adhesion molecules, such as cadherins, affects actomyosin dynamics and apical folding. Similarly, adhesion of the cells to the basement membrane (BM), via integrins, is required for proper basal epithelia folding. Nevertheless, how changes in cell adhesion and actomyosin organization interact with each other to drive epithelial folding during morphogenesis remains unknown. Here, we use the primordium of the wing, the Drosophila wing imaginal disc epithelium, as a model system to analyse the role of integrins on the regulation of the actomyosin rearrangements and cell shape changes underlying basal tissue folding. The wing imaginal disc is an epithelial sac contacting on its basal side with a BM. During its development, the disc folds basally along a row of cells, the future wing margin cells. In this work, we show that basal disc folding involves four interconnected events: reduction on integrin levels, increase of F-actin accumulation basolaterally, cell shortening and detachment from the BM. We show that ectopic maintenance of high levels of integrins in wing margin cells prevents these changes, leading to abnormal folding of the epithelium. Contrariwise, reduction of integrin levels in an ectopic location in the disc recapitulates the events that take place at the wing margin, resulting in the formation of an ectopic fold. Finally, through computational exploration, we have found that the reduction of adhesion to the BM must precede the changes in F-actin reorganization and cell shortening to allow proper folding of the epithelium. Based on these results, we propose that changes in the adhesion of cells to the BM mediated by integrins trigger modifications in the actomyosin network necessary for basal epithelial folding during organogenesis.

634V  Apterous Regulates the Formation of Stable Myotendinous Junctions in the Drosophila Embryo  
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Skeletal muscles are required for locomotion such as walking, grasping, blinking and chewing. To accomplish these movements, muscles must form secure tendon attachments that can withstand the force of contraction. It is vital that strong attachments form between the right pairs muscle and tendons during development. These myotendinous junctions link transmembrane Integrin proteins to the cytoskeleton and sarcomere. In the Drosophila embryo thirty distinct muscles make up each abdominal hemisegment. While certain muscle subsets are known to have specific guidance cues (for example, ventrolateral muscles use the protein Kon-tiki), a gap in our knowledge exists about how the lateral transverse muscles regulate where and how their attachments are formed. We have identified the transcription factor Apterous as important for the formation of robust muscle attachments. Apterous is expressed in and required for the proper patterning of all four lateral transverse muscles in each abdominal hemisegment. We find that embryos with gain or loss of Apterous function display attachment defects, including missing or incorrect attachments. Misexpression of Apterous leads to severe disruptions to the muscle pattern, including an increase in lateral transverse-like muscle morphologies. Moreover, using time-lapse confocal imaging, we show that misexpression of apterous in the somatic musculature leads to loss of muscle attachment upon the onset of contractions, resulting in an inability to hatch and embryonic death. We demonstrate that overexpression of apterous leads to loss of both beta-PS Integrin and alpha-PS2 Integrin from myotendinous junctions. Our work establishes a clear function for apterous in the regulation of muscle attachment, linking changes in gene expression to alterations in muscle morphology.

635V  The Thanos Requirement for Transdetermination Leads to an End Game on Wing Cell Fate as Ectopic Eyes  
Develop Alison Smith, Justin Kumar Indiana University Bloomington

As development begins cells are in a pluripotent state, but via cell signaling pathways and morphogen concentration...
gradients cells begin to determine towards their final cell fate. Various developmental anomalies arise when a cell does not adopt or retain its correct fate. A process where a determined cell switches its fate to that of another without dedifferentiation is via transdetermination. Transdetermination can be studied in *Drosophila* via ectopic eye formation. Ectopic eyes can be formed through the misexpression of the master regulator gene of the retinal determination network *eyeless* (*ey*). Though it is more complex than a wing imaginal disc’s cell immediately switching its fate to that of a retinal fate solely with the initiation of *ey* misexpression. There are also certain requirements that must be met - the Thanos requirement. The Thanos requirement has five pillars including: spatial misexpression, co-expression of Dpp, magnitude of misexpression, initiation and duration of *ey* misexpression. I hypothesize that during this transdetermination as the cells move from their wing fate to that of retinal fate the cells proceed through a biphasic state where the cell is not solely fated as either wing or retinal. This is hypothesized due to my work using the UAS-GAL4 misexpression system. With this system I uncovered a previously undiscovered pillar of the Thanos requirement: duration of *ey* misexpression. This uninvestigated requirement is as follows, for an ectopic eye to form there must be GAL4 expression to drive *ey* misexpression not only in the wing disc but also the in the eye field of the eye antennal disc. This is such that as the cells begin to take on a more retinal fate than wing if the GAL4 driver is active in only the wing imaginal disc it will turn off, ceasing *ey* misexpression and no ectopic eye will form. Though if the GAL4 driver is active in both the wing imaginal disc and the eye antennal disc even as the cell takes on a more retinal fate the GAL4 enhancer will remain on, *ey* misexpression will continue, and an ectopic eye will form (if the four other pillars are also met). In the cases where all of the five pillars of the Thanos requirement are met there is an end game on wing fate as the cells adopt their new retinal state. I am further investigating the Thanos requirement using immunofluorescence and quantitative PCR. By understanding the requirement of transdetermination we may better understand the mechanism of transdetermination. By better understanding the mechanism of transdetermination we can not only gain a better understanding of cell fate determination, but what goes awry when cells switch their fates during development.

**636V**  The JNK and Hippo pathways regulate an overlapping transcriptome to control neoplastic tissue growth *Katrina Mitchell*, *Joseph Visser*, *Kieran Harvey* 1) Peter MacCallum Cancer Centre, Melbourne, Australia; 2) Sir Peter MacCallum Department of Oncology, University of Melbourne, Australia; 3) Depart of Anatomy and Developmental Biology, Monash University, Australia; 4) Department of Clinical Pathology, University of Melbourne, Australia

The final size of an organ is determined by both organ-extrinsic and organ-intrinsic factors, which can adapt to changes in environmental conditions and tissue damage. Homeostatic mechanisms eliminate damaged or abnormal cells to prevent tumour growth, which in turn facilitates compensatory proliferation to maintain organ size. In growing *Drosophila melanogaster* epithelial tissues, cells with disrupted apical basal polarity are eliminated by neighbouring wild type cells through cell competition. The JNK pathway limits the growth and viability of neoplastic cells and has been reported to do this by suppressing the key Hippo pathway transcriptional regulatory protein Yorkie. Using targeted DamID and transcriptomics, we have identified target genes of the JNK and Hippo pathway transcription factors and found that these pathways act in parallel to regulate overlapping target genes. Yorkie activates the transcription of genes that promote cell proliferation and survival, whilst the JNK pathway transcription factors Jun and Fos repress transcription of these genes specifically in neoplastic cells. Additionally, we have identified a co-repressor that Jun/Fos cooperates with to prevent the uncontrolled growth of neoplastic tumours. These findings shed new light on organ size control, tissue damage and tumour suppression.

**637V**  The Osiris family genes regulate endocytic trafficking during *Drosophila* tracheal maturation *Lan Jiang*, *Aaron Scholl*, *Istri Ndoja*, *Doria Morante*, *Abigail Ivan* Oakland University

The *Drosophila* trachea is a ramifying network of epithelial tubes with a monolayer of epithelial cells surrounding an apical lumen. Following the formation of continuous tubes, the tube maturation process follows. It is a multistep process: (a) apical secretion to form an apical lumen containing chitin-based extracellular matrix, during which tubes expand in both diameter and length, (b) solute clearance to remove solid luminal matrix and liquid in the lumen, and (c) air filling to inflate the entire trachea. These changes occur at the apical side of the tracheal tubes with little or no changes to the basolateral side.

The *Osiris* (*Osi*) gene family is located at a locus that exhibits dosage sensitivity (triplo-lethal and haplo-lethal). The protein sequence annotation of *Osi* proteins identified an endo/lysosomal signal, a transmembrane domain, a domain of unidentified function DUF1676, and an AQXLAY motif, suggesting that these proteins are likely involved in vesicular trafficking. However, the functional analyses of *Osi* genes are very limited. Only a few studies indicate the potential role of *Osi* genes in endosome-mediated protein trafficking. For example, RNAi knockdown of *Osi 21* suggests its involvement in lysosome-mediated degradation of endocytosed rhodopsin in *Drosophila* eye. In addition, *Osi23* is essential for the formation of nanopores lining the olfactory sensillum in *Drosophila*, potentially through endo/lysosome-mediated trafficking. However, the role of *Osi* genes in *Drosophila* trachea is still unknown.

Previously we identified several *Osi* genes with obvious expression in the *Drosophila* trachea from stage 14 until the end of embryogenesis. In addition, expressing HA-tagged *Osi* proteins in trachea revealed their localization in vesicle-like
structures, suggesting their potential roles in vesicular trafficking. Immunohistochemistry using Osi specific antibodies further confirmed that the endogenous Osi proteins are localized at various endosomes in tracheal cells. The potential functional redundancy of Osi genes prompted us to generate Osi single, double, and triple loss-of-function mutants using CRISPR-cas9. No obvious phenotypes were observed in single mutants. However double and triple mutants showed strong defects in trachea tube maturation. Genetic interaction studies suggest that Osi genes are required for trachea tube maturation through endosome-mediated protein trafficking.

638V  The adult Drosophila salivary gland exhibits an unusual mode of cell division Caitlin van Ree, Nicole Dominado, Nicole Siddall, Gary Hime Department of Anatomy and Physiology, University of Melbourne, Parkville, VIC, Australia

Larval salivary glands of Drosophila are well known to exhibit polytene salivary glands formed via endoreplication and have been long used to study this process. In contrast, less is known of the development of adult Drosophila salivary glands except that they consist of a single layer, tubular epithelium that originates from a population of diploid cells found as an imaginal ring near larval salivary gland ducts. We have shown that the adult salivary glands contain three distinct epithelial domains, two of which are comprised of cuboidal epithelial cells and one of squamous epithelial cells. These cell types develop during the pupal period and after eclosion secretory cells develop extensive apical membrane invaginations. The junctional polarity of the epithelial cells exhibits an unusual change soon after eclosion as E-cadherin localisation migrates from a position apical to the septate junction to a more basal position. The epithelial cells are polyploid and larger than the imaginal ring cells from which they derive. The adult salivary epithelial cells do not undergo mitosis as they do not express phosphorylated Histone H3, yet total cell numbers increase within 2 days of eclosion. By using genetic tools designed for the MARCM lineage tracing technique we have shown that the polyploid cells of the cuboidal epithelium lose chromosomes during the division period and appear to be using amitosis as a mechanism to increase cell number. Amitosis is a form of cell division undertaken by polyploid cells that does not require establishment of a mitotic spindle and results in chromosome loss. Amitosis may be an unusual mechanism to be utilised in primary formation of a tissue but it may be the only efficient way for polyploid cells to increase in number. The adult Drosophila salivary gland will serve as a model for genetic analysis of this mode of division.

639V Cling film – a novel regulator of epithelial morphogenesis Clara-Maria Ell1,2,3, George Pyrowolakis1,2,3 1) Spemann Graduate School of Biology and Medicine (SGBM), Albert-Ludwigs University of Freiburg, Germany; 2) CIBSS - Centre for Integrative Biological Signalling Studies, Albert-Ludwigs University of Freiburg, Germany; 3) Faculty of Biology, Institute for Biology I, Albert-Ludwigs University of Freiburg, Germany

Epithelial morphogenesis is essential for the transformation of simple epithelial sheets into complex organs. The Drosophila larval wing undergoes drastic changes during pupal development and has served as a valuable model to address the underlying mechanisms at tissue-, cell- and subcellular-level. Changes include proliferation, cell rearrangements, cell shape modulation and drastic extracellular matrix (ECM) remodeling, all of which are tightly regulated by intrinsic and extrinsic cues and forces. Here we describe Cling film (Cling), a target of patterning cues in larval wing discs, as a novel regulator of wing morphogenesis. Cling encodes a transmembrane protein with a large, multi-domain extracellular region which predominantly localizes to the apical cell surface. Expression of Cling starts in a spatially defined pattern in late larval wing discs and becomes more uniform in the early pupal stages. Adult wings of cling mutants are severely malformed and strongly folded. We could show that defects in cling mutants manifest in early pupal stages and that cling mutant pupal wings increase their surface normally but fail to stretch out along the proximo-distal axis resulting in heavily folded wings. The phenotype is accompanied by failure to degrade the apical ECM, while genetic ablation of the apical ECM component Dumpy can fully restore wing expansion in cling mutants. Our results indicate that Cling is critically involved in patterned ECM remodeling during early wing morphogenesis. We are currently investigating interactions of Cling with structural ECM components and the ECM degrading proteases Stubble and Notopleural.

640V dysfusion negatively regulates JAK/STAT signaling to constraint the invasive cell population Jhen-Wei WU1, Chuheh-Wen Wang1, Ruo-Yu Chen3, Liang-Yi Hung1, Yu-Chen Tsai1, Yu-Ting Chan1, Yu-Chiuang Chang2, Anna C-C Jang1 1) Department of Biotechnology and Bioindustry Sciences National Cheng Kung University, 1 University Rd, Tainan City 70101, Taiwan; 2) Institute of Biomedical Sciences, National Sun Yat-sen University, 70 Lien-Hai Rd, Kaohsiung 804, Taiwan; 3) Department of Life Science and Life Science Center, Tunghai University, No.1727, Sec.4, Taiwan Boulevard, Taichung City 407224, Taiwan

Cell migration is a critical process for embryonic development and cancer metastasis. How the epithelial cells are selected and adapt migratory cell fate remains unclear. To decipher the underlying mechanism, we apply border cells (BCs), a small group of cells disseminating from the epithelium and migrating through germ cells Drosophila oogenesis, as a model to study collective cell movement. In a forward genetic screen, we found that overexpression of dysfusion (dysf) which severely impeded BC migration. Functional analysis further shows that depletion of dysf in BCs completely blocked cell mobility but that in polar cells, a specialized pair of cells secreting Upd to transactivate JAK/STAT signaling in the neighboring cells to form a cluster, leads to 64% of BC migration delay. These results indicate the
requirement of Dysf in both border and polar cells during migration. Interestingly, overexpression of UAS-dysf in BCs hampered the recruitment, only 3.2 BC observed in the cohort, in comparison to 6 BC in the wild type. To test whether the reduction of BC number result from JAK/STAT signaling hypoactivation, we examined STAT activity by STAT-GFP reporter under dysf overexpression. In wild type BCs, the intensity of STAT-GFP gradually increased during migration, but ectopically upregulated in the dysf mutant cells, and extra border cells were also observed simultaneously. Consistently, the ectopic BC phenotype caused by JAK or Upd overexpression can be suppressed by overexpression of UAS-dysf. Interestingly, Dysf protein is resided at the nuclear membrane of all germline and follicle cells but specifically reduced in BCs upon migration. When BCs reach the destiny, oocyte, their Dysf becomes undetectable. Moreover, we observed that nuclear/cytosol (N/C) ratio of STAT was greatly reduced up overexpression of dysf in the salivary gland. To elucidate how Dysf regulates STAT nuclear transport, we carried out a biochemical screen to seek for interacting proteins with endogenous knock-in tagged Dysf. We found that Pendulin, a member of the Importin-alpha protein family, interact with Dysf. Therefore, we propose that dysf may determine the size of border cell clusters by regulating JAK/STAT signaling.

641V Characterization of adhesion and secretin GPCRs in the salivary glands and germ cells during Drosophila embryogenesis Sean Riccard, Caitlin Hanlon Quinnipiac University

Embryogenesis requires coordinated cell migration directed by molecular guidance cues for proper tissue development. G-protein coupled receptors (GPCRs) are widely conserved transmembrane receptors that relay extracellular signals to the intracellular environment. Despite their prominence, GPCRs are understudied during Drosophila embryonic development. The adhesion and secretin GPCR subfamilies, characterized by their long N-terminal domains, are an interesting topic for further study because of their role in other developmental processes including cell adhesion and endocrine signaling. Adhesion GPCRs are highly expressed in mouse embryonic kidney structures, promote angiogenesis in cell culture, and are implicated in human skeletal development and disease. Secretin GPCRs are also implicated in human skeletal development as well as retinal angiogenesis in mice. Here, we used RNAi to tissue-specifically knockdown the adhesion and secretin GPCRs to determine their role in the embryonic salivary glands (SG) and germ cells (GC). We found that adhesion GPCRs influence Drosophila SG structure in the late embryo. Knockdown of CG11318 or CG15556 has resulted in irregular SG placement (p<0.05). Furthermore, Crumbs-stained SGs in embryos with a knockdown for CG15556 have shown that the apical domain of these SGs are uneven, not smooth and straight like the wild type SG. Interestingly, knocking down either of these gene types in the GCs has had no effect on GC migration. These findings suggest a role for adhesion GPCRs in proper embryonic SG migration. Our next steps are to observe SG migration in embryos with deficiency and insertion lines that cover CG11318, as well as characterize embryonic expression of CG11318 by in situ hybridization.

642V Dunk Regulates Cortical Localization of Myosin II during Drosophila Cellularization through Interaction with the Scaffolding Protein Anillin Jiayang Chen, Bing He, Melissa Wang Dartmouth College

Cleavage is a common step of early embryonic development, generating a monolayer of epithelial cells at the surface of the embryo called “blastoderm”. In Drosophila, this process is achieved by cellularization, a special form of cytokinesis. Similar to typical cytokinesis, cellularization is initiated by recruiting non-muscle myosin II (“myosin”) to the cleavage furrows. Myosin first forms an interconnected hexagonal array at the base of the invaginating cleavage furrows and subsequently reorganizes into individual contractile rings. We have previously identified a gene dunk that promotes myosin retention at the basal array during early cellularization, but the underlying mechanism is unclear. In this work, we performed a genome-wide yeast two-hybrid screen and identified anillin (Scraps in Drosophila), a conserved cytokinesis scaffolding protein, as the primary binding partner of Dunk. Anillin has been reported to regulate the formation of cytokinetic rings during cellularization, but it is unclear whether it regulates the basal myosin array before ring formation. We found that anillin extensively colocalized with the basal myosin array during early cellularization. This colocalization could be detected as early as the onset of cellularization when myosin was first recruited to the cortex as discrete puncta. During the formation of nascent cleavage furrows, anillin and myosin remained colocalized in the cortical puncta as they moved to the cleavage furrows. In anillin mutant, myosin showed a biased localization to the vertices of the basal array and was depleted from edges, closely resembling the myosin phenotype in dunk mutant embryos. Dunk and anillin also display functional interactions. In dunk mutant embryos, the localization of anillin at the basal array was severely disrupted. Furthermore, embryos doubly heterozygous for anillin and dunk showed synthetic defects during cellularization. Finally, dunk and anillin mutants showed similar synthetic phenotypes when combined with mutations in Bottleneck, an actin-bundling protein functioning to restrain actomyosin contractility during cellularization. Together, our results suggest that Dunk regulates myosin at the basal array by interacting with anillin and regulating its cortical localization. Our work demonstrates a previously unappreciated role for anillin in regulating cortical myosin dynamics in early cellularization and may shed light on the regulation of myosin in other cytokinetic processes.

643V Mechanical bistability of the mesoderm facilitates mesoderm invagination during Drosophila gastrulation Hanqing Guo1, Michael Swan2, Bing He1 1) Dartmouth College; 2) Princeton University

Apical constriction driven by non-muscle myosin II (“myosin”) provides a well-conserved mechanism to mediate
epithelial folding. It remains unclear how contractile forces near the apical surface of a cell sheet drive out-of-plane bending of the sheet and whether myosin contractility is required throughout folding. We developed a CRY2-CIB based optogenetic system (“Opto-Rho1DN”) to inhibit the myosin activator Rho1 during Drosophila mesoderm invagination, a typical epithelial folding process mediated by apical constriction. Stimulation of Opto-Rho1DN resulted in rapid myosin dissociation from the cell cortex and disassembly of apical actin, causing acute loss of actomyosin contractility. Interestingly, we find that inhibition of actomyosin contractility during the early, “priming” stage of invagination causes reversal of invagination (the “Early group”). In contrast, invagination continues when inhibition is imposed during the actual folding step after the tissue passes through a stereotyped transitional configuration (the “Late group”). This binary response to actomyosin inhibition suggests that the mesoderm is mechanically bistable during gastrulation. Through 3D reconstitution, we show that apical relaxation happens in both groups after stimulation but has distinct impact on cells that have bent towards the ventral midline due to apical constriction. In the Early group, the bent cells unbend. In the Late group, however, the bent cells keep bending towards the ventral midline as invagination continues, suggesting the presence of additional mechanical input other than actomyosin contractility. Computer modeling suggests that the observed mechanical bistability can arise from an in-plane compression from the surrounding ectoderm, analogous to a buckling process. In line with this model, we find that the transitional state coincides with an apical-basal cell shortening in ectoderm, which may generate in-plane compression through cell volume conservation. Importantly, ectodermal shortening still occurs in the absence of apical constriction, indicating that they can provide distinct mechanical input for mesoderm folding. Taken together, our results demonstrate mechanical bistability in Drosophila mesoderm during gastrulation and suggest that mesoderm invagination is mediated by a joint action of local apical contractility and global tissue compression to trigger epithelial buckling.

644V  Shaping 3D geometry in tubulogenesis: a PDZ domain-containing protein Arc regulates Crumbs to determine salivary gland morphology in Drosophila embryogenesis Ji Hoon Kim, Kwon Kim, Devin Vertrees, Rika Maruyama, Deborah Andrew Johns Hopkins University School of Medicine, Baltimore, MD

The proper architecture of an organ is inseparable from its optimal functionality and physiology. During development, three dimensional tubular organs arise from two dimensional primordia through dynamic changes in cell shape and arrangement. By using embryonic salivary gland (SG) development as a model system to study tubulogenesis, we have discovered that the FoxA transcription factor Fork head (Fkh) is essential for tube formation and internalization. Thus, a subset of Fkh target genes should have roles in SG morphogenesis. arc, an early-expressed SG gene whose expression in the SG and other tubular epithelia requires Fkh, encodes a large cytoplasmic protein containing two PDZ domains. We have discovered that Arc contributes to overall SG dimensions; loss of arc results in shorter, stubbier SG tubes with more cells in circumference and Arc overexpression results in highly elongated SG tubes with fewer cells in circumference. Similar SG phenotypes were induced by perturbing function of non-muscle myosin II (MyoII) and its antagonistic transmembrane protein Crumbs (Crb) in SG. Both the hyperactivation of MyoII and the suppression of Crb activity resulted in shorter SGs with more cells in circumference, as seen in arc loss-of-function mutants. Moreover, Crb levels were significantly reduced in arc null mutant SGs and Arc overexpression resulted in the mis-localization of Crb protein. Importantly, Arc co-localized with Crb at the cell-cell junctions of SG placode cells, suggesting that the two proteins could interact directly, likely through one of the Arc PDZ domains and the cytoplasmic PDZ-binding motif found at the C-terminus of Crb. Indeed, we have discovered that the first PDZ domain of Arc is required for its co-localization with Crb. Based on these observations, we propose that Arc functions as a newly discovered player in SG morphogenesis. We hypothesize that Arc modulates Crb dynamics by affecting its membrane stability and/or localization through direct physical interactions. We further propose that the inhibitory actions of Crb on MyoII serve to limit MyoII activity, consequently limiting the number of SG cells that internalize at any given time and ultimately shaping the final dimensions of the mature SG tube.

645V  Cell polarity determinant Dlg1 regulates the spatial organization and contractile behavior of non-muscle myosin II during tissue morphogenesis Melissa Fuentes, Bing He Dartmouth College

Contractile forces generated by non-muscle myosin II (“myosin”) are critically involved in many morphogenetic processes in animal development. In this work, we found that the cell polarity determinant Dlg1 regulates the spatial distribution and contractile behavior of myosin during tissue folding and tissue elongation in early Drosophila embryos. During Drosophila gastrulation, ventrally localized mesodermal cells undergo apical constriction and invagate to form a ventral furrow. We found that depletion of Dlg1 disrupts the transition between apical constriction and invagination without affecting the rate of apical constriction, the main driving force for furrow formation. In Dlg1 deficient embryos, cells adjacent to the constriction domain (“flanking cells”) display a reduced level of apical adherens junctions and ineffective apical myosin contractions. These defects result in overstretching of the apical domain of the flanking cells and weakening of mechanical coupling between the mesoderm and the neighboring ectoderm, which we show is sufficient to cause delay in invagination. After ventral furrow formation, the germband epithelium doubles in length along the anterior-posterior axis by shortening along the dorsal-ventral axis, a process mediated by cell intercalation. During cell intercalation, planar polarized localization of myosin facilitates patterned shrinking of anterior-posterior
junctiional myosin, an increase in medio-apical myosin, and ectopic accumulation of myosin along the lateral membrane. Further mutant analysis demonstrates that the SH3 and GUK domains of Dlg1 are both important for the proper spatial organization of myosin during germband extension. The myosin regulator Rho-associated kinase (Rok) displays a similar mis-localization pattern as myosin upon loss of Dlg1. These defects are accompanied by ectopic apical constriction in the germband and a reduced rate of tissue extension. Together, our findings reveal the function of Dlg1 in regulating subcellular distribution and/or contractile behavior of myosin in multiple tissues under various developmental contexts.

646V  The role of Scabrous in long distance Notch signaling during bristle patterning **Adam Presser**, Ginger Hunter Clarkson University

Correctly ordered thoracic bristles are necessary for the fly to effectively sense its environment. Notch signaling is critical for the patterning of these bristles: cells in the developing notum epithelium self-organize into Delta-expressing bristle precursor cells and Notch-activated epithelial cells. In order to maintain wildtype bristle spacing, long-distance signaling is necessary. This can occur via either filopodia-mediated Notch signaling or paracrine signaling by diffusible Notch regulators. Scabrous is a secreted protein expressed in bristle precursor cells, and the dynamics of Scabrous signaling in live notum are under-characterized. Based on the literature, we hypothesized that Scabrous acts as a positive regulator of Notch signaling at a distance. Furthermore that Scabrous activity is independent of filopodia-mediated, long-range Notch signaling. Here, we describe several aspects of Scabrous signaling including the location of Scabrous within the patterning tissue using live confocal imaging of Drosophila pupae and fluorescence-stained, dissected pupal nota. We also analyze Notch signaling in response to changes in Scabrous expression levels. These data will help to improve current models of tissue patterning driven by Notch-mediated lateral inhibition.

647V  How to form and maintain a monolayered epithelium: the role of integrins **Lourdes Rincón-Ortega**, Acaimo González-Reyes, María Dolores Martín-Bermudo Centro Andaluz de Biología del Desarrollo, CSIC-Univ. Pablo de Olavide, Sevilla, Spain

Integrins are essential proteins that connect the cell with the extracellular matrix (ECM) and have an important role in the development and maintenance of epithelia. They are implicated in many epithelial diseases and also in tumor progression. To study integrin function in epithelial homeostasis, we use the follicular epithelium (FE) of the *Drosophila* ovary as model system.

The monolayered FE present, in the developing egg chambers of the ovary, follicle cells (FCC), the most abundant cell type, and polar cells, that act as signaling centers for the patterning of the FE. Individual egg chambers are connected by a line of stalk cells. The FE is surrounded by a specialized ECM known as basement membrane (BM), which supports egg chambers growth during oogenesis. The FE is connected to the BM through integrins. Integrins are essential to maintain the architecture of the FE, since the absence of integrins induces extra-layers of FCC at the poles of the egg chambers (Fernández-Miñán et al., 2007). In addition, we have recently observed that lack of integrins produces alterations in the number and distribution of polar and stalk cells, which results in aberrant follicles.

In this project, we aim to understand how these phenotypes arise, thus gaining insight into the role of cell-ECM interactions mediated by integrins during epithelial morphogenesis and maintenance. Our results show that integrins control the number of polar and stalk cells by regulating the proliferation rate and activation of signaling pathways in their precursors. Furthermore, analysis of cell division in *vivo*, leads us to propose that the formation of extra-layers could be due to a combination of aberrant spindle orientation, cellular tension and BM mechanical properties. Taking all together, our findings show that integrins regulate epithelia development and maintenance by different molecular and cellular mechanisms, including activity of signaling pathways, spindle orientation and mechanical properties of cells and their surrounding BM.

648V  Myosin XV regulates basal filopodia formation during bristle patterning **Rhiannon Clements**¹, Aidan Cahill¹, Luke Cowert², Ginger Hunter² 1) Clarkson University, Potsdam, NY; 2) The Pennsylvania State University, University Park, PA

The ability of Notch signaling to drive the formation of a broad range of biological patterns relies, in part, on the activity of cellular protrusions that allow contact between cells at a distance. One example of this is the patterning of sensory bristles on the thorax of Drosophila melanogaster. In this tissue, thin, dynamic, actin-based, filopodia which extend from the basal surface of the patterning epithelia and are essential for long-range Notch signaling. We have identified Myosin XV as a regulator of the formation and maintenance of basal filopodia in the notum. Myosin XV has previously been shown to localize to, and play a role in the dynamics of, filopodia in Drosophila as well as in other cellular protrusions in mammals, including stereocilia. Using a combination of genetics, cell biology, and confocal imaging, we find that Myosin XV contributes to the density of sensory bristles through its role in regulating basal filopodia dynamics. Currently we are using a CRISPR-based strategy to generate a null mutation in the myosin XV gene. We observe that Notch signaling is affected by specific targeting of basal filopodia, through the use of a transcriptional reporter of Notch signaling. Together these results support a role for basal filopodia in lateral inhibition during bristle patterning.
649V Scraps, an anilin, and Nebbish, a kinesin, are integral components of a Fox transcription factor-regulated subnetwork that mediates specific cardiac progenitor cell divisions Md Rezaul Hasan1,2,3, Rajnandani Katariya1,2, Andrew Kump1,2,2, Manoj Panta1,2,3, Kristopher Schwab1,2,3, Mark Inlow1-4, Shaad Ahmad1,2,3 1) Department of Biology, Indiana State University, Terre Haute, IN; 2) The Center for Genomic Advocacy, Indiana State University, Terre Haute, IN; 3) Rich and Robin Porter Cancer Research Center, Indiana State University, Terre Haute, IN; 4) Department of Mathematics and Computer Science, Indiana State University, Terre Haute, IN

Forkhead/Fox transcription factors (TFs) mediate multiple cardiogenic processes in both mammals and Drosophila. We showed previously that the Drosophila Fox genes jumeau (jumu) and Checkpoint suppressor 1-like (CHES-1-like) control three categories of cardiac progenitor cell division—asymmetric, symmetric, and cell division at an earlier stage—by regulating Polo kinase activity, and mediate the latter two categories in concert with the TF Myb. Those observations raised two questions: whether other Fox TF-controlled genes mediating cardiac progenitor cell divisions were also regulated by both CHES-1-like and jumu in a polo-like manner and whether such Fox-regulated genes mediated all three categories of cardiac progenitor cell division or a subset thereof. By comparing transcriptional expression profiles of wild-type, jumu loss-of-function, and CHES-1-like loss-of-function mesodermal cells, we identified multiple genes transcriptionally activated by jumu, but not regulated by CHES-1-like. Phenotypic analysis of mutations showed that two of these exclusively jumu-regulated targets, the kinesin-encoding gene nebbish (neb) and the anilin-encoding gene scraps (scra) are required for only two of the three categories of jumu-regulated cardiac progenitor cell division: symmetric and cell division at an earlier stage. Synergistic genetic interactions between neb, scra, jumu, and polo; between neb and Myb; the absence of such synergistic interactions between either scra and CHES-1-like or neb and CHES-1-like; and the rescue of solely symmetric and earlier cardiac progenitor cell division defects in jumu mutants by ectopic cardiac mesoderm-specific expression of neb demonstrate that scra and neb comprise an exclusively jumu-regulated subnetwork mediating a specific subset of cardiac progenitor cell divisions. Preliminary data from our phenotypic analysis of other exclusively jumu-regulated genes suggests that the kinesin-encoding gene pavarotti, the citron kinase-encoding gene sticky, and the Rho GTPase-encoding gene tumbleweed may be other components of this subnetwork. Using additional genetic interaction and rescue assays, we are attempting to position neb and scra topologically relative to each other and these other potential subnetwork components. Collectively, our results illustrate how an individual regulator can utilize different combinations of downstream effectors to control distinct developmental processes.

650V Exploring the mechanistic roles of APC in the Armadillo/β-catenin destruction complex Katherine Gerber, Julia Kiefer, Emily Errickson, Sonia Hafiz, Matthew Krause, Carmen Navia, Hannah Salvucci, David Roberts Franklin & Marshall College, Lancaster, PA

The tumor suppressor, Adenomatous Polyposis Coli (APC), is an important negative regulator of the Wg/Wnt signaling pathway and is inactivated in nearly 80% of all colon cancer cases. APC participates in a multi-protein “destruction complex” that phosphorylates the proto-oncogene, Armadillo/β-catenin (Arm/β-cat), thereby targeting Arm/β-cat for ubiquitin-mediated proteolysis. Despite nearly 30 years of research on APC, its precise mechanistic role in the destruction complex remains unknown. APC contains several Arm/β-cat binding sites, and prior research from several groups has suggested that these sites play important mechanistic roles in the destruction complex, suggesting specific models of how APC contributes to the destruction complex. We have been testing proposed models using a structure/function approach using Drosophila APC2 as a model system. Surprisingly, our previous research demonstrated that Arm/β-cat binding sites in APC2 are dispensable for Arm/β-cat destruction in cells of the embryonic epidermis that do not receive Wg/Wnt signaling, but are required in cells that receive Wg/Wnt signaling. These findings suggest that Arm/β-cat binding sites on APC and Axin could be redundant in cells that do not receive Wg/Wnt signaling, but that APC’s Arm/β-cat binding sites may play a unique mechanistic role(s) in cells that receive Wg/Wnt signal. To test the redundancy hypothesis, we have generated APC2 and Axin transgenes either containing or lacking Arm/β-cat binding sites. Additionally, to explore that possibility that APC’s Arm/β-cat binding sites have a unique mechanistic role, we have generated APC2 transgenes that replace the Arm/β-cat binding sites with Arm/β-cat binding sites from other Arm/β-cat-interacting proteins such as α-catenin, TCF, and Axin. To date, our results indicate that APC’s Arm/β-cat binding sites are partially replaceable by Arm/β-cat binding sites from TCF and Axin, but not the Arm/β-cat binding site from α-catenin. Collectively, these findings suggest that APC’s Arm/β-cat binding sites do not function simply to recruit Arm/β-cat into the destruction complex, but rather they also likely play a more complex mechanistic role(s). Furthermore, it suggests that specific contacts to Arm/β-cat are required for efficient Arm/β-cat destruction.

651V C. elegans Notch proteins are tuned to lower force thresholds than Drosophila Notch, bypassing the requirement for Epsin-mediated ligand endocytosis. Paul Langridge1,2, Alejandro Garcia-Diaz1, Jessica Chan1, Iva Greenwald1, Gary Struhl1 1) Augusta University, Georgia, GA; 2) Columbia University, New York, NY

The conserved transmembrane receptor Notch has diverse and profound roles in controlling cell fate during animal development. In the absence of ligand, a Negative Regulatory Region (NRR) in the Notch ectodomain adopts an autoinhibited confirmation, masking a protease cleavage site; ligand binding induces cleavage of the NRR, leading to
Notch ectodomain shedding as the initiating step of signal transduction. In *Drosophila* and vertebrates, recruitment of transmembrane protein ligands by the endocytic adaptor Epsin, and their subsequent internalization by Clathrin-mediated endocytosis, exerts a «pulling force» on Notch that is essential to expose the cleavage site in the NRR. Here, we show that Epsin-mediated endocytosis of transmembrane ligands is not required to activate the two *C. elegans* Notch proteins. Using an *in vivo* force sensing assay in *Drosophila*, we present evidence (i) that the *C. elegans* NRRs are tuned to lower force thresholds than the NRR of *Drosophila* Notch, and (ii), that this difference depends on the presence of a “Leucine plug” that occludes the cleavage site in the *Drosophila* and vertebrate Notch NRRs but is absent from the *C. elegans* Notch NRRs. Our results thus establish an unexpected evolutionary plasticity in the force-dependent mechanism of Notch activation and implicate a specific structural element, the Leucine plug, as a potential determinant.

652V  Structural basis of the Calpain A:Cactus (IκB) complex reveals fit induced and competition based mechanisms that alters NFκB activity in embryonic patterning and the immune response Alison Julió, Paloma Dias e Vasconcellos, Priscila Gomes, Maira Cardoso, Pedro Pascutti, Ethan Bier, Paulo Bisch, Helena Araujo. 1) Instituto de Ciências Biomédicas, Universidade Federal do Rio de Janeiro, RJ, Brazil; 2) Instituto de Biofísica Carlos Chagas Filho, Universidade Federal do Rio de Janeiro, RJ, Brazil; 3) Section of Cell and Developmental Biology, University of California, San Diego, La Jolla, CA, USA; 4) Tata Institute for Genetics and Society, University of California, San Diego, La Jolla, CA, USA; 5) Instituto Nacional de Entomologia Molecular (INCT-EM), Universidade Federal do Rio de Janeiro, RJ, Brazil.

Toll pathway activation during embryogenesis and during the innate immune response leads to N-terminal phosphorylation and subsequent degradation of inhibitor Cactus/IκB proteins through the proteasome. This event releases Dorsal/NFκB transcription factors that translocate to the nucleus and regulate gene expression. Cactus is also a target of proteolysis by Calpain A, producing a Cactus E10 fragment and regulating the levels of free Cactus that may interact with the Toll pathway. However, structural and biophysical requirements involved in the interaction of Calpain A and Cactus remains unknown. Here, we report the structural requirements for the action of Calpain A in the control of NFκB in Drosophila. We have modeled the Calpain A:Cactus, Cactus E10 and Dorsal:Cactus complexes, to define the previously unknown domains involved for complex establishment and maintenance. Molecular dynamic simulations suggested that the N-terminal region of Cactus undergoes wide conformational rearrangements upon interaction with Calpain A. Our results indicate that Cactus acts as a modulator in the cleavage process, by approximating the cleavage site to the catalytic site of Calpain A supporting a fit induced model. In addition, our simulations indicate that the Calpain A:Cactus complex is sustained by salt-bridges formed along the Cactus alpha helix, anti-parallel ankyrin repeats and Calpain A EF hands 2, 3, 5 and CBSW domains. Subsequent mutational analysis confirms our structural predictions. Functional analysis using Cactus alleles mutated at the cleavage site and catalytically-dead Calpain A mutants demonstrates that the Cactus cleavage event by Calpain A is fundamental in modulating NFκB responses triggered by Dorsal nuclear translocation. Interestingly, our results revealed an unexpected function for Calpain A independent of catalysis, suggesting that Calpain A may also modulate Toll signal transduction by sequestering Cactus. These results indicate that competition between Calpain A:Cactus and Dorsal:Cactus complexes is an important aspect in Toll responses. New CRISPR Calpain A alleles reinforce the action of Calpain A to modify the Dorsal gradient, affecting target genes during early embryogenesis, and to modulate immune responses mediated by Toll. Taken together, these results provide a mechanistic view for the structural requirements involved in Calpain A modulating NFκB signaling.

653V  Analysis of pMad and Medea Expression in BMP Pathway in Drosophila with Multiple Fluorescent Proteins Hung-Yuan (Zeke) Chen, Gregory Reeves Texas A&M University

Morphogen gradients are important in early Drosophila embryo development. One such gradient, the BMP/Dpp gradient, patterns the dorsal region of the embryo, which induces a series of protein interactions. After BMP-like morphogen Dpp transports to the dorsal region and binds to type I receptors Thickevins (Tkv), Tkv is activated by type II receptor Punt and then phosphorylates Mothers against Dpp (Mad). The phosphorylated Mad (pMad) and Smad4 homolog Medea translocate together to the nucleus to regulate the target gene. Compared to the Dpp gradient, pMad and Medea are procedurally closer to gene regulation. However, the transport and expression of pMad and Medea are rarely discussed. Their expression depends on the morphogen concentration which is determined by the cell location. In this work, we tag eGFP and mScarlet I to Mad and Medea and mtagBFP to Histone 2A for locating the nuclei using Crispr-Cas9 techniques and analyze the embryo images after injection into flies and crossing. Finally, we then do image analysis of 14-cycle embryos to see the expression of these proteins.

654B  Abl tyrosine kinase controls the distribution and propagation of cellular forces by regulating the coherence of an actin network Aravind Chandrasekaran, Akanni Clarke, Philip McQueen, Hsiao-Yu Fa, Garegin Papoian, Edward Giniger. 1) NINDS, NIH, Bethesda, MD; 2) University of Maryland, College Park, MD

Cytoplasmic signaling pathways regulate cellular morphogenesis, but how do the nanoscale dynamics of a signaling protein create structures and direct forces at the multi-micron spatial scales of a cell? We have now compared experimental measurements of actin distribution and dynamics in single axonal growth cones in vivo in the fly wing with the results of single-molecule computational simulations of actomyosin dynamics. The results suggest a simple
We have performed live imaging of the tip of an axon – its growth cone – in vivo in the developing fly wing. This reveals that the core of the growth cone is a mass of non-polarized actin that is undergoing constant, stochastic fluctuations. Those fluctuations are the engine of growth cone motility, as we have shown that net axon advance comes from a spatial bias in the actin fluctuations that causes the network to take more and longer steps forward than steps back. The bias comes from the cytoplasmic signaling pathways that control actin polymerization and branching. Our data show that Abl tyrosine kinase, a key downstream effector of axon guidance receptors, controls the spatial spread of the actin network: increasing Abl activity causes the growth cone actin core to expand, while decreasing Abl causes the network to contract. Computational simulations now reveal that the mechanism by which Abl controls network size is mediated through modulation of the lengths of actin filaments. Activating Abl causes a net decrease in the lengths of actin filaments, reducing their ability to link the actin network together. Consequently, the network as a whole fragments and spreads out in the growth cone, just as we observe experimentally. Conversely, reduction of Abl promotes extension of actin filaments, allowing myosin-dependent contractility to act across long length scales to condense the entire actin network, again, just as we observe experimentally.

These data provide a framework for interpreting the effects not only of Abl, but of many regulators of growth cone function. Moreover, the principles revealed here should apply to many other developmental contexts that rely on cytoplasmic signaling mechanisms to control the spatial distribution of actin-dependent structures and forces.

655C Pelado, a conserved protein that regulates actin dynamics Claudia Molina1, Patricio Olguin2, Álvaro Glavic3, Marek Mlodzik1 1) Icahn School of Medicine at Mount Sinai; 2) Faculty of Medicine, University of Chile; 3) Faculty of Sciences, University of Chile

Actin dynamics are essential in every cell, primarily for cell division, migration, and morphological changes. Actin polymerization can be branched or linear, and it depends on the associated regulatory proteins. There are hundreds of proteins that regulate the actin cytoskeleton dynamics. Competition for actin monomers also occurs between proteins that induce branched or linear actin polymerization. Cell specialization requires actin cytoskeleton transformation to allow the formation of cell structures, like cuticular hairs in *Drosophila*. This is a well characterized and useful study model to analyze proteins that regulate actin dynamics. Structurally, these hairs are mainly formed by linear actin filaments. A screen was performed to identify wing development regulators. A lethal mutation was found on a gene that was named *pelado*. Pelado is a protein conserved throughout the animal kingdom but has no clear function. Epithelial mutant cells for *pelado* shows hair elongation defects. This phenotype was reversed by inducing linear actin polymerization, by reducing branched actin polymerization and by increasing the amount of actin monomers. This suggests that Pelado is involved in the competition for actin monomers. A similar phenomenon occurs in hemocytes, where Pelado is essential to induce filopodia formation in a process that also requires Diaphanos and Profilin. Pelado’s function was also evaluated in A549 human cell line, in a wound healing assay. It has been described that this cell type requires branched actin polymerization to achieve efficient migration. In *pelado* knockout cells, the wound closes faster and the opposite occurs if we introduce the *pelado* fly gene in those cells. Immunoprecipitation assays suggest that Pelado is part of a multiprotein complex that includes Scar/WAVE, Diaphanos and Profilin. These data indicate that Pelado’s function in regulating the actin cytoskeleton is conserved, preventing branched actin polymerization at the same time as favoring linear actin polymerization, through actin monomer competition mechanisms that modulate actin cytoskeleton dynamics. To evaluate the domains of Pelado that are important for its function, a C-terminal deletion was generated that can induce filopodia formation but does not rescue hair formation. This indicates that the C-terminal portion of Pelado has a regulatory function that is essential for certain processes, including hair formation in *Drosophila*.

656A Spd-2 gene duplication suggests cell type-specific mechanisms of pericentriolar material assembly Ryan O’Neill1, Afeez Sodeinde2, Frances Welsh3, Carey Fagerstrom1, Brian Galletta3, Nasser Rusan1 1) National Heart, Lung, and Blood Institute, NIH; 2) Yale, New Haven, CT; 3) University of Washington, Seattle, WA

Centrosomes are the major microtubule organizing center (MTOC) of the cell, ensuring proper spindle formation and chromosome segregation during cell division. Hundreds of proteins make up the centrosome, including many proteins that form the pericentriolar material (PCM) which nucleates microtubules. The specific requirements for MTOC activity varies across cell types, a concept we are just starting to investigate and understand. In this study, we took an evolutionary cell biological approach to gain insight into cell type-specific regulation of centrosome proteins, reasoning that gene duplication could lead to the evolution of centrosome gene duplicates with cell type-specific functions. We first used BLAST to screen 35 sequenced *Drosophila* genomes for duplications of centrosome genes, finding five that were duplicated at least once. Here, we focus on the duplication of *Spd-2* in *D. willistoni*: we refer to the parental gene as *Spd-2A* and the new copy as *Spd-2B*. In *D. melanogaster*, *Spd-2* is known to function by organizing PCM in neuroblasts and spermatocytes. We found that *Spd-2B* is rapidly evolving and lacks the C-terminal 116 amino acid tail of *Spd-2A*. To explore expression and function we made GFP-tagged *Spd-2A* and *Spd-2B* transgenes, including their *D. willistoni* native
regulatory elements, in *D. melanogaster*. Spd-2A is ubiquitously expressed, whereas Spd-2B is only expressed in spermatogenesis. Consistent with their expression, Spd-2A, but not Spd-2B, rescues *spd-2* mutant neuroblast PCM. In contrast, Spd-2B rescues *spd-2* mutant spermatocyte PCM, whereas even when ectopically expressed Spd-2A fails to do so. Thus, after gene duplication Spd-2A lost the ability to organize PCM in spermatocytes, leading to complementary functions for Spd-2A and Spd-2B. We further mapped the changes responsible for this difference in function, finding that removing the C-terminal tail from Spd-2A allowed it to properly organize spermatocyte PCM, whereas adding a C-terminal tail to Spd-2B prevented it from organizing spermatocyte PCM. We infer a model where the C-terminal tail of Spd-2 mediates a priming step that precedes PCM recruitment. Together, these results show that somatic and germline cells have different requirements for PCM, and suggest that Spd-2 is differentially regulated across cell types to satisfy these distinct requirements.

657B  Developing tools to study the actin mesh during *Drosophila* oogenesis Hannah Bailey, Margot Quinlan University of California Los Angeles, Los Angeles, CA

The process of egg development, oogenesis, is highly conserved and crucial for producing offspring. *Drosophila melanogaster* have long served as a model system to understand aspects of egg development including stem cell and germ cell development, meiosis, cell migration, intercellular signaling and mRNA localization. An essential component of oogenesis in *Drosophila* is the presence of a cytoplasmic actin meshwork that persists during mid-oogenesis. This complex actin network is built by the collaboration of actin nucleators, Spire and Cappuccino (Spir and Capu). The composition, organization, stabilization, and removal of the mesh remains unknown. This is, in part, due to the requirement for actin binding proteins in early oogenesis and our inability to visualize removal of the actin mesh because egg chambers expire *ex vivo* just prior to this transition. To overcome these obstacles, I am developing methods to directly observe this meshwork and study the underlying regulatory mechanisms of the mid to late oogenesis transition. Specifically, I am developing methods for long-term *in vivo* imaging of oogenesis in *Drosophila melanogaster*. In principle, this approach will make possible study of mesh removal. Studying the roles of Spir and Capu has also been challenging due to differences in endogenous temporal control and the tools available, including the commonly used drivers of the bipartite GAL4/UAS system. I am developing improved drivers for the GAL4/UAS system to better match endogenous expression timing, allowing for careful characterization of these actin nucleators *in vivo*. Lastly, using the Auxin Inducible Degradation (AID) system, I am testing candidates to identify mesh components and regulators. Altogether, my work will lead to a greater understanding of dynamic actin rearrangements during development and facilitate detailed characterization of the actin mesh that can translate to studies in other organisms.

658C  Dynein acts to cluster glutamate receptors and traffic the PIP5 kinase, Skittles, to regulate postsynaptic membrane organization at the neuromuscular junction Amanda L. Neisch1, Thomas Pengo1, Adam W. Avery1,2, Min-Gang Li2, Thomas S. Hays1 1) University of Minnesota, Minneapolis, MN; 2) Oakland University, Rochester, MI

Cytoplasmic dynein is essential in motoneurons for retrograde cargo transport that sustains neuronal connectivity. Little, however, is known about dynein’s function on the postsynaptic side of the circuit. Using genetic, immunolocalization, and electrophysiology studies, we have identified distinct postsynaptic roles for dynein at neuromuscular junctions (NMJs). We have found that dynein punctae accumulate specifically on the postsynaptic side of glutamatergic synaptic terminals. Postsynaptic dynein is required for the localization of P(4,5)P2, a phospholipid membrane component, and a number of membrane-associated proteins including components of the spectrin cytoskeleton. Skittles, a phosphatidylinositol 4-phosphate 5-kinase that produces P(4,5)P2, organizes the spectrin cytoskeleton, also localizes specifically to glutamatergic synaptic terminals and this localization is dynein dependent. Further, depletion of postsynaptic dynein results in enlarged ionotropic glutamate receptor (iGluR) clusters and an increased amplitude and frequency of mEJPs. P(4,5)P2 levels do not affect iGluR clustering and dynein does not affect the levels of iGluR subunits at the NMJ, suggesting a unique transport independent function of dynein at the NMJ in clustering iGluRs. As dynein punctae closely associate with iGluR clusters, we propose that dynein physically stabilizes iGluRs at the postsynaptic membrane for proper synaptic transmission.

659A  βH-spectrin Recruits PP2AWaldorf to Crumbs where it Regulates Growth and Apical Domain Stability In *Drosophila* Kristen Browder1,2, Seung-kyu Lee1,3, Elizabeth Klipfell1, Mark Travoni1, Katelyn Wolfgang1, Claire Thomas1 1) Penn State, University Park, PA; 2) Genentech, South San Francisco, CA; 3) National Institute of Aging (NIH/NIA/IRP), Baltimore, MD

Spectrin is a large F-actin crosslinking protein that most famously forms 2D networks in association with the plasma membrane of red blood cells. This ‘membrane skeleton’ confers cell shape and membrane strength during the rigors of circulation. In NON-erythroid tissues, spectrin has additional roles in the endomembrane system. We have previously shown that the apically polarized βH-spectrin (βH), encoded by the *karst* locus in *Drosophila*, is required for the stability of several apical proteins, through the promotion of endosomal recycling to the plasma membrane - so called, ‘dynamic protein stabilization’. The apical protein determinant Crumbs recruits βH to the apical membrane and is itself trafficked in a βH-dependent manner. βH binds to the Hippo/Warts pathway (HWP) regulator Expanded, which mediates Crumbs...
Here we report that a yeast 2-hybrid (Y2H) screen identified the PP2A substrate-specificity subunit Waldorf (a PP2A-PR72/B" isoform) to be a binding partner of βH, via a short, conserved sequence in its C-terminal globular domain. Genetic interaction and molecular epistasis experiments strongly suggest that PP2A with the Waldorf specificity-subunit bound to it (PP2A<sup>Waldorf</sup>) acts as a negative regulator of Crumbs by acting to displace aPKC from Crumbs. Consistent with this notion, mutant versions of Crumbs lacking target residues for aPKC in the FERM-domain binding site do not respond to changes in Waldorf levels. Knockdown of Waldorf leads to reduced growth suggesting that Crumbs crosstalk with the HWP is a primary target of this homeostatic regulation. Overexpression of Waldorf specifically destroys the apical domain suggesting that Waldorf may also play a role in modulating the apical domain.

We also show that Waldorf modulates protein trafficking in a similar way to βH, and its previously reported partner Annexin B9 in that knockdown of Waldorf leads to an increase in Rab7-positive and acidic compartments, suggesting that PP2A<sup>Waldorf</sup> also normally acts by suppressing lysosomal trafficking, most likely in favor of recycling pathways. Our results support a model in which Crumbs recruits βH in a complex with the HWP activator Expanded, and PP2A<sup>Waldorf</sup> bound to βH acts in a homeostatic fashion to limit Crumbs activation of the HWP by displacing aPKC. This in turn limits the amount of growth suppression caused by Crumbs-dependent HWP activation.

660B  Cullin 3 promotes polarization of aPKC phosphorylated differentiation determinants during asymmetric neuroblast division Cheng-yu Lee, Hideyuki Komori, Noemi Rives-Quinto, John Bugay Univ Michigan

The aPKC/Par-6 complex is widely used in polarizing cortical localization of protein determinants that promote proper specification of cell identity and cell functionality. The prevailing model suggests that aPKC phosphorylation displaces its target proteins from the aPKC/Par-6 cortical domain into cytoplasm by perturbing their interactions with phospholipids or adaptors, leaving unphosphorylated proteins enriched in the opposite cell cortex. It remains unclear if phosphorylation-induced exclusion from the cortex is a generalizable mechanism for aPKC-mediated polarization of numerous downstream proteins during development and homeostasis. During asymmetric neuroblast division, aPKC kinase activity in the apical cortex regulates basal localization of Notch antagonists including Numb and their segregation into neuroblast progeny where they promote differentiation by downregulating Notch signaling. In contradiction of the prevailing model, we found that aPKC kinase activity levels positively correlate with polarized Numb accumulation in the basal cortex of mitotic neuroblasts. Analyses of novel fly or human numb alleles that encode missense mutations at two conserved aPKC phosphosides indicate that the phosphomimetic but not the non-phosphorylatable form of Numb protein asymmetrically localizes to the basal cortex of mitotic neuroblasts. These results indicate that aPKC phosphorylation positively induces polarization of Numb. We screened for genes that are required for Numb segregation during asymmetric neuroblast division in numb-hypomorphonic brains, which show a mild supernumerary neuroblast phenotype due to reduced Numb activity in neuroblast progeny destined to differentiate. We identified the cullin 3 (cul3) gene as a novel regulator of Numb polarization in mitotic neuroblasts. Mitotic cul3-null neuroblasts show aPKC and Numb localized uniformly throughout the cortex, reducing Numb levels in neuroblast progeny and leading to their reversion into supernumerary neuroblasts due to defects in downregulation of Notch signaling. Consistent with defects in aPKC-induced Numb polarization, increased aPKC kinase activity levels restore Numb polarization in the basal cortex of mitotic cul3-null neuroblasts and restore differentiation in their progeny. We propose that the non-proteolytic function of Cul3 promotes aPKC phosphorylation-induced Numb polarization in mitotic neuroblasts by facilitating efficient Numb phosphorylation by aPKC.

661C  Unraveling Positive and Negative Feedback in Planar Cell Polarity Alexis Weiner, Kaye Suyama, Jeffrey Axelrod Stanford University

Planar Cell Polarity (PCP) polarizes cells along an axis parallel to the tissue plane, and this results in organized cell polarity across entire tissues. PCP includes both positive and negative feedback mechanisms to cause the polarization of two core protein complexes. However, the interdependency between molecular players and lack of genetic techniques has left the role of feedback mechanism mired in mystery. To uncover a more detailed mechanism of polarization we have developed a novel genetic ‘Velcro’ approach to force molecular players in the pathway to cell-cell junctions. This method allows us to parse the specific steps of polarization and differentiate between the role of positive and negative feedback in PCP. It also lets us ask the important evolutionary question of whether cells within PCP retain the single cell system of polarization when we use this tool together with a mutant that removes cell-cell PCP communication. Using the novel ‘Velcro’ tool, we have shown by proof of concept that forcing a single component, Dsh, to cell-cell junctions at the clonal boundary we can reverse cell polarity.

We can dissect the role of clustering as the mechanistic function underlying positive feedback by expressing clustering mutants of Dsh, Pk and Fmi. We have engineered flies expressing proteins that are oligomerization deficient and will position them using ‘Velcro’. This will determine if self-assemblies are required for polarization. Finally, we can test if this is also required for cell autonomous polarization. These experiments will reveal the role of positive feedback during cellular polarization and if PCP retains the ability of single cells to use this positive feedback to polarize autonomously.

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We will also tackle the proposed long-range negative feedback and role of ‘mutual’ exclusion in the segregation of proximal and distal complexes in PCP. Using our novel ‘Velcro’ technique, we are testing if proximal proteins such as Vang and Pk are excluded when Dsh is forced to localize asymmetrically at clonal boundaries. We also will ask if this exclusion requires clustering by introducing oligomerization deficient Dsh mutants. Second, we will ask if this exclusion can happen cell autonomously by removing cell-cell connections with the Fmi mutant. Furthermore, we will investigate the minimal requirements for this exclusion by removing other molecular players such as Fz, Dgo, Vang, and Pk. Lastly, we will attempt to perform the same exclusion experiments by forcing Pk, the proximal counterpart to Dsh. We will evaluate if distal complexes are excluded in alignment with the idea of ‘reverse’ mutual exclusion. Ultimately this will complement the first set of experiments and reveal the role of long-range negative feedback in PCP signaling. With these tools and experiments, we aim to address the previously inaccessible questions concerning the contributions of positive and negative feedback in PCP.

662A The Establishment and Maintenance of Centrosome Asymmetry in Neural Stem Cells Roberto Segura, Clemens Cabernard University of Washington

Asymmetric cell division (ACD) is an evolutionarily conserved process where proteins, organelles, and mRNAs are unequally partitioned between daughter cells. This maintains the stem cell population, where progenitor cells undergo ACD to produce a self-renewing stem cell and a differentiating daughter cell. Organelles, such as centrosomes, can be partitioned asymmetrically between daughter cells, and this is observed in many diverse cell types. In Drosophila neural stem cells, called neuroblasts, the differentiating ganglion mother cell receives the older mother centrosome, while the renewing neuroblast receives the younger daughter centrosome. This biased inheritance is determined by differences in molecular identity, which is mediated by the phosphorylation activity of Polo kinase. Preliminary data has implicated the involvement of protein phosphatase 4 (PP4) in this cascade, and the extent of PP4’s molecular involvement in this process is not fully understood. These molecular identities are speculated to impact the other factors during ACD. Additionally, centrosomes retain distinct clusters of mRNA, and this retention is theorized to be facilitated by the molecular identity of the centrosome. However, the spatiotemporal mechanisms that regulate centrosome asymmetry and its functional impact are unknown. I hypothesize that (1) dephosphorylation of cytoskeletal proteins, such as gamma-tubulin, contribute to the establishment and maintenance of centrosome asymmetry, and (2) biased mRNA localization is dependent on centrosome asymmetry. To address these hypotheses, I will utilize fluorescent in vivo imaging and novel optogenetic methods to elucidate the spatiotemporal regulation and functional importance of centrosome asymmetry. These experiments will reveal the regulation of this process within the context of ACD, which will broaden our understanding of its importance in other processes, such as stem cell differentiation and the onset of cancer.

663B Regulated demolition in muscle remodeling: a T-tubule membrane disassembly pathway maintains muscle function shravan girada, jen Nguyen, Tzu-Han Lin, Amy Kiger Section of Cell and Developmental Biology, Division of Biological Sciences, University of California, San Diego

Muscles are complex cells with specialized structures for contraction. While arrays of sarcomeres perform contraction, it is the network of plasma membrane invaginations, called Transversal (T)-tubules, that coordinate sarcomere contractions. We study T-tubule requirements in body wall muscles of Drosophila at different growth stages, which allows live imaging of the membrane network. We uncovered an endogenous requirement for shibire, a Dynamin large GTPase, in the initiation of T-tubule disassembly within a wildtype developmental muscle remodeling program. Consistent with its role in disassembly, overexpression of shibire led to inappropriate T-tubule fragmentation and defective muscle function for larval mobility. These results indicate the importance for regulation of a normal dynamin function in T-tubule disassembly that may also hint at how dominant mutations in a conserved human DNM2 lead to centronuclear myopathy (CNM).

Previously, we implicated Class II PI3-kinase, Pi3K68D (or PI3KC2), in abdominal muscle remodeling defects associated with loss of Mtm PI3-phosphatase, a homolog of human MTM1 linked to a recessive (X-linked) form of CNM. We now identify a requirement for Pi3K68D in T-tubule disassembly at initiation of regulated muscle remodeling. Remarkably, disruption of Pi3K68D function also prevented the ectopic T-tubule fragmentation induced by shibire overexpression. Interestingly, mammalian PI3KC2A activity has been shown to recruit DNM2 for its role in endocytic vesicle formation, raising the question whether a shared pathway is also involved in T-tubule fragmentation. Our current studies unveil Pi3K68D phosphoinositide dependence, sites of Pi3K68D and Shibire function and the contributions of other endocytic pathway members in T-tubule disassembly. Altogether, our results suggest shared elements between the mechanism for dynamin-mediated endocytosis and T-tubule membrane disassembly and elevate the prospects of human PI3KC2A-targeted therapies for both dominant DNM2- and recessive MTM1-related myopathies.

664C Systematic functional analysis of Rab GTPases in neuronal development and maintenance Ilsa-Maria Daumann, Friederike Kohrs, P. Robin Hiesinger Freie Universitaet Berlin
Neurons are morphologically elaborate and long-living cells. Both features put special demands on membrane trafficking. Rab GTPases are regulators of membrane trafficking and the *Drosophila* genome encodes 26 rab genes. All rabs are expressed in the nervous system and 13 of the 26 rabs exhibit nervous system-enriched expression patterns. To facilitate systematic functional analyses of neuron-specific membrane trafficking, we have generated a complete rab null mutant collection. Remarkably, the complete loss of any of the 13 nervous system-enriched Rabs did not cause obvious defects concerning fly survival, fertility, or morphology under laboratory conditions. However, under challenging conditions including continuous neuronal stimulation or temperature variation all rab mutants revealed specific sensitivities affecting development, function, or maintenance of the nervous system. Our observations suggest that the majority of Rabs serve modulatory functions that ensure robustness of development or function to challenging environmental conditions.

We followed up on our initial systematic characterization with a more detailed analysis of four neuronal or neuron-enriched rabs: rab26, rab19, rabX1 and rabX4. Rab26 has previously been proposed to link synaptic vesicle recycling and autophagy. However, our characterization of the rab26 mutant revealed no autophagosomal or synaptic vesicle defects in flies. Instead, we found an activity-dependent role of Rab26 in receptor trafficking at cholinergic synapses. Our analyses of rab26, rab19, rabX1 and rabX4 revealed synthetic lethality for double mutants of rabX1 with each of the other three, but no other double mutant combination. We will present an update on our characterization of the functional contributions of each of these rabs.

### 665A The STRIPAK complex and microtubule protein transport in *Drosophila* muscle tissue

**Yungui Guo, Erika Geisbrecht Kansas State University**

In humans, Bcl-2-associated athanogene 3 (BAG3) is essential for proteostasis in stressed cells. Some of the known functions of BAG3 include promoting chaperone activity, facilitating aggresome formation, and initiating the destruction of proteins via macroautophagy. Our lab has discovered that *Drosophila* Starvin (Stv), the ortholog of mammalian BAG3, biochemically and genetically interacts with the evolutionarily conserved NUAK serine/threonine kinase, and this NUAK-Stv-Hsc70-4 complex plays a role in the autophagic clearance of proteins. However, the mechanism of how damaged proteins get transported to the sites of degradation is not clear. Based upon literature in neuronal cells that describe a role for the Striatin Interacting Phosphatase and Kinase (STRIPAK) complex in the axonal transport of autophagosomes, we propose the hypothesis that the STRIPAK complex may be required to transport proteins along microtubules (MTs) to the lysosome in *Drosophila* larval muscles. To test for a possible role of the supramolecular STRIPAK complex in BAG3-mediated transport, we have employed a sensitized genetic assay using RNA interference (RNAi) technology. Thus far, our data shows that stv genetically interacts with genes that encode for some members of the STRIPAK protein complex, including Striatin interacting protein (Strip1), MOB kinase activator 4 (Mob4), and Connector of kinase to AP-1 (Cka). Moreover, preliminary observations suggest perturbations in the MT network within muscle cells upon RNAi knockdown of STRIPAK complex members. This project will give us a better understanding of how the NUAK-Stv-Hsc70-4 complex functions with STRIPAK complex in the autophagic clearance of protein and maybe lead to finding some potential drug targets for treating protein aggregate diseases in human.

### 666B MDIS, a mitochondrial DNA exonuclease enforces uniparental inheritance of mitochondrial genome

**Zhe Chen¹, Christian A Combs¹, Yong Chen¹, Annie Lee², Hong Xu¹ 1) National Institutes of Health; 2) Walter Reed National Military Medical Center**

Mitochondrial genome is exclusively transmitted through maternal lineage in animals. The uniparental inheritance was once regarded as a passive outcome of distinct cytoplasmic contents of eggs and sperms. Recent studies demonstrated active mechanisms to remove mitochondrial DNA (mtDNA) during spermatogenesis in various species. However, the physiological significance of mtDNA clearance, or mitochondria uniparental inheritance in general remains a mystery and the factors involved in this process are largely unknown. Using proximity-labeling based proteomic screen, we recovered a putative mitochondrial nucleoid protein MDIS (mitochondrial DNA in sperm), encoded by *CG12162* locus. MDIS is highly enriched in *Drosophila* testis, and specifically in late spermatogenesis stages, when the mtDNA clearance takes place. *MDIS* null flies are male semi-sterile, otherwise completely healthy. Spermatogenesis progresses normally in *MDIS* flies but produces immotile mature sperms that often contain multiple copies of mtDNA nucleoids, indicating that MDIS is required for mtDNA clearance. *In vitro* assay suggests that MDIS is an exonuclease that degrades both single-stranded DNA and double-stranded DNA with overhangs. Importantly, ectopic expression of a mitochondrially targeted *E.coli* exonuclease III in *MDIS* flies, effectively removes mtDNA remnants and largely restores male fertility, demonstrating that failure to eliminate mtDNA causes immotile sperms and male infertility. Surprisingly, we observed pronounced nuclear DNA fragmentations in mature sperms of *MDIS* flies, suggesting a potential detrimental consequence of persisting mtDNA in mature sperms. Our results demonstrate that MDIS play essential roles in preventing the transmission of paternal mtDNA to the offspring, and in safeguarding the nuclear genome integrity. MDIS, to our knowledge, represents the first-ever identified factor that specially enforces maternal inheritance of mitochondrial genomes. This will allow future studies to understand physiological significances and underlying mechanisms of this highly conserved, yet mysterious phenomenon.
667C Roles for CG5755, a SLC25A46 ortholog, in mitochondrial morphogenesis during Drosophila spermatogenesis Claire Olson, Caroline Phan, Tommy Mason, Kyle Kowalewski, Sam Kavarana, Vivienne Fang, Karen G Hales Davidson College

Drosophila spermatogenesis involves significant morphological changes in mitochondria and thus provides a useful system to study genes that regulate mitochondrial movement and shaping. The Z2-3738 recessive male sterile fly strain, originating from the Zuker collection, displays defects in spermatogenesis, including male sterility and disorganized sperm bundles with scattered short mitochondrial aggregates, due to a nonsense mutation in CG5755, a testis-specific gene. Its paralog, CG8931, is expressed in most other tissues. The human ortholog, SLC25A46, a member of a mitochondrial solute carrier family, has been shown to localize to the mitochondrial outer membrane and function in mitochondrial fission. SLC25A46 has been implicated in several human neurodegenerative diseases; thus, elucidating the function of its Drosophila orthologs may provide valuable insight into these conditions. Here, we characterize the function of CG5755 by exploring its localization, interactions, and putative role in the nervous system as proposed by others. We observed localization of a CG5755-GFP transgenic protein in the mitochondria of sperm bundles and potentially in the nebenkern. We investigated genetic interactions of CG5755 with genes involved in mitochondrial morphology, including the fusion gene fzo and cristae shaping gene mic60, and observed a synthetic phenotype in combined Z2-3738 and fzo mutants, suggesting that CG5755 may function in a conserved mitochondrial fusion pathway with fzo. Additionally, although current RNA-seq data report CG5755 as a testis-specific gene, another group reported evidence of neurological defects in neuron-specific CG5755 knockdown flies. We conducted experiments to determine whether CG5755-GFP localizes to nervous system tissues and whether Z2-3738 flies show defects in locomotor function. Together, these results confirm the role of CG5755 in mitochondrial shaping in spermatogenesis, provide preliminary evidence about its function in mitochondrial dynamics and its interactions with other relevant genes, and demonstrate whether it is functionally active in the nervous system.

668A Moonlighting of the Golgi protein, Gorab, at the centriole is regulated by its high affinity for centriolar protein Sas6 Levente Kovacs1, Jennifer Chao-Chu2, Sandra Schneider3, Agnieszka Fatalska2,3, Magdalena M Richter2, Emma Stepina2, Marco Gottiordo2, George Tzolovsky3, Nikola S Dzhindzhev3, Maria Giovanna Riparbelli3, Giuliano Callaini3, Michal Dadlez2, Gang Dong4, David M Glover1,2 1) Division of Biology and Biological Engineering, California Institute of Technology, Pasadena, California, USA; 2) Department of Genetics, University of Cambridge, UK; 3) Laboratory of Mass Spectrometry, Institute of Biochemistry and Biophysics, Polish Academy of Sciences, Warsaw, Poland; 4) Max Perutz Labs, Vienna BioCenter, Vienna, Austria; 5) Department of Life Sciences, University of Siena, Siena, Italy

Centrioles are evolutionarily conserved organelles required to assure the fidelity of cell division, motility and signaling. Although the majority of key proteins ensuring proper centriolar structure, duplication and maturation have been described, several new candidates have emerged from protein interaction studies. In a search for interacting partners of the core centriole cartwheel protein, Sas6, of Drosophila, we recently identified Gorab, better known as a Golgi protein mutated in the human disease Gerodermia osteodisplastica. We discovered that Gorab is a moonlighting protein – whereas dimeric Gorab localizes to the trans-Golgi, monomeric Gorab directly interacts with Sas6 dimers at the centriole’s core where it is essential for centriole duplication and symmetry. Here we show that Gorab’s high binding affinity to the limited amount of centriolar Sas6 is a key determinant of Gorab’s localization. Thus, upon Gorab overexpression, there is no increase in its levels at centrioles and the surplus Gorab associates with the trans-Golgi. Meiotic spermatocytes are particularly sensitive and moderate Gorab overexpression results in a Golgi-related cytokinesis defect and consequently male sterility. Upon Sas6 overexpression, however, the amount of Gorab dramatically increases both in the centrioles and in non-centriolar Sas6 aggregates and Gorbi-associated Gorab decreases. Moreover, targeting elevated levels of Sas6 to mitochondria re-localizes endogenous Gorab to this organelle while depleting Gorab at the Golgi. In line with these observations, in vitro interaction assays reveal that Sas6 can disrupt preformed Gorab homodimers and disassociate Gorab from its Golgi-associated interactor Rab6. Strikingly, Sas6 overexpression can rescue the cytokinesis defects of Gorab-overexpressing males and restore their fertility. Thus, the high affinity of the Sas6 dimer for Gorab monomers ensures that Gorab gives priority to its moonlighting function at the centriole allowing remaining Gorab to dimerize and localize to the Golgi for its official duties.

669B Essential functions of gish in nuclear positioning during early embryogenesis Lingkun Gu, Mo Weng University of Nevada, Las Vegas

Nuclei are often actively positioned for specialized cellular functions or developmental purposes and failures in correct positioning can lead to dysfunctions and defects. Early embryos of Drosophila melanogaster are tasked with positioning hundreds of nuclei because the embryogenesis begins with 13 synchronized nuclear divisions within a syncytium. Although nuclei initially localize in the anterior half of the embryo, during nuclear cycle 4-6, they are evenly spread along the anterior-posterior axis in an actomyosin-dependent process called axial expansion. Then, during cycle 7-9, nuclei migrate and distribute evenly in the syncytial cortex, followed by the last four nuclear divisions. The even nuclear spacing ensures that cells of similar sizes are generated during the cleavage stage. However, the molecular mechanisms of axial expansion and cortical migration are still poorly understood. Here, we identify a plasma membrane-associated
kinase *gilgamesh (gish)* as an important player in this complex process. *gish*-depleted embryos show defects in axial expansion: nuclear spreading to the posterior half of the embryo is delayed and the nuclei preferentially localize to the anterior half. Interestingly, the nuclei also show premature localization to the syncytial cortex, suggesting *gish* may be involved in the positioning of active myosin or in a mechanism that balances myosin-induced cytoplasmic flow. During the last four nuclear divisions on the cortex, *gish* depletion leads to defects in mitotic furrow formation, which further disrupts nuclear spacing and is accompanied by desynchronization of the nuclear cycle. In summary, we show that *gish* is required for the even positioning of nuclei in fly syncytium at multiple stages.

**670C** Why are axonal endoplasmic reticulum tubules so narrow? *Kishen Chahwala*, Cahir O’Kane University of Cambridge

Axonal endoplasmic reticulum tubules are much narrower in axons than in most other locations. But what does this mean for luminal protein diffusion? To test whether the small lumen diameter is limiting for protein diffusion, we compared the rate of diffusion of a GFP-tagged luminal protein through wildtype larval *Drosophila* axonal ER tubules, with diffusion through mutant axonal ER tubules with a larger diameter, in a triple mutant lacking the ER-shaping proteins, Rtn1, ReepA and ReepB. We performed FRAP experiments on axons to look at fluorescence recovery and thus the freedom of GFP to diffuse through the ER lumen.

We found that luminal diffusion of GFP was significantly faster in the larger mutant ER tubules. The time to half recovery in wider axonal ER tubules (lacking Rtn1, ReepA and ReepB) was approximately half as long as in wildtype.

The result implies that narrow axonal ER tubule diameter limits luminal protein diffusion. This leaves open the question of why it might be important to limit luminal ER diffusion along axons, for example, whether the diffusion of the much smaller (relative to GFP) Ca\(^{2+}\) ion is also constrained, and whether altered ER luminal diffusion might be a factor in the pathology of diseases like Hereditary Spastic Paraplegia that are caused by mutations in ER-shaping proteins.

**671A** EMC is required for biogenesis and membrane insertion of Xport-A, an essential chaperone of Rhodopsin-1 and the TRP channel *Catarina Gaspar*, *Ligia Vieira*, *Cristiana Santos*, *John Christianson*, *David Jakubec*, *Kvido Strisovsky*, *Colin Adrain*, *Pedro Domingos* 1) Instituto de Tecnologia Química e Biológica da Universidade Nova de Lisboa (ITQB-Nova); 2) Membrane Traffic Lab, Instituto Gulbenkian de Ciência (IGC); 3) University of Oxford; 4) Czech Academy of Sciences; 5) Queen’s University

The ER membrane complex (EMC) is required for the biogenesis of a subset of tail anchored (TA) and polytopic membrane proteins, including Rhodopsin-1 (Rh1) and the TRP channel. To understand the physiological implications of EMC-dependent membrane protein biogenesis, we performed a bioinformatic identification of *Drosophila* TA proteins. From 254 predicted TA proteins, screening in larval eye discs identified 2 proteins that require EMC for their biogenesis: fan and Xport-A. Fan is required for male fertility in *Drosophila* and we show that EMC is also required for this process. Xport-A is essential for the biogenesis of both Rh1 and TRP, raising the possibility that disruption of Rh1 and TRP biogenesis in EMC mutants is secondary to the Xport-A defect. We show that EMC is required for Xport-A TMD membrane insertion and that EMC-independent Xport-A mutants rescue Rh1 and TRP biogenesis in EMC mutants. Finally, ER resident, N196-glycosylated Rh1 TMD1-5 accumulates in Xport-A and EMC6 mutants, revealing a role for Xport-A in a glycosylation-dependent triage mechanism during Rh1 biogenesis in the endoplasmic reticulum.

**672B** Developing a *Drosophila* genetic screen for mutations that disrupt axonal ER organization *Nishani Jeyapalan*, Cahir O’Kane Genetics Department, University of Cambridge

Developing a *Drosophila* genetic screen for mutations that disrupt axonal ER organization  
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Axonal endoplasmic reticulum (ER) consists of a continuous tubular membrane network. Its continuity and potential as a channel for long-distance communication have earned it the description “a neuron within a neuron”. Mutations affecting ER-modeling proteins can cause the axon degeneration disease, hereditary spastic paraplegia (HSP), implying the importance of this network in axon maintenance. While the assembly, distribution, transport, and density of the network must be regulated to meet the needs of the neuron, we understand hardly anything of this regulation. To address this, our goal is to identify novel proteins involved in organizing and maintaining the ER network, using targeted CRISPR/CAS9 mutagenesis.

To allow easy visualization of axonal ER for screening, we identified *Drosophila* lines expressing GAL4 in small numbers of motor neurons innervating the adult leg, and used them to express fluorescent ER and plasma membrane markers, observed through undissected cuticle in adult legs. To allow tissue-specific CRISPR/CAS9 mutagenesis in the labelled motor neurons, we introduced a *UAS-CAS9* construct into these flies, targeting CAS9 expression only to the labelled
motor neurons utilising the *Drosophila* GAL4 system. Crossing this stock to any transgenic guide-RNA (gRNA)-expressing line allows tissue-specific somatic mutagenesis in the labelled neurons, and screening for axonal ER phenotypes as a consequence of tissue-specific gene disruption.

We have performed a pilot mutagenesis screen to observe axonal ER phenotypes in CAS9-mutated *Drosophila* motor neurons. We selected a small library of gRNAs, targeting a small selection of genes encoding ER-associated proteins, HSP protein orthologs, and motor proteins. We crossed these gRNA lines to the line expressing ER and PM markers and CAS9 in leg motor neurons, and tested the progeny for alterations in ER distribution in adult leg femoral motor axons.

Confocal screening showed a highly discontinuous axonal ER network as well as differences in ER and PM average intensities using several of these gRNA lines, in contrast to the more uniform continuous ER structure observed in control motor axons.

Having shown proof-of-principle for detecting axonal ER phenotypes caused by gRNAs in motor axon ER, we are now individually investigating gRNAs that cause axonal ER phenotypes via *in vivo* larval imaging, to ascertain whether axonal ER dynamics are also affected. Furthermore, we can now expand the pilot screen to encompass genes involved in several biological pathways, testing the roles of various gRNAs on axonal ER architecture.

**673C** A Dominant modifier Screen for Genetic Interactor of Jagunal in the *Drosophila* compound eye *Gerson Ascencio, Judy Abuel, Jorge Inojoza* San Francisco State University, San Francisco, CA

The endoplasmic reticulum (ER) is a continuous network of membrane tubules and flattened cisternae involved in protein and lipid synthesis, protein secretion, and post-translational protein modification. A recent study showed that the ER is partitioned asymmetrically in proneuronal cells during mitosis in early *Drosophila* embryos prior to cell fate determination. Furthermore, this asymmetric ER partitioning relies on the highly conserved ER transmembrane protein Jagunal (Jagn), however, the molecular pathway that drives Jagn-dependent ER partitioning is currently unknown. Here, we hypothesize that Jagn interacts with pathways involving cell fate selection to drive the generation of neuronal cell diversity. To identify possible genes that interact with Jagn, we performed a dominant modifier screen in the *Drosophila* compound eye. Expression of JagnRNAi in the *Drosophila* compound eyes results in a rough eye phenotype in 80% of the eyes examined. Based on this, we crossed a collection of deficiency lines (DF) covering the entire 3rd chromosome and examined for either enhancement or suppression of the rough eye phenotype. We have identified ten suppressors and twelve enhancers of Jagn-induced rough eye lines phenotype. We examined these DF lines and selected eight genes involved in functions such as organelle assembly, microtubule attachment, and organelle movements. We then performed a secondary dominant modifier screen in the *Drosophila* compound eye, using mutants of the selected target genes and mutant gene Dally gave us a similar rough eye phenotype percentage. Dally is important for neuron projection morphogenesis, and positive regulation of signal transduction. We hypothesize that Dally can be working with Jagn as a cell fate determinant. Targets will provide important insight into the molecular pathway involved in ER organization and partitioning.

**674A** Endosomal maturation in *Drosophila* nephrocytes depends on a trimeric Rab7 GEF complex *Maren Janz*, Lena Dehnens*, Jitender Kumar Verma*, Olympia Ekaterini Psathaki*, Lars Langemeyer*, Florian Fröhlich*, Jürgen J. Heinisch*, Heiko Meyer*, Christian Ungermann*, Achim Paululat† 1) University of Osnabrück, Department of Biology and Chemistry, Zoology and Developmental Biology; 2) University of Osnabrück, Department of Biology and Chemistry, Biochemistry; 3) University of Osnabrück, Center of Cellular Nanoanalytics, Integrated Bioimaging Facility Osnabrück (iBiOs); 4) University of Osnabrück, Department of Biology and Chemistry, Molecular Membrane Biology; 5) University of Osnabrück, Department of Biology and Chemistry, Genetics

*Drosophila* nephrocytes are specialized cells displaying highest endocytic activity. As a functional adaptation they display enormous cell membrane expansions and membrane tubulation where the endocytic active site is situated. The huge size of the cells also ensures enormous storage capacity, e.g. for degraded proteins or metabolic components. We use nephrocytes to investigate the trimeric Rab7 guanine nucleotide exchange factor (GEF) complex Mon1-Ccz1-Bulli. During endocytosis, Rab5 decorated early endosomes mature into Rab7 positive late endosomes. Subsequently, fusion with lysosomes leads to acidification and degradation of the lysosomal content/cargo. To function on membranes, Rab proteins are activated by GEFs and further inactivated by GTPase activating proteins (GAPs). Maturation of early endosomes to late endosomes and the fusion with lysosomes also depend on the two heterohexameric tethering complexes CORVET and HOPS. The CORVET complex is an effector of Rab5 whereas HOPS interacts with activated Rab7. In *Drosophila*, a trimeric Rab7 GEF complex, containing Mon1, Ccz1 and Bulli is active. We have shown that the absence of Bulli results in impaired endosomal maturation and enlarged Rab7 positive and negative endosomes with clustered Rab5 inside. Furthermore, the lack of either Mon1 or Ccz1 reveals defective Rab5 and Rab7 localization and internalization of Rab5 into endosomes. Bulli has been identified in insects and mammalian cells, indicating that the trimeric GEF is common in metazoan.
Overall, our data demonstrate the importance of the trimeric Rab7 GEF complex for proper localization of Rab5 and Rab7 in *Drosophila*. Our work aims to provide fundamental insight into the regulation of the endocytic pathway via this complex in nephrocytes. Especially the mechanism leading to Rab5 clustering in mutant flies will be investigated in future studies.


**675B  Peroxisome metabolism in enterocytes regulates the diet-gut-brain axis and lead to neurodegeneration** Francesca Di Cara, Stephanie Makdissi, Julia Kalinowski, Eden Bishop, Alex Liaukovich, Smita George

Peroxisomes are metabolic organelles that contribute to maintain the healthy metabolic status of the cell. Emerging evidence suggests that peroxisomes contribute in different ways to the function, development and survival of different tissues. Recent studies demonstrated that the peroxisome is essential in intestinal epithelial cells for the metabolic status of the organism, to regulate tissue inflammation and to maintain host-commensal/pathogen interactions in the gut. Considering the importance of gut-derived signaling for interorgans communication such as the gut-brain axis, we performed transcriptomic and metabolomic analyses to determine whether peroxisome metabolism in the gut controls secretion of trophic factors that contribute to neuronal health in adult flies. Integration of metabolomic and peptidomic analyses of the hemolymph integrated to transcriptomic and metabolomic analyses of control guts and peroxisome-depleted guts, revealed that peroxisomes modulate the production and secretion of gut-specific neuropeptides and inflammatory lipids in the circulation and in turn affect the gut-brain communications. Alteration of peroxisomal β-oxidation causes lipotoxicity in enterocytes triggers systemic release of inflammatory lipids (Cer22/Cer18) and inhibits the secretion of trophic neuropeptides such as Bursicon and Npl1 that are necessary to promote metabolic adaptation and stress responses in the organism. The transcriptional inhibition of these genes and the accumulation of inflammatory lipids together lead to brain inflammation, locomotor defects and reduce animal lifespan. Treatment with the free fatty acids scavenger, Niacin, rescues the described phenotypes in adult flies that have depleted peroxisomes in the enterocytes. Conversely, treatment with the established inhibitor of peroxisomal β-oxidation Thoridazine, or feeding an high fat diet lead to brain inflammation, dopaminergic neurons death, locomotion defects and reduces lifespan in control flies.

Considering that emerging evidence suggests that changes in diet-gut-brain axis influence the metabolic and inflammatory status of the neurons that is linked to Neurodegenerative Diseases (NDs) and that mild mutations in peroxisome genes are associated to the onset of NDs, we believe this work advanced our understanding of the the impact of peroxisomal metabolism on gut-brain axis, neuro-inflammation and the onset of NDs.

**676C A neuroprotective role of select peroxisome proteins at the fat body of Drosophila melanogaster** Kazuki Ueda1, Matthew Anderson-Baron1,2, Nathan Hoeven3, Julie Haskins1, Andrew Simmonds1 1) Department of Cell Biology, Faculty of Medicine and Dentistry, University of Alberta, Edmonton, AB, Canada T6G 2H7; 2) Future Fields Ltd. Edmonton, AB, Canada T6G 2H7; 3) Ross Shepard High School, Edmonton, AB, Canada T5M 2P2

A systemic lipid homeostasis is crucial for proper functioning and development of organ systems. Peroxisome is a membrane-bound organelle and is one of the central hubs of lipid metabolism. A congenital, autosomal recessive disorders affecting one of 14 core peroxisome genes, *Peroxins (Pex)*, leads to a spectrum of peroxisome biogenesis disorders (PBDs) exhibiting neurological deficits and developmental delays in children currently with palliative treatments as only available options. Peroxisomes are thought to associate and coordinate with lipid droplets (LDs) (cellular fat stores) to regulate β-oxidation of fatty acids and promote synthesis of plasmalogens and docohexaenoic acids. Our lab has recently identified peroxisome-independent functions of two Pex proteins (or Peroxins), Pex13 and Pex14, in promoting lipid storage at LDs in *Drosophila melanogaster*. In particular, we observed that Pex14 prevented localization of hormone sensitive lipase (Hsl), but not brummer (adipose triglyceride lipase homologue), to the surface of the LDs when co-overexpressed in *Drosophila* S2 cells under a condition that induces lipolysis. The Drosophila larval fat body (human adipose tissue and liver equivalent) stores lipids in large LDs and releases them during later development or during starvation conditions. We employed various *Pex* mutant and fat body-specific *Pex* knockdown (Pex FKD) flies to examine roles Peroxins have in lipid storage in vivo. When wild-type (WT) and Pex FKD (Pex1, Pex13, and Pex14) were fed on holidic food (low fat), they remained relatively stable and exhibited similar survival rates. When raised on holidic+lard food (high fat), the survival rates of both WT and Pex1 FKD larvae remained similar to the previous feeding condition while Pex13 and Pex14 FKD displayed a reduction in their survival rates suggesting lipotoxic effects due to an abnormal increase in systemic lipid levels. Compared to the WT larvae, Pex13, Pex14, and Pex16 FKD showed reduced brain size while Pex11 and Pex19 did not seem to be affected.
Compared to the WT larvae, only Pex13 and Pex14 mutants displayed reduced fat storage in the fat body indicated by reductions in size and quantity of LDs while Pex1 and Pex5 mutants seemed to display increased fat storage. Intriguingly, the dsRNA-mediated KD of Pex16 in S2 cells dramatically reduced Pex14 mRNA expression suggesting co-regulation of the two Pex genes. Overall, the evidence seems to support the unique roles of both Pex13 and Pex14 in promoting lipid storage in the fat body of Drosophila melanogaster. Moreover, these roles are independent of peroxisomes as other Pex FKD and mutants that also similarly affect peroxisome functions were seemingly unaffected.

677A  Identifying the minimal sequence that enables protein trafficking to the B-body, a novel nuclear domain Shania Kalladanthyil, Miranda Adams, Kaveh Kiani, Luis Rodriguez Kennesaw State University

Nuclear domains (ND) are the areas inside the cell nucleus with sharp boundaries and specific protein composition. Mechanisms that control the existence of various nuclear domains are not fully understood. The B-body is a prominent ND that was recently discovered in developing Drosophila flight muscles. We used B-body's resident protein Bruno 1 (Bru) as a model to dissect out the minimal protein sequence that is required to enable trafficking to this ND. Our study was facilitated by the relatively simple domain structure of Bru, which consists of one paired and one single RNA-recognizing motifs (i.e., RRM1/2 and RRM3), separated by two intrinsically disordered regions (IDR1 and IDR2). RNA-binding ability is critical for B-body affinity since ectopic RNAase treatment releases Bru from B-bodies in muscle cryosections. Using transgenic flies expressing GFP-tagged Bru mutants, we determined that B-body affinity depends on the functionality of RRM1/2, but not RRM3. Next, a series of truncation mutants were deployed to probe for the minimal sequence required for B-body accumulation. Neither the N-terminal IDR1 nor RRM1/2 alone nor in combination could accumulate in the B-body. However, RRM1/2 combined with the C-terminal IDR2 retained B-body affinity. Our study reaffirms the importance of domain combinations for the proper ND trafficking while highlighting the surprising selectivity of IDRs in this process. Future studies should determine if the affinity to B-body can be decoupled from the nuclear functions of the Bru protein.

678B  Characterization of the physical and functional connection between CNK and Misshapen Eloise Duramé1,2, Caroline Baril1,2, Malha Sahmi1,2, Marc Therrien1,2 1) Université de Montréal; 2) Institut de Recherche en Immunologie et Cancérologie (IRIC)

Connector eNhancer of KSR (CNK) is an evolutionarily conserved scaffold protein acting as a key regulator of the RAS-MAPK signaling pathway in Drosophila. To discover new functions of CNK, we captured the proximal interactome of CNK in Drosophila S2 cells using an improved version of the BioID technique called miniTurboID. This identified components of the RAS-MAPK pathway known to interact with CNK such as KSR, MEK, RAF and HYP. In addition, we identified the Ste20-related kinase Misshapen (MSN), which is involved in several biological events related to cell migration and planar cell polarity. To decipher the connection between CNK and MSN, we first tested whether CNK and MSN could physically interact in S2 cells. Co-immunoprecipitation experiments demonstrated their ability to associate in S2 cells. This interaction depends on the N-terminal part of CNK (containing the SAM, CRIC and PDZ domains) and the C-terminal part of MSN (containing the CNH domain). In another set of experiments, we obtained the common proximal interactors of CNK and MSN in S2 cells using miniTurboID. Strikingly, we identified 72 common interactors to CNK and MSN, which are largely associated to vesicular trafficking, cytoskeletal organization, cell polarity and migration. Finally, we used the UAS/GAL4 system to look for a genetic link between CNK and MSN. We found that knockdown of MSN during eye development had no effect on the level of phosphorylated MAPK and photoreceptor differentiation in third instar larva suggesting that the function of MSN is not related to RAS-MAPK signalling. Interestingly, we found that depletion of MSN during wing development produced wing blisters in ~40% adult Drosophila analyzed while depletion of CNK had no effect. However, co-depletion of MSN and CNK produced wing blisters in ~70% of adult flies, thus providing the first evidence of a genetic link between CNK and MSN. Blisters formation in adult Drosophila wing generally results from improper apposition/adhesion of the wing epithelial bilayer. Imbalance in integrin signalling, vesicular trafficking and cytoskeletal organization have been linked to blister formation. We are now investigating the role of CNK and MSN in those biological events. Our findings suggest that CNK is not exclusively devoted to RAS-MAPK signaling, but that it might work in other signaling events that play critical roles during vesicular trafficking, cytoskeletal organization and cell migration.

679C  Septins are necessary for detachment and protrusion formation in border cell migration Allison Gabbert, James Mondo, Joseph Campanale, Denise Montell UC Santa Barbara

Collective cell migration is crucial for development and the preferred mode of migration by metastatic tumors, but much about it is unknown. The border cell cluster in the Drosophila ovary is an ideal model for collective cell migration, as border cells migrate on and between nurse cells. The cytoskeleton is a critical regulator of cell migration. Septins, now considered the fourth cytoskeletal element, remain unexplored in collective systems. The objective of this study is to understand the functions and necessity of septins in border cell migration. Using RNAi, mutants, and over-expression, we investigated the impact of septins on border cell migration and cluster morphology through fixed and live imaging. We used high-resolution Airyscan imaging paired with Tissue Cartography to generate 3D models of the surface of the
cluster. We found that knocking down any of the five Drosophila septins significantly impacted border cell migration and cluster morphology through detachment failure, failure to form stable forward-directed protrusions, and a loose blebbly morphology. Overexpressing septins also dramatically impacted migration, with an inverse effect on morphology. As septin subunits function by forming higher order structures with each other, we investigated if septin monomers interacted. Clonal knockdown of Septin 1 (Sep1) or Septin 2 (Sep2) led to a significant loss of Peanut (Pnut, or Sep3) in the follicle and border cells. Similarly, knockdown of Sep1 or Pnut caused a loss of Sep2. These findings suggest that Sep1, Sep2, and Pnut form structures with each other in the follicle and border cells independently of Sep4 and Sep5, which have no impact on Pnut or Sep2 expression. To uncover the mechanistic role of septins in border cell migration, we tested candidates that may interact with septins. We observed co-localization between septins and nonmuscle myosin II in fixed imaging, and then explored further through live imaging. Amazingly, dynamic myosin flashes completely co-localized with dynamic septin expression. This suggests an interaction between septins and myosin. For example, septins may act as a scaffold for myosin, recruiting it and stabilizing protrusions. In conclusion, we found that septins are necessary for the detachment of the border cell cluster and for protrusion formation, while too much septin induces an excess of curvature and prevents migration. Sep1, Sep2, and Pnut interact to form structures and may interact with myosin.

**680A  Nuclear lamins promote collective cell migration and coordinate protrusion dynamics** Lauren Penfield, Denise Montell University of California, Santa Barbara

Cells migrate collectively during embryonic development and cancer metastasis. Determining how groups of cells detect and respond to confined tissue environments *in vivo* is a major goal. The nucleus, which is generally the stiffest organelle, deforms and sometimes ruptures to move through confined environments. While *in vitro*, the stiffness of the nucleus impedes movement of single cells through confined spaces, the effects of nuclear stiffness *in vivo* are unknown. Further, the roles of nuclei in collective cell migration are not clear. Here, we use border cell migration in the fly ovary as an *in vivo* model to investigate the effects of confined migration on nuclei. We found severe yet transient nuclear deformations occur, particularly in the leading cell, as border cells squeeze through tiny crevices between germline cells, termed nurse cells. These spaces are narrower than even a single border cell nucleus. Leading cells extend protrusions between nurse cells, which may pry open spaces to allow the cluster to traverse. The nucleus in the leading cell deforms as it moves through the neck of the protrusion and restores a more circular shape as the protrusion widens. In contrast, nuclei in the following cells had less dynamic movement and shape changes. These data suggest that nuclei in leading cells may widen protrusions to expand the size of the migration path and allow the cluster to move forward. To experimentally test how nuclei contribute to border cell migration, we investigated nuclear lamins, proteins that assemble into intermediate filaments to structurally support the nucleus. Depletion of the Drosophila B-type lamin, Lam, from the outer motile border cells, but not the inner, nonmotile polar cells, impeded border cell migration. Surprisingly, perturbations of the Drosophila A-type lamin, LamC, did not impair migration. While wildtype border cell clusters normally have one leading protrusion as they migrate, border cells depleted of B-type lamin had multiple, short-lived protrusions, resulting in unproductive cluster movement and failure to progress down the migration path. Further, border cell nuclei depleted of B-type lamins were smaller, formed blebs, and underwent rupture. Together, these data indicate that B-type lamins maintain nuclear structure, and this is required to stabilize the leading protrusion and promote collective cell migration through confined spaces.

**681B  Investigating the initiation of collective cell migration in the Drosophila follicular epithelium** Sierra Schwabach, Sally Horne-Badovinac University of Chicago

The collective migration of cells is a critical process during development, wound healing, and cancer metastasis. We study the highly coordinated collective migration of follicular epithelial cells in the Drosophila egg chamber, which helps to create the elliptical shape of the mature egg. During the formation of a new egg chamber, follicle cells initiate a collective migration perpendicular to the anterior-posterior (AP) axis. In many models of collective migration, a subset of the cells contact free space which can act as an external polarizing cue for the cells to start migrating. In contrast, the follicle cells form a topologically closed epithelium that lacks a free edge. As a result, the direction of follicle cell migration in each egg chamber is stochastic and can occur either clockwise or counterclockwise with respect to the AP axis. Follicle cell migration depends on a type of planar polarity, in which the atypical cadherin Fat2 localizes to the trailing edge of each cell and signals to the cell behind to promote migration. Interestingly, we know that the maintenance of Fat2’s planar polarized localization also requires migration of the tissue, which is driven by SCAR/WAVE-dependent protrusions at the leading edge of each follicle cell. This leads to our model that polarized Fat2 signaling and migration promote one another in a self-reinforcing, mutually dependent feedback loop. While we have a grasp on the mechanisms that promote follicle cell migration once it has started, it is unknown how follicle cells initiate collective migration without the existence of an external polarizing cue. We hypothesize that it is the interplay between motility and polarized Fat2 signaling that allows the tissue to self-organize and initiate collective migration. We have taken two different approaches to address this hypothesis. The first includes improving upon current live imaging methods to watch migration initiation unfold in its natural context. The second approach utilizes the GAL4/UAS system to delay migration.
The basement membrane is a sheet-like extracellular matrix that underlies epithelia and surrounds muscles. In the
initiation until the egg chambers are older and easier to image. Preliminary results from both methods will be discussed
at the poster.

682C Control of Crag’s localization and activity in the polarized deposition of basement membrane proteins in
epithelial cells. Hemin Shah¹, Alex Hoover¹, Megan Gladwin², Trudi Schüpbach², Olivier Devergne¹ 1) Department of
Biological Sciences, Northern Illinois University, DeKalb, IL; 2) Department of Molecular Biology, Princeton University,
Princeton, NJ

Epithelial cells play critical roles in the development and maintenance of an organism, and the establishment and
maintenance of apical-basolateral polarity (ABP) are essential to their function and integrity. ABP is established,
maintained, and tightly regulated via intracellular trafficking and environmental cues, such as those provided by the
basement membrane (BM). The BM is a specialized sheet of extracellular matrix accumulating and underlying epithelial
cells on their basal side. Despite the important roles of the BM in the architecture and functions of epithelial cells, little is
known about the mechanism ensuring the exclusive basal restriction of the BM components. To study BM deposition, we use the follicular epithelium (FE) of the Drosophila ovary as our model system. The GEF/RabGTPase complex Crag/Rab10 is a key regulator of the biological pathway specifically dedicated to the basal restriction of BM components. However, the exact mechanism responsible for Crag’s polarized intracellular localization and its activity in BM polarity remains yet to be elucidated. In FE, it assumes a polarized localization and accumulates apically and laterally through yet unknown mechanisms. Importantly, specific localization of the GEF Crag is thought to be critical for the localized activation of Rab10. Thus, to understand how Crag controls the polarized deposition of BM proteins, it is critical to determine how Crag’s activity and localization are controlled. Crag is a multidomain protein containing the DENN domains at the N-terminus responsible for its GEF activity, a Calmodulin Binding Site (CBS) domain responsible for its calmodulin-binding activity, and a conserved C-terminus domain. To determine the domain(s) necessary for Crag’s localization and activity, we performed a structure-function analysis. Our data suggest that the CBS domain, but not the DENN domains, is important for the localization of Crag to the apical and lateral domains, suggesting a role of calmodulin in the subcellular localization of Crag. The DENN domains, however, are required, but not sufficient, to control the basal restriction of the BM components to the basal side of the epithelial cells, suggesting that the proper localization of Crag is required for its control of the polarized secretion of basement membrane proteins. Altogether, our data shed a light on the regulation of the activity and localization of Crag, a key component of the biological pathway that controls BM polarity.

683A Basement membrane repair dynamics in the Drosophila midgut Aubrie Stricker, Kimberly LaFever, Katherine
Peebles, M. Shane Hutson, Andrea Page-McCaw Vanderbilt University, Nashville, TN

Basement membranes are the oldest, most conserved forms of extracellular matrix and serve to separate tissue layers,
provide mechanical support, direct signals to neighboring cells, and insulate tissues from signals. Further, basement
membranes are subject to mechanical damage and require dynamic repair mechanisms. Faulty basement membrane
repair mechanisms can aid in the progression of diseases such as asthma and diabetes, and diseases of the basement
membrane itself, including Alport’s syndrome and Goodpasture’s syndrome. Therefore, understanding how basement
membranes repair will be vital to treating these conditions. Our work utilizes the Drosophila midgut basement
membrane to probe repair dynamics. In Drosophila, all major basement membrane components have been conserved
but with less redundancy than mammals. Our lab has developed an assay to reproducibly damage the basement
membrane and study the repair process. Previously we reported that many aspects of basement membrane repair are
shared during homeostasis. Thus, it is unclear whether basement membrane damage is actively detected, or instead,
passively repaired by homeostatic mechanisms.

Our recent data suggests basement membrane damage is actively detected. Following damage, there is an increase in
the number of enteroendocrine (EE) cells, the major secretory cell type in the gut, and we find that guts without EE cells
cannot repair the basement membrane. Additionally, the EE cells are a significant contributor of collagen IV needed to
repair damage. Importantly, EE precursor cells express a mechanosensory stretch-activated ion channel, Piezo, raising
the possibility that a change in stiffness of damaged basement membranes signals the initiation of repair. Excitingly, Piezo
knockout flies are able to maintain basement membrane homeostasis in the adult fly but cannot repair it after damage.
This is our first evidence that there is a unique mechanism to detect basement membrane damage and initiate repair.
However, to our surprise, Piezo knockout flies also showed an increase in EE cells following damage, suggesting that EE
cells are regulated independently of Piezo. This result suggests that Piezo and EE cells function independently to repair
basement membrane, raising the question of how Piezo functions in basement membrane repair.

684B The mystery of the Peroxidasin mutant: why does this catalytically dead Drosophila mutant survive?
Katherine Peebles¹, Kimberly LaFever¹, Gautam Bhave¹,², Andrea Page-McCaw¹ 1) Vanderbilt University; 2) Vanderbilt
University Medical Center

The basement membrane is a sheet-like extracellular matrix that underlies epithelia and surrounds muscles. In the
gut of *Drosophila*, the stiff basement membrane surrounds the muscles used in peristalsis to keep them flat and smooth. Collagen IV is one of the main components of the basement membrane, where it adds structure and stiffness. The *Peroxidasin (Pxn)* gene encodes an enzyme that crosslinks collagen IV at the NC1 domain. This crosslinking supports basement membrane stability and contributes to its stiffness. Hypomorphic P-element mutations of *Pxn* exist in *Drosophila* that survive to the larval stages with decreased viability to adulthood, but no null mutation has been reported. We expected a null mutation to be lethal at the end of embryogenesis, when collagen IV mutants die. Using CRISPR, we created a mutant (*Pxn<sup>11</sup>*) that deletes a portion of the catalytic domain, eliminating its activity. Further, this deletion also caused a frameshift mutation that inserted a stop codon soon after the deletion. Contrary to our expectations, about twenty percent of expected homozygotes survive and live an apparently normal lifespan. These homozygotes exhibit muscle defects in the gut consistent with loss of stiffness in the basement membrane. A few hypotheses that could account for their viability will be discussed.

**685C The role of ZP domain proteins in controlling corneal lens architecture** Neha Ghosh, Hongsu Wang, Jessica Treisman Skirball Institute of Biomolecular Medicine, NYU School of Medicine

The shape of organs can be critical to their function; for example, changes in the curvature of the human cornea result in visual disorders such as myopia, hypermetropia, astigmatism or keratoconus. A simpler model system like *Drosophila* may provide insights into the development of curved refractive structures such as the cornea. Each ommatidium of the *Drosophila* eye contains a biconvex corneal lens that focuses light on the underlying photoreceptor cells. The central part of the corneal lens is thought to be secreted by the underlying cone and primary pigment cells, while the peripheral portion is attached to the secondary and tertiary pigment cells. Here we show that loss of the zinc-finger containing transcription factor Blimp-1 specifically from the secondary and tertiary pigment cells causes flattening of the external corneal lens surface, dramatically reducing its refractive power. One possible mechanism for this effect of Blimp-1 would be a change in the pattern of attachments of the corneal lens to the peripheral pigment cells. The fly corneal lens is a cuticular structure containing the polysaccharide chitin, and the morphology of other chitinous structures such as denticles and tracheal tubes has been shown to depend on the localization of specific zona pellucida (ZP) domain containing proteins, which attach the apical extracellular matrix to the plasma membrane. In a transcriptomic study of the effects of *Blimp-1* knockdown in the mid-pupal retina, we observed changes in the expression levels of several genes encoding ZP domain proteins; *dusky-like (dyl)*, *CG10005* and *cypher (cyr)* were downregulated, while *neyo*, *piopio* and *quasimodo* were upregulated. We find that loss of *dyl* or *cyr* also causes flattening of the outer surface of the corneal lens. These observations suggest that Blimp-1 may influence corneal lens architecture in part by ensuring the appropriate distribution of the ZP proteins that attach its periphery to the secondary and tertiary pigment cells.

**686A Neural IgCAMs at work in epithelia: phylogeny and function** Colleen Maillee, Dan Bergstralh University of Rochester

Epithelial cells are linked together by adherens junctions, which are largely responsible for cell-cell adhesion, and occluding junctions, which form a barrier to paracellular diffusion. Our published results show that another epithelial adhesion module, comprised of Ig-domain adhesion proteins – Neuroglian/L1CAM, Fasciclin II/NCAM, and a three Ig-domain protein, either Fas3 (insects) or potentially CADM1 (vertebrates) – helps to maintain epithelial integrity during proliferation by ensuring that the tissue grows as a monolayer. These factors are not only found in proliferating epithelia, where they can localize along the length of the cell-cell border, but also in the vertebrate excitatory synapse and *Drosophila* neuromuscular junction. Furthermore, they have been extensively studied in both flies and mammals for their role in axon guidance. We want to know how they work. L1CAM is evolutionarily conserved at the sequence level, and the function of its cytoplasmic domain is well studied. Remarkably, whereas the extracellular C-terminal structures for NCam/Fas2 and CADM1/Fas3 are similar, their cytoplasmic N-termini are not. These findings suggest conservation of function without conservation of sequence. We are using phylogenetic analysis, structural prediction algorithms, and *Drosophila* genetics to study this problem.

**687B Fatty acid trafficking during Drosophila oogenesis** Roger White, Michael Welte University of Rochester, Biology department, Rochester, NY

Lipid droplets (LDs) are ubiquitous fat storage organelles with essential roles in lipid metabolism, including storage and trafficking of fatty acids. During Stages 9-10B of oogenesis, *Drosophila* egg chambers accumulate hundreds of thousands of LDs containing triacylglycerol (TAG) and sterol esters. Most of the fatty acids (FAs) in these stored lipids ultimately come from lipophorin (LPP) particles circulating in the hemolymph. Previous work has led to the following working model for how these FAs are funneled into LDs: LPP particles dock on the nurse cell plasma membrane via lipophorin receptors and are broken down by extracellular lipases. The liberated FAs are then taken up via FA transporters, activated by acyl-CoA synthetases and converted to TAG by DGAT1/Midway. The long-term goal of this project is to critically test this model and identify the molecular players and regulatory steps involved. To this end, we have developed a protocol to monitor FA trafficking during oogenesis using fluorescently labeled fatty acid (FLFA). When flies are fed food
supplemented with various FLFAs, FLFA accumulates in nurse cell LDs. Preliminary evidence suggests that when flies are starved prior to FLFA feeding FLFAs do not enrich in LDs, but appear to accumulate in mitochondria, suggesting that egg chambers direct FLFA to different intracellular locations according to the physiological state of the animal. When isolated egg chambers are incubated in FLFA containing media, we observe incorporation in LDs already after 15 mins, providing an inroad for observing FA trafficking to nascent LDs. In mutants for midway, no LDs form and egg chambers arrest at Stage 9. Using FLFA supplementation, we found that in the mutant egg chambers FLFA is still taken up, but accumulates in mitochondria, even for well-fed flies. In addition, nurse cell mitochondria take on an abnormal morphology. We hypothesize that when incoming FAs are not properly sequestered in LDs, they inappropriately accumulate in mitochondria, causing dysfunction. Intriguingly, reducing the dosage of Eip75B, a Drosophila PPAR homolog, allows midway mutant egg chambers to develop further, implicating dysregulated lipid signaling in the developmental arrest. In summary, our results suggest that during oogenesis LDs are critical for regulating FA trafficking and usage and are key for mitigating lipotoxicity.

688V  Rap1 acts via Canoe and Rho1 to control the adhesion and cytoskeletal rearrangements that drive rapid wound repair Katheryn Rothenberg, Rodrigo Fernandez-Gonzalez University of Toronto

Coordinated cell movements contribute to tissue development and repair and to the spread of metastatic disease. We investigate collective cell migration during wound healing in the Drosophila embryonic epidermis, which allows for genetic, pharmacological, and biophysical manipulations. Upon wounding, the cells immediately adjacent to the wound become polarized: cell-cell adhesion molecules are internalized from the wound edge, and actin and the molecular motor non-muscle myosin II accumulate at the interface with the wounded cells, forming a supracellular cable around the wound that coordinates cell movements. The cable is thought to assemble from linking anchor to cell at former tricellular junctions (TCJs) along the wound edge, which are reinforced during wound closure through accumulation of cell-cell adhesion components. However, the mechanisms that coordinate the adhesive and cytoskeletal rearrangements required for rapid wound closure are unclear. The small GTPase Rap1 is a mechanosensitive molecular switch that promotes cytoskeletal polarization and regulates cell adhesion turnover in other systems. Using quantitative time-lapse microscopy of fluorescently tagged proteins, we found that Rap1, the Rap1 effector Canoe/Afadin, and E-cadherin were simultaneously depleted from the wound edge and accumulated at TCJs. Reducing Rap1 activity by overexpressing a dominant-negative Rap1 (Rap1DN) slowed wound repair by 45%. The slower wound closure was accompanied by defective actomyosin polarization to the wound edge and a 25% loss in E-cadherin accumulation at TCJs compared to a 25% increase in controls. Together, our data indicate that Rap1 is necessary for rearranging cell-cell adhesions and creating a stable force-generating actomyosin cable to drive rapid wound closure. Consistent with this model, Canoe knock-down led to 44% slower wound repair and 32% lower E-cadherin fluorescence at TCJs. These results suggest that Rap1 signals through Canoe to reinforce TCJs during wound closure. To understand how Rap1 may be affecting assembly of the actomyosin cable, we measured Rho1 activity in Rap1DN embryos and found that Rho1 activity was reduced by 72% in Rap1DN embryos, which is accompanied by a 54% reduction in tension at the wound edge. However, Canoe knock-down does not affect Rho1 activity. Our data support a model in which Rap1 simultaneously drives actomyosin cable assembly via activation of Rho1 and reinforced adhesion at TCJs via Canoe/Afadin.

689V  Cell wound repair requires the coordinated action of linear and branched actin nucleation factors Justin Hui, Mitsutoshi Nakamura, Susan Parkhurst Basic Sciences Division, Fred Hutchinson Cancer Research Center, Seattle, WA

During cellular wound repair, actin and other cytoskeletal proteins act in concert to coordinate wound closure in a timely fashion. A critical step is to form an actomyosin ring that contracts to close the wound. This process is dynamically regulated in part by Rho family GTPases, which through their different effectors can regulate both linear and branched actin nucleators to facilitate cytoskeletal dynamics. We have previously shown that knockdown of the formin Diaphanosus, a linear actin nucleator and downstream effector of Rho, results in prolonged wound closure and the inability to form a proper actomyosin ring. However, when all linear nucleation factors are removed, some actin organization at the wound persists. We investigated the contribution of branched nucleation factors to cell wound repair. We show that the Wiskott-Aldrich Syndrome (WAS) protein family members (WASp, Wash, and SCAR), and their cofactor Arp2/3, play non-redundant roles in regulating actin organization and architecture of the contractile actomyosin ring. Notably, we observe stark differences in the spatial and temporal recruitment patterns to wounds among the WAS proteins during the wound repair process. In particular, WASp is recruited early, SCAR recruitment peaks late, and Wash is present throughout the process. Further, individual knockdown of the WAS proteins resulted in different actomyosin ring architectures, including differences in mesh density and filament orientation. Dynamically, WASp knockdown exhibits slower contraction whereas SCAR contracts faster and Wash contracts similarly to wildtype. Interestingly, in the absence of branched actin, wounds exhibited an abundance of elongated linear filaments that colocalize with Diaphanosus, as well as other unique actin structures, which colocalize with Myosin. When both linear and branched nucleation factors are simultaneously knocked down, we observe a significant decrease of these elongated filaments and the absence of previously seen actin structures. We also inhibited myosin activity in an Arp3 knockdown and observed a surprising wound repair phenotype consisting of spiraling linear filaments and excessive actin bundling. Our results

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emphasize the strong requirement for balance and crosstalk among linear and branched actin nucleators, as well as myosin, to facilitate proper actin filament architecture, organization, and dynamic contractile closure of the actomyosin ring during cell wound repair.

690V Molecular regulation of centrosome stability Ana Pimenta-Marques, Tânia Perestrello, Patrícia Rodrigues, Paulo Duarte, Mariana Faría, Mónica Bettencourt-Dias Instituto Gulbenkian de Ciência

An important feature for cell homeostasis is how structures such as different organelles are maintained in the cell. This is particularly relevant for organelles whose number and function are under a tight control, as deregulation of these properties has critical implications for the cell. This is the case of the centrosome. The centrosome is the main microtubule-organizing center of eukaryotic cells. This organelle is composed of two centrioles, surrounded by a multiprotein matrix called the pericentriolar material (PCM) which is critical for microtubule nucleation.

Centrosomes are thought to be very stable and can persist over long periods of time. However, these structures can disappear in certain developmental stages and their numbers and structure can be deregulated in several diseases. Moreover, some centrosome components are quite dynamic. While a large body of knowledge has been produced regarding the biogenesis of these structures, little is known about how they are maintained. Our previous work showed that both POLO kinase and PCM are critical players in a centriole maintenance program which is shut off in the female germ-line to allow fertility. We have now investigated the players involved in this program, at the level of the centriole. By studying centrosome stability in the female germine and in Drosophila cultured cells, we found that several centriole components play a role, amongst them the centriolar wall protein ANA1 has a very strong phenotype associated with the centrosome maintenance program. We then further show that the stability conferred by Polo and the PCM to centrosomes is dependent on ANA1, and that ectopic targeting of ANA1 can prevent centrosome loss in oogenesis. Our work suggests that the PCM functions as a concentrator of components, in particular ANA1, which are required for centriole maintenance. We will discuss our data and how it provides a deeper understanding on the molecular mechanisms by which centriole stability is regulated.

691V Evolutionary Diversification of Drosophila Arp2 for Specialized Actin Branching Sarah Tomlin², Tristan Spain¹, Harmit Malik²,³, Courtney Schroeder⁠;¹ 1) UT Southwestern Medical Center; 2) Fred Hutchinson Cancer Research Center; 3) Howard Hughes Medical Institute

The actin cytoskeleton, which is composed of force-generating polymers, often forms branched networks that are critical in many fundamental cellular processes, including cell motility, cell division and vesicular movement. Branched actin networks are generated by the Arp2/3 complex, a 7-membered protein complex including actin-related proteins (Arps) 2 and 3. Similar to actin and most Arps, Arp2 is evolutionarily ancient among eukaryotes and under stringent sequence conservation, yet we surprisingly discovered two clade-specific gene duplications of Arp2 in Drosophila: Arp2D in the obscura clade and Arp2D2 in the montium clade. Our targeted sequencing and phylogenetic analyses of Arp2D and Arp2D2 show these duplicates have evolved independently and arose ~14 million years ago at the origin of their respective clades. The two duplicates exhibit distinct sequence diversification from canonical Arp2, and unlike the ubiquitously expressed Arp2, both duplicates are testis-enriched in expression. Why would evolution recurrently select for a divergent Arp2 for potential roles in the male germline? To elucidate the function of these duplicates, we investigated whether both duplicates can polymerize branched actin networks similar to canonical Arp2. We replaced canonical Arp2 in D. melanogaster with D. pseudoobscura Arp2D or D. auraria Arp2D2 and found they can both rescue the Arp2 knockout lethality phenotype, suggesting the duplicates can indeed polymerize branched actin networks despite their sequence divergence. Cytological analyses further confirmed the two duplicates localize to branched actin networks. Surprisingly, however, Arp2D-expressing flies have significantly reduced fertility unlike Arp2D2-expressing flies. Our findings suggest that canonical Arp2’s roles can largely be partitioned between somatic versus germline roles, and we hypothesize that while both Arp2 gene duplicates can polymerize branched actin in somatic cells, Arp2D fails to generate proper actin networks for germ cell development. We are currently investigating the integrity of germline actin in the transgenic flies and exploring how Arp2D’s sequence divergence has perhaps led to specialized actin polymerization for germline biology.

692V Centrosome-induced membrane infolding linked to Rac pathway and Arp2/3 network recruitment during actin cap formation in the Drosophila embryo Rebecca Tam, Nicolas Vergara Ruiz, Tony Harris University of Toronto

The cell cortex is a dynamic network of proteins that allows cells to change shape during processes like migration and division. Local activation of the Rac-GEF pathway leads to the assembly of Arp2/3 networks which change the cell cortex. Moreover, centrosomes influence the cortex during cell migration and division. How centrosomes activate the assembly of actin networks is not well understood. During early Drosophila embryogenesis, an actin cap forms above the nucleus and the growth of the cap creates a dome-like structure that houses the mitotic apparatus. The actin cap forms below the plasma membrane and requires centrosomes and the Rac-GEF, Sponge. What remains unclear is how centrosomes promote the local recruitment of the Rac-GEF pathway. My current research shows that the plasma
Membrane trafficking describes the vesicular transport of proteins and macromolecules by cells. Trafficking defects impair signaling events and have dramatic consequences on cell homeostasis. Enterocytes (ECs), intestinal differentiated cells, are the main cell type of the gut epithelium. Given their absorptive functions, they display a high membrane trafficking flux. Notably, defects in endocytic pathways can affect EC function and lead to intestinal bowel diseases. However, how does trafficking regulate intestinal tissue maintenance is poorly understood. Rab21 is a small GTPase involved in early endosomal trafficking. It was first identified in human intestinal cells and is expressed in ECs. Importantly, Rab21 levels are drastically decreased in ECs upon intestinal inflammation.

Using the Drosophila intestine as an in vivo model system, we investigated EC-specific functions for Rab21 in gut tissue maintenance. We monitored and assessed Rab21 loss of function in ECs of adult flies. Rab21 depleted guts showed severe intestinal morphology abnormalities. Furthermore, cellular equilibrium was affected, and a gain in mitotic cells, associated with increased cell death was observed. Interestingly, we found that an increased level of the IL-6 like cytokine, Upd3, in ECs was responsible for overproliferation, through activation of the JAK/STAT pathway mediated via yki activity and apoptosis.

To further characterize Rab21 functions, we performed a TMT-based quantitative proteomic analysis for Rab21 depleted guts and identified 101 proteins significantly modulated. We investigated the functional link between Rab21 and two overabundant proteins: ApoLpp, an apolipoprotein involved in diacylglycerol mobilization and Tret1-1, a trehalose transporter. We observed that Rab21 silencing in EC affects lipid content in the gut, as well as in the fat body, while circulating lipids did not appear affected, highlighting an unappreciated lipid-related function for Rab21. Furthermore, we demonstrated that loss of Rab21 in EC impaired trehalose levels in hemolymph, suggesting the involvement of Rab21 in the regulation of trehalose distribution, likely via specific trafficking of the trehalose transporter. Altogether, our data indicate that EC Rab21 plays an important role in intestinal tissue maintenance and functions. Although Rab21 specific roles on lipids and trehalose regulation remain to be defined, we show that Rab21 contributes to ECs survival and their correct function.

Macrouautophagy (autophagy) is a trafficking pathway that relies on delivery of membrane-bound cytoplasmic contents for degradation by the lysosome, referred to as the autolysosome. The lysosome is a very important component in autophagy regulation as it must be able to accommodate the convergence of cargos from both autophagy and endocytosis pathways, as well as coordinate autolysosome maturation with changing levels of basal versus stress-induced autophagy. In order for to replenish the lysosomes during prolonged starvation-induced autophagy, one-way new lysosomes can be made is through the autophagic lysosome reformation (ALR) pathway. Tubule-like projections from lysosomes pinch-off to form proto-lysosomes, eventually maturing into new functional lysosomes. We identified two phosphoinositide lipid regulators, class II PI3-kinase (PI3KC2) and Myotubularin PI3-phosphatase (Mtm), that act in a shared autolysosome inhibitory pathway to maintain basal levels of autophagy in the fat body of fed Drosophila larvae. Strikingly, we discovered that wildtype flies also co-express a truncated, noncatalytic PI3KC2 splice variant (PI3KC2-short) that, converse to PI3KC2 and Mtm, is required for autolysosome maturation through repression of their catalytic activities. We have shown that defects in autolysosome maturation correlate with aspects of (ALR). Our work addresses the importance of coordinated activity between specific phosphoinositide kinase and phosphatase activities. In addition, this study reveals a novel role for their joint regulation by a noncatalytic splice variant to respond to changing homeostatic demands in membrane flux.
Stem cell differentiation is essential for development and tissue homeostasis. Often differentiation is accompanied by a rewiring of organelles and alterations of the metabolome. However, how mitochondrial dynamics contributes to, and influences differentiation in vivo remains poorly understood. Using the Drosophila germline as a model system, we find that remodelling of the inner mitochondrial membrane is coupled to germline stem cell (GSC) differentiation. During the early stages of differentiation, mitochondrial Complex V is essential to drive the remodelling of cristae and in its absence the immediate progeny fail to survive. The failure to remodel mitochondria during early germ cell differentiation results in activation the Integrated Stress Response through the ER membrane kinase, PERK. PERK does so by detecting alterations in the ER membrane. Furthermore, the loss of Complex V and the failure to remodel mitochondria, induces the formation of lipid droplets that is partially mediated through PERK activity. Taken together, our data suggest that the loss of inner mitochondrial remodelling during early germline stem cell differentiation destabilizes the ER membrane thereby activating PERK and inducing cell death.

697V Cell-extracellular matrix adhesion is necessary for rapid embryonic wound closure Michelle Ly1,2,3, Katheryn Rothenberg1,2, Clara Schimmer1,2, Raymond Hawkins1,2, Rodrigo Fernandez-Gonzalez1,2,3,4,5 1) Institute of Biomedical Engineering, University of Toronto, Canada; 2) Translational Biology and Engineering Program, Ted Rogers Centre for Heart Research, University of Toronto, Canada; 3) Collaborative Specialization in Developmental Biology, University of Toronto, Canada; 4) Department of Cell and Systems Biology, University of Toronto, Canada; 5) Developmental and Stem Cell Biology Program, The Hospital for Sick Children, Canada

Embryos repair wounds rapidly, with no inflammation or scarring. In embryos, wound repair is driven by the collective movement of the cells around the wound to seal the lesion. Embryonic wound healing is associated with reorganization of the actomyosin cytoskeleton and cell-cell adhesive structures in the cells adjacent to the wound. Cell migration often involves adhesion to and remodeling of the extracellular matrix (ECM) through integrin-based adhesions. However, the role of integrin-based adhesion in embryonic wound closure has not been investigated. To establish the role of cell-ECM adhesion in embryonic wound repair, we imaged and wounded the epidermis of embryos mid-way through development (12-14 hours) using a pulsed ultraviolet laser mounted on a spinning disk confocal microscope. Using embryos expressing fluorescently labeled talin, we found that upon wounding, the cells adjacent to the wound polarized talin to the wound edge, suggesting increased cell-ECM interactions at the wound margin. To determine the impact of cell-ECM adhesion on embryonic wound closure, we injected embryos with RGDS, a peptide that binds integrins and inhibits ECM binding. RGDS treatment delayed wound closure by 43% with respect to controls. Imaging of fluorescently labeled myosin II revealed that the delay in wound healing occurred with no significant effects on the myosin rearrangements associated with wound repair. We obtained similar results when we knocked down integrins using RNA interference against the β integrin subunit: wound closure displayed a 54% delay compared to controls with no changes in myosin dynamics around the wound. However, we found a 71% decrease in actin accumulation around the wound edge relative to controls. We also noticed a defect in the E-cadherin reorganization necessary for actomyosin remodelling during wound healing. In control embryos, E-cadherin accumulated at former tricellular junctions at the wound margin, displaying a 29% increase in tricellular junction E-cadherin fluorescence 15 minutes after wounding. In contrast, RGDS treatment resulted in a 13% decrease in tricellular junction E-cadherin at the same time point. We obtained similar results in integrin RNAi embryos, which displayed a 20% decrease in tricellular junction E-cadherin. Together, our results suggest that cell-ECM adhesion contributes to wound repair and reveal a previously unrecognized interplay between cell-cell and cell-ECM adhesion in the collective cell movements that drive embryonic wound closure.

698V Cell extrusion during starvation-induced intestinal shrinkage Aparna Sherlekar Banerjee, Elsa Su, Samantha Thomas, Keerthana Yellapragada, Lucy Erin O’Brien Stanford University School of Medicine

The adult Drosophila midgut shrinks as a result of nutrient deprivation. Shrinkage is in part due to reduction in enterocyte cell volume and cell loss by enterocyte cell extrusion - there was two-fold increase in extrusions upon fasting. At steady-state most cells activated executioner caspases before they extruded. Inhibition of caspase activation blocked extrusions at steady-state. However, in fasted guts, ~70% of extruded cells were negative for cleaved caspase. TOR signaling is a conserved pathway that senses and responds to nutrient signals. Paradoxically, TOR signaling activation by inhibiting Tsc2 or Rheb overexpression increased cell extrusions in the midgut. Tsc2 inhibition showed dramatic resistance to fasting compared to control flies, while flies with blocked caspase activation had significantly shorter lifespans. Further studies can help understand the role of TOR signaling in mediating extrusions to achieve organ shrinkage in the adult Drosophila midgut.

699V Basal intercellular junctions integrate local cytoskeletal forces to regulate Hippo signalling in growing epithelia Benjamin Kroeger1, Samuel Manning1, Yoshana Fonseka2, Kieran Harvey1,2 1) Monash Biomedical Discovery Institute, Monash University, Melbourne, Australia; 2) Peter MacCallum Cancer Centre, Melbourne, Australia

The Hippo signalling pathway is an evolutionarily conserved regulator of organ growth and homeostasis. By the assembly of pathway components in discrete subcellular domains, Hippo signalling can be either amplified or attenuated. The main subdomains previously associated with Hippo signalling in fly tissues are found at the apical portion of epithelial cells,
and include the adherens junctions and apical membranes. Hippo signalling can be modulated by cytoskeletal tension, which exerts forces at the adherens junctions and influences the subcellular localisation of the key Hippo pathway kinase, Warts.

Using a combination of molecular genetic approaches and high-resolution imaging, we discovered a novel subcellular domain where Warts is enriched: punctate, basal intercellular junctions. Using APEX2 transmission electron microscopy we found that in addition to its known enrichment at apical junctions, Warts accumulates in focal intercellular junctions at the basal-most region of epithelial tissues. Our genetic analyses show that the basal junctions are similar in composition to the belt-like adherens junctions. We reveal that Warts is dynamically recruited by LIM-family proteins to punctate basal junctions in response to local cytoskeletal forces, driven by myosin activity. By recruiting Warts basally, it is spatially restricted from activators of Hippo pathway signalling, which are enriched at the apical portion of epithelial cells. Our study sheds light on how the Hippo pathway integrates mechanical signals, and highlights previously overlooked aspects of cell biology and tissue dynamics that operate at the basal region of growing epithelial tissues.

700C Rab1 suggests a role for ER regulation in chromosomal separation during mitosis Katie Rollins, J. Todd Blankenship University of Denver

Mitotic spindles are often observed to be found in close association with the endoplasmic reticulum – however, the purpose of this association is less clear. The syncytial stage of fly embryogenesis is a period of rapid mitotic divisions, with the blastoderm stage marked by four rounds of divisions in a membrane-less syncytium near the cell cortex. These rapid divisions demonstrate major remodeling events of various membranous organelles, such as the nuclear envelope, the endoplasmic reticulum, and the deformation of the plasma membrane into cytokinetic-like furrows. Work from our lab, as well as others, has shown the contribution of multiple Rab-directed pathways to these processes, but relatively little work has examined how transformations in the ER contribute to successful mitoses. Here, we demonstrate that Rab1 controls ER to Golgi trafficking and is highly active within the Drosophila syncytium. Interestingly disrupting Rab1 results in severe mitotic failures, suggesting a role for the ER in permitting appropriate spindle function. In Rab1 knockdown embryos the endoplasmic reticulum accumulates in large spindle associated aggregates. This aggregation appears to be dynein-dependent, and disrupting the aggregation rescues mitotic function. We will also report on the mechanisms by which the over-accumulated ER affects the integrity of the spindle.

700A Understanding the role of Matrimony in suppressing the drive of the B chromosomes Kaylah Samuelson, Stacey Hanlon Department of Molecular and Cell Biology, University of Connecticut, Storrs, CT

The genome is constantly under attack by selfish genetic elements that bias their transmission to the next generation. A classic type of selfish element B chromosomes - supernumerary, nonessential chromosomes that can be detrimental to their host- were recently discovered in a single laboratory stock of D. melanogaster that carries a null mutation in matrimony (mtrim^{TM3, Sb Ser}). This combination allows the B chromosomes to be transmitted through female meiosis at a higher-than-expected frequency, a phenomenon referred to as meiotic drive. Though it is clear that having only one functional copy of mtrim is necessary but not sufficient for this drive, the mechanism by which a reduction in Mtrim protein leads to drive of the B chromosomes is unknown. Mtrim has a critical function during female meiosis where it regulates Polo kinase (Polo) to promote centromeric cohesion of achiasmate (non-crossover) chromosomes and ensure their proper segregation during the meiotic divisions. This regulation requires Mtrim to bind to Polo through a highly conserved ST/P domain on Mtrim. One residue, T40, is particularly important for this interaction, as expression of a transgenic copy of Mtrim that has single amino acid change at that position (mtrim^{T40A}) is unable to rescue defects in the segregation of achiasmate chromosomes in a mtrim^{TM3} heterozygote. Since Mtrim’s interaction with Polo is necessary to promote the proper segregation of achiasmate chromosomes, I hypothesize that this interaction may also be essential for Mtrim’s ability to suppress the drive of the B chromosomes. To address this hypothesis, we will compare B chromosome transmission frequencies in a drive-competent (mtrim^{TM3, Sb Ser}) genetic background both in the presence and absence of Mtrim transgenes that affect Mtrim's ability to bind Polo (e.g., mtrim^{T40A}). Our investigation will determine if Mtrim’s interaction with Polo is critical for the suppression of B chromosome meiotic drive, indicating that Mtrim’s role in drive suppression and achiasmate chromosome segregation may be mechanistically similar. If the Mtrim-Polo interaction is not required for suppressing B chromosome meiotic drive, this result would lead us to test the necessity of other conserved Mtrim domains in its role as a drive suppressor. Overall, our work is uncovering the genetic mechanisms behind how the host genome protects itself against the proliferation of selfish genetic elements.

702B Crossover interference through ATR phosphorylation of Mei218 leading to phase separation of RING finger proteins Susan McMahan, Colette Anikwue, Jeff Sekelsky University of North Carolina at Chapel Hill

Sturtevant reported in his classic 1913 paper that the presence of one meiotic crossover (CO) discourages others, a phenomenon Muller termed interference. More than 100 years later we still don’t know the mechanism of interference. We propose a model in which induced phase separation of pro-CO proteins from the synaptonemal complex (SC), a
protein structure that forms between paired homologous chromosomes, followed by coarsening, these pro-CO proteins to accumulate at a single site or widely separated sites.

In Drosophila, ~20-24 double-strand breaks (DSBs) are made per meiosis, distributed uniformly throughout the euchromatin (at large scale). About 1/4 of these are designated to become COs and the rest are repaired as non-crossovers. In her EM studies of recombination Drosophila, Carpenter discovered a CO-associated structure she called the recombination nodule (RN), a 100 nM sphere that sits on top of the SC. The proposal by Dernburg and colleagues that the SC had liquid-like properties and that recombination factors can move within this compartment suggest the RN may be a droplet containing CO-designating proteins.

We propose that the pro-CO mei-MCM complex (Mei-217, Mei-218, and Rec) is localized to DSB sites. A 2nd set of pro-CO factors, the RING-finger proteins Vilya, Narya, and Nenya (here abbreviated VNN), move within the liquid SC. Phosphorylation of the intrinsically disordered domain of Mei-218 by Mei-41 (ATR kinase) promotes physical interaction with VNN. VNN accumulates and dewets from the SC. Exchange between nearby occurs through the SC, but larger accumulations become more stable, so that each SC (i.e., each chromosome arm) eventually has a single VNN or widely separated VNN foci. These sites are designated as COs. This process naturally leads to crossover assurance - the guarantee that each pair of homologous chromosomes (excepting the 4 in Drosophila melanogaster) almost always get at least one crossover to promote accurate segregation. In our view, interference is merely a consequence of the assurance process.

We made numerous in situ mutations in mei-218 to alter ATR phosphorylation sites. Mutation of S/T to the phosphomimetic residue D results in decreased COs and loss of interference, with increasing severity for 3, 5, or 8 sites mutated. In the most severe case there is high negative interference, where double-COs in two adjacent intervals are significantly more frequent than expected if the intervals are independent of one another. In addition to these data, progress toward dissection of different phosphorylation sites and mapping of interactions between VNN proteins and phospho-Mei-218 will be presented.


703C  Meiotic Crossover Patterning: The Centromere Effect Nila Pazhayam, Jeff Sekelsky University of North Carolina at Chapel Hill

Crossing-over between homologous chromosomes is a critical part of meiosis that prevents aneuploidy by promoting proper segregation of chromosomes. By facilitating accurate disjunction of homologs, crossing-over forestalls miscarriages and chromosomal disorders such as Down syndrome, the risk of which increases with maternal age. Meiotic crossovers (COs) are formed from programmed double-strand breaks (DSBs) that undergo homologous recombination. Although the DSBs that initiate crossing-over are distributed throughout the chromosome, intricate patterning governs where COs are placed. Three types of patterning events have been observed, one of which is the centromere effect (CE) that ensures CO exclusion in the regions surrounding the centromere. The CE is crucial to the meiotic cell, as centromere-proximal COs increase risk of nondisjunction. Despite its importance, the mechanisms behind this patterning event are poorly understood. To address this gap in knowledge, I will investigate the mechanisms underlying the CE using Drosophila as a model system.

Pericentric heterochromatin in Drosophila is not homogenous and is instead divided into two classes: the heterochromatin closer to the centromere (alpha heterochromatin) consists of highly repetitive satellite arrays, while that adjacent to euchromatin (beta heterochromatin) is less repetitive with some amount of unique sequence. Recent work from our lab has shown that the CE in Drosophila manifests as a complete exclusion of COs in alpha heterochromatin, and we hypothesize that this could be due to an absence of DSBs in this region. To test this, I will study differences in the number and positioning of DSBs in flies with heterochromatin defects, as compared to wild type. Recent work from our lab has also suggested that the CE in beta heterochromatin and proximal euchromatin is distance dependent. I will investigate whether this distance effect is based on distance from the centromere, or alpha heterochromatin, by deleting a large satellite array from the pericentric regions of the X chromosome. This will not only remove a large segment of alpha heterochromatin, but also move beta heterochromatin and proximal euchromatin closer to the centromere. Measuring CO rates in these regions will determine how the CE in proximal euchromatin and beta heterochromatin is influenced by proximity to alpha heterochromatin and the centromere. Another avenue that I plan on investigating is the sensitivity of the CE to the number of centromeres. I will study this by measuring crossover rates in flies that have a reduced total number of centromeres through the use of compound chromosomes. Through these and other methods, my overarching goal is to gain more understanding of the mechanisms at work behind crossover patterning, particularly the exclusion of crossovers near the centromere.

704A  Mechanisms and regulation of meiotic recombination: a whole-genome approach Carolyn Turcotte, Jeff Sekelsky University of North Carolina - Chapel Hill
Meiosis is a specialized form of cell division that generates haploid gametes from diploid cells. Crossovers between homologous chromosomes are essential to accurate chromosome segregation during meiosis, and chromosome missegregation leads to miscarriage and chromosomal disorders such as Down syndrome. Crossovers are generated via homologous recombination, a process that repairs double-strand DNA breaks (DSBs) using a homologous template. Many more breaks than crossovers are formed, and the number and position of crossovers are highly regulated in a phenomenon termed “crossover patterning.” Despite crossover patterning’s discovery over a century ago, the mechanisms that govern it remain unclear. A classical model for homologous recombination suggests that all crossovers are derived from a single symmetrical intermediate, with noncrossover products being derived from an earlier intermediate. The multiple pathways within homologous recombination can be traced via heteroduplex DNA (hDNA), DNA in which the strands are derived from different parent molecules (e.g. homologous chromosomes). Normally, SNPs and indels between the strands in hDNA are repaired by mismatch repair to form gene conversions or restorations of the original sequence. Our lab has found that Drosophila Xpc; Msh6 mutants, deficient in both canonical and short-patch mismatch repair, exhibit continuous hDNA tracts, making Drosophila the only metazoan thus far in which this has been achievable. Our lab has studied these hDNA tracts at a test locus and found hDNA signatures conflicting with the classical recombination model. However, much more extensive analysis than is possible at this test loci is required to revise this model. I am conducting whole-genome sequencing of hDNA to determine if the signatures seen at a single locus are consistent genome-wide, and whether certain hDNA signatures are localized to specific regions of the genome. This analysis will shed light on meiotic recombination mechanism and will inform if and how pathway choices in recombination differ in different genomic contexts. Understanding meiotic recombination mechanism and pathway choice is crucial to determining its connection to crossover patterning and proper homolog segregation.

**705B Tissue specific requirements of the Rcd4:Ana3 sub-complex in Drosophila centriole assembly** Pallavi Panda1, Levente Kovacs1, Nikola Dzhindzhev2, Agnieszka Fatalska1, Veronica Persico3, Maria Giovanna Riparbelli2, Giuliano Callaini3, David M Glover1 1) California Institute of Technology, United States; 2) University of Cambridge, United Kingdom; 3) University of Siena, Italy

Centrosomes are macromolecular protein complexes assembled through precise protein-protein interactions. Such proteins regulating centriole duplication appear to assemble and function in sub-complexes; Sas6 bound to phosphorylated Ana2 initiates pro-centriole formation; Sas4 interacts with centriolar microtubules; the Cep135:Ana1:Asl network promotes centriole to centrosome conversion enabling recruitment of Plk4 and peri-centriolar material. We have recently described a new sub-complex formed in the Drosophila centriole between Rcd4 and the C-terminal half of Ana3. As a poorly characterized Drosophila centriolar protein, we describe null and hypomorphic rcd4 mutant flies that exhibit fewer centrioles and aberrant mitoses in several somatic tissues and show structural defects in the basal bodies of sensory organs. Structured illumination microscopy of cultured cell centrioles reveals that Rcd4 and Ana3 load onto zone I of the centriole during interphase, where Ana3 recruitment precedes Rcd4. We find that neither Ana3 nor Rcd4 participates directly in the mitotic conversion of centrioles to centrosomes, but assembly of the Rcd4:Ana3 sub-complex is a pre-requisite for the recruitment of Ana1 and subsequently for the transition to the final stage of centriole to centrosome conversion. In contrast, we show differing requirements of the sub-complex for centriole development in the male germline. Whereas ana3 mutants are male sterile indicating that Ana3 is essential for centriole formation during spermatogenesis, rcd4 mutants are male fertile, yet display a multitude of centriolar defects. Contrary to our findings in somatic cells, elongated centrioles of ana3 and rcd4 mutant spermatocytes still accumulate Ana1 and Asl. Moreover, it appears that while Rcd4 is not required for the initial recruitment of Ana3 to spermatogonia centrioles, it is needed for the recruitment/maintenance and accurate localization of Ana3 upon centriole elongation in primary spermatocytes. Taken together, our results suggest tissue-specific roles of Rcd4 and Ana3 in centriole and basal body formation.

**706C Functional domains of the Ana1 centriole protein and their regulation by mitotic protein kinases and phosphatases** Agota Nagy1, Nikola Dzhindzhev2, Pallavi Panda1, Levente Kovacs1, Helene Randone2, Jingyan Fu1,2, Zoltan Lipinszki1,2, David Glover1,2 1) Division of Biology and Biological Engineering, California Institute of Technology, Pasadena, California, USA; 2) Department of Genetics, University of Cambridge, Cambridge, UK; 3) State Key Laboratory of Agrobiotechnology, College of Biological Sciences, China Agricultural University, Beijing, China; 4) Biological Research Centre, Institute of Biochemistry, MTA Lendület Laboratory of Cell Cycle Regulation, Szeged, Hungary

Centrioles are evolutionary conserved, 9-fold symmetrical structures that form the core component of centrosomes and also serve as basal bodies for cilia and flagella. They play critical roles in cell division, motility and signalling. The assembly of centrioles through the cell cycle is tightly regulated, leading to their elongation in the G2 phase followed by their conversion to centrosomes during mitosis. These processes require the progressive assembly of a network of interacting proteins, which extend from the inside to the outside of the centriole. Here we focus on Ana1, the protein that links the inner and outer centriole components of this molecular network. We wish to understand the independent function of Ana1’s domains and to this end, have created transgenic flies expressing N- and C-terminal fragments of different lengths. We separately express these fragments in different ana1 mutant backgrounds and determine the extent to which they can function independently to drive centriole duplication and elongation in different tissues. We are
determining how the interactions of Ana1’s different functional domains with other key centriole proteins are dependent upon phosphorylation by mitotic protein kinases for completion of the procentriole, its disengagement from the mother centriole, and recruitment of peri-centriolar material and how protein phosphatases ensure the cyclical nature of some of these interactions.

707A  **Cohesin dynamics during meiotic prophase in Drosophila oocytes** Muhammad Abdul Haseeb, Sharon Bickel Dartmouth College

Premature loss of sister chromatid cohesion during the meiotic arrest in human oocytes is a major contributor to the maternal age effect, a phenomenon in which the probability of an aneuploid pregnancy increases as a woman ages. Our laboratory has previously uncovered evidence for a cohesion rejuvenation program in Drosophila oocytes that operates during prophase, after cohesive linkages are originally established during S phase. To better understand the behavior of rejuvenation-specific cohesin, we designed an “allele-switch” transgene that expresses endogenous levels of the cohesin subunit Smc1 tagged with mCherry which is flanked by FRT sites. Induction of Flippase using the prophase specific mata-Gal4 driver causes excision of the mCherry tag, resulting in expression of Smc1 tagged with superfolder GFP (sGFP). This tool allows us to specifically visualize chromatin-associated cohesin that is synthesized after the canonical cohesion establishment that occurs during oocyte DNA replication. Upon switch induction with the mata driver, newly synthesized Smc1-sGFP begins to load onto the meiotic chromosomes as early as oogenesis stage 2. Meiotic prophase lasts ~6 days in Drosophila oocytes. Surprisingly, almost the entire cohesin population appears to turn over in < 2 days (by oogenesis stage 5). Turnover at centromeres is slower than at arms. Mata-induced knockdown (KD) of the cohesin subunit Smc3 abolishes chromosomal loading of newly synthesized Smc1-sGFP. This failure to load rejuvenation-specific cohesin during prophase leads to premature loss of arm cohesion in metaphase I arrested oocytes, as evidenced by fluorescence in situ hybridization that uses Oligo-PAINT probes to visualize X-chromosome arm cohesion. Together these results indicate that rejuvenation in Drosophila oocytes is required to maintain levels of cohesion that are sufficient for accurate chromosome segregation. Future experiments will utilize the allele-switch transgene to investigate cohesin loading dynamics when cohesion regulators or rejuvenation specific proteins are knocked down during prophase. These experiments will allow us to better understand the mechanism(s) underlying cohesion rejuvenation during meiotic prophase in Drosophila oocytes.

708B  **Discs large licenses Pins to orient mitotic spindles** Kathryn Neville, Dan Bergstralh University of Rochester

Oriented cell divisions are critical to animal development. Division orientation determines the placement of daughter cells, and thereby promotes cell diversity and helps to organize and expand tissues. The direction in which a cell divides is established by the orientation of the mitotic spindle at metaphase. Spindle orientation in animals typically relies on an evolutionarily conserved machine comprised of at least four proteins: in flies, these are called Partner of Insuteable (Pins), Gai, Mushroom body defective (Mud), and Dynein. The canonical model is that Pins acts as a cortical anchor for Mud and Dynein, which exert a pulling force on astral microtubules that reels the spindle into alignment. Another established, but poorly understood, component of this machinery is Discs large (Dlg), an apical-basal polarity factor that interacts with phosphorylated Pins. Whereas previous work from our lab and others suggests that Dlg is required for Pins localization, I show here that Pins localization is unaffected by the loss of Dlg. My results indicate that Dlg works to activate, rather than localize Pins function. These results suggest that the canonical model is incomplete, raising the question of how the spindle orientation machinery is regulated.

709C  **Evolutionarily conserved midbody reorganization precedes ring canal formation during gametogenesis** Kari Price, Dyuthi Tharakan, Lynn Cooley Yale School of Medicine

During gametogenesis, male and female germ cells develop as syncytia connected by intercellular bridges that facilitate cell-cell communication and the transport of molecules and organelles between sister cells. Functional studies demonstrate the requirement of germline bridges for fertility, yet the mechanism underlying their formation is still unclear. Immunofluorescence studies of fixed Drosophila tissues have suggested that these bridges, otherwise known as ring canals, arise from arrested constriction of the contractile ring. In contrast, studies in the mouse testis have suggested that germline bridges are derived from the midbody matrix. These observations, combined with the compositional differences between Drosophila male and female ring canals and mouse and Drosophila male ring canals, raise the question of whether there is one underlying, conserved mechanism of ring canal formation. To investigate how ring canals are formed, we performed time-lapse imaging of the conserved ring canal component Pavarotti/MKLP1/kinesin-6 during the incomplete mitotic divisions of the male and female Drosophila germlines. In both testes and ovaries, rather than contractile ring arrest, contractile rings constrict completely to form a dense, transient midbody intermediate that reorganizes into an open ring canal in less than one hour. The midbody-to-ring canal transition appears to be a conserved feature of gametogenesis as we also observed kinesin-6-labeled midbodies in fixed preparations of mouse and Hydra vulgaris male germlines.

To further characterize the midbody-to-ring canal transition, we imaged known midbody ring components in the
Drosophila testis. We observed that the germline midbody is compositionally similar to, but functionally distinct from, the midbodies formed during complete cytokinesis. Known midbody ring components Septin 2 and Citron kinase/Sticky localize to the germline midbody, but as expected, germline midbodies fail to initiate microtubule severing and abscission. Interestingly, in addition to the midbody ring localization, Sticky localizes to a subset of ring canals, presumably nascent ring canals, but not mature ring canals suggesting a possible role in the midbody-to-ring canal transition. In fact, knockdown of sticky in the male germline (nos>sticky RNAi) results in persistent midbody-like foci. We are now working to understand the precise role of Sticky during the midbody-to-ring canal transition and identify additional components required.

710A Functional Analysis of Bloom Syndrome Helicase in Development and DNA Repair Colleen Bereda, Evan Dewey, Jeff Sekelsky University of North Carolina at Chapel Hill, Chapel Hill, NC

Bloom Syndrome is a rare autosomal recessive disorder in humans caused by mutation of the BLM gene that leads to increased genome instability and cancer. The BLM gene codes for a helicase (BLM) that works together with Topoisomerase 3-alpha (Top3α) in homology-directed repair of DNA. Top3α assists by directly binding to BLM and helping to release the torsional stress on DNA as BLM helicase unwinds recombinatory intermediates. These proteins preserve genome stability and have been shown in many organisms to operate together in the prevention of detrimental mitotic (non-meiotic) crossovers via two main DNA repair pathways, Synthesis-Dependent Strand Annealing and double-Holliday Junction dissolution. In Drosophila, BLM (known as Blm) also has roles in proper meiotic chromosome segregation and rapid cell cycle progression of the developing embryo. To investigate the Blm-Top3α interaction in DNA repair, I performed a yeast 2-hybrid (Y2H) assay using the Drosophila proteins. I found interaction was specific to certain regions of Blm, with the strongest observed for a C-terminal region conserved among several Drosophila species, amino acids (aa) 1381-1487. To further decipher Blm roles, I created specific Blm deletions of regions from my Y2H data via CRISPR/Cas9 editing and tested various Blm roles in vivo. First I examined the role of Blm in promoting embryonic development. While deletion of aa 1-240 did not have an effect, deletion of aa 576-720 had substantial impact with most embryos failing to hatch. I then assessed the effects of Blm deletions on known Blm roles in meiotic chromosome segregation via a nondisjunction assay. Both aa 1-240 and aa 576-720 deletions were not significantly different from our wild-type control in contrast to my embryonic development data. This result suggests deletion of aa 576-720 is a separation-of-function mutation. Future studies will assess importance of these Blm regions in preventing mitotic crossovers and determining their importance to Blm function in DNA repair. I will also further explore Blm aa 576-720 and the relevance of predicted ATR/ATM (Drosophila Mei-41 and Tefu; regulators of DNA repair) phosphorylation sites within this region for proper Blm function in embryonic development, meiotic segregation, and DNA repair. By characterizing the functions of Blm in Drosophila, we will better understand BLM function within humans and the detrimental health effects of BLM mutations.

711B Alternative End Joining Preferences in RPA-Deficient Drosophila Jacob Zuckerman, Terrence Hanscom, Nick Woodward, Vicki Do, Mitch McVey Department of Biology, Tufts University, Medford, MA

DNA double-strand breaks can be repaired using several different repair pathways. These include homologous recombination, classical-non homologous end joining, and a third, less well-understood set of mechanisms known as alternative end-joining. Alternative end joining repair is extremely inaccurate, causing insertions, deletions, and/or indels and has been linked to several different types of cancers. One type of alternative end joining, known as synthesis-dependent microhomology mediated end-joining (SD-MMEJ), occurs when short repeated sequences near the break site form secondary structures that prime nascent synthesis of microhomologies that are then used to repair the DNA. In yeast, the replication protein A (RPA) heterotrimer has been shown to prevent another similar alternative end-joining pathway, microhomology mediated end-joining. To bind efficiently, RPA requires approximately 30 base pairs of single stranded DNA. Previous data from our lab demonstrate that movement of the short-repeated sequences to a distance greater than 30 bp from the break site reduces the frequency of SD-MMEJ usage from 15% to <1%. Therefore, we hypothesize that RPA that may inhibit SD-MMEJ by binding to short stretches of single stranded DNA and preventing secondary structures from forming. To test this, we have obtained a fly stock with a P element inserted in the 5’ UTR of the RPA70 gene, which severely decreases RPA70 expression. Cas9-induced breaks will be generated in a set of reporter plasmids that are injected into embryos from this stock. Deep sequencing and computational analysis of the inaccurate repair products will allow us to discern the role of RPA in SD-MMEJ repair. In addition, these studies may shed light on alternative end joining repair processes in cancer cells, which likely experience RPA exhaustion due to the accumulation of single-stranded DNA during perturbed DNA replication.

712C Defining Mitotic Crossover Mechanisms Using CRISPR/Cas9 and Bloom Syndrome Helicase Evan Dewey, Jeff Sekelsky University of North Carolina-Chapel Hill

Genome stability is key to longevity of multicellular organisms and avoidance of disease. Despite daily challenges from numerous sources of DNA damage threatening this stability, cells regularly repair DNA to maintain genomic resilience. Improper or misregulated repair causes accumulation of “scars” in the form of detrimental mutations within...
the genome, eventually leading to genome instability, cancer, and other genetic disease. Homology directed repair (HDR) of DNA double strand breaks is one DNA repair pathway that, if improperly regulated, leads to accumulation of mutations via mitotic (non-meiotic) crossovers and loss of heterozygosity. Therefore, better understanding of mitotic crossover mechanisms and regulators is critical to prevention of cancer and other genetic disease. CRISPR/Cas9 has also become increasingly reliant on accurate HDR to integrate desired mutations or corrections in genome editing, but the precise CRISPR/Cas9 HDR mechanisms remain elusive. I have begun to test mitotic crossover mechanisms using both CRISPR/Cas9 induced double strand breaks and mutation of Bloom Syndrome Helicase (Blm), a key HDR regulator that prevents mitotic crossovers. Through combination with total mismatch repair (MMR) knockout only possible in Drosophila (through both Msh6 canonical and backup Xpc short-patch pathways), it is now possible to analyze resulting mitotic crossover products using Sanger sequencing for CRISPR/Cas9 induced double strand breaks and whole genome sequencing for Blm mutants. Using these tools, I have begun to accurately define mitotic crossover mechanisms for the first time in a multicellular organism, with an unligated double Holliday junction resolution model indicated as the primary mechanism. Unexpectedly, combination of Blm and MMR knockout also appears to substantially increase the rate of mitotic crossovers compared to knockout of either alone, indicating a previously uncharacterized genetic interaction between Blm and MMR components in HDR and prevention of mitotic crossovers. This work has begun to enhance understanding of how DNA is repaired in both CRISPR and Blm mutant contexts, expanding knowledge of how mitotic crossovers lead to genome instability, identifying previously uncharacterized interactions in HDR and mitotic crossover regulation, and providing better understanding of how to beneficially utilize mitotic crossover mechanisms in genome editing.

713A  The Krüppel-like factor Cabut has cell cycle regulatory properties similar to E2F1 Peng Zhang1,2, Alexia J. Katzaroff1,2, Laura A. Buttitta1,2, Yiqin Ma1,2, Huaqi Jiang3, Derek W. Nickerson4, Jan Inge Øvrebø1,2, Bruce A. Edgar1,2,3,4 1) Huntsman Cancer Institute, University of Utah, Salt Lake City, UT 84112; 2) Department of Oncological Sciences, University of Utah, Salt Lake City, UT 84112; 3) Molecular and Cellular Biology Program, University of Washington, Seattle, WA 98195; 4) Division of Basic Sciences, Fred Hutchinson Cancer Research Center, Seattle, WA 98109

Using a gain-of-function screen in Drosophila, we identified the Krüppel-like factor Cabut (Cbt) as a positive regulator of cell cycle gene expression and cell proliferation. Enforced cbt expression is sufficient to induce an extra cell division in the differentiating fly wing or eye, and also promotes intestinal stem cell divisions in the adult gut. Although inappropriate cell proliferation also results from forced expression of the E2f1 transcription factor or its target, Cyclin E, Cbt does not increase E2F1 or Cyclin E activity. Instead, Cbt regulates a large set of E2F1 target genes independently of E2F1, and our data suggest that Cbt acts via distinct binding sites in target gene promoters. Although Cbt was not required for cell proliferation during wing or eye development, Cbt is required for normal intestinal stem cell divisions in the midgut, which expresses E2F1 at relatively low levels. The E2F1-like functions of Cbt identify a distinct mechanism for cell cycle regulation that may be important in certain normal cell cycles, or in cells that cycle inappropriately, such as cancer cells. That Cabut has E2F-like activity is especially interesting when one considers that the E2Fs, which are the best characterized transcriptional regulators of the eukaryotic cell cycle, are not actually essential for cell cycle gene transcription or cell cycle progression. Hence, it is clear that other transcription factors must also regulate the transcription of the 300 to 400 genes necessary for DNA replication and mitosis. Cabut appears to be one of these missing transcription factors that can complement E2F1.

714B  Excess histone H3 is a Chk1 inhibitor that controls embryonic cell cycle progression Yuki Shinso, Amanda Amodeo Dartmouth College

The early embryos of many species including Drosophila spend the first few hours of development undergoing a series of rapid cell cycles to generate the large number of cells required for embryogenesis. These early reductive divisions are largely transcriptionally silenced and therefore depend upon pools of maternally provided components. Cessation of the rapid cell cycle program at the mid-blastula transition (MBT) is dependent on achieving a sufficient nuclear to cytoplasmic (N/C) ratio and the activity of the DNA-damage checkpoint kinase, Chk1 (grapes). How Chk1 activity is coupled to the N/C ratio has remained unclear. We have shown that large stores of extra-nucleosomal histone H3 accumulate in the early nuclei because the maternally provided histone pools vastly exceed the amount of DNA in the early embryo. As the N/C ratio increases these pools are exhausted. At the same time, histone H3 is a known Chk1 substrate. Therefore, we hypothesized that in the early nuclei H3 may compete with Chk1’s cell cycle regulatory targets (CDC25-string and twine) to prevent cell cycle slowing before the correct N/C ratio is reached at the MBT. To test this we expressed a version of the H3 N-terminal tail that contains the Chk1 binding site, but cannot be incorporated into chromatin in the pre-MBT embryo. We found that this 47-amino acid fragment is sufficient to delay cell cycle slowing and reduce Chk1 activity in vitro and in vivo. Mutation of the Chk1 phosphosite in the context of the full length H3 protein is sufficient to lengthen the cell cycle and accelerate the onset of the MBT. Together our data support a model in which hyper-abundant histone H3 in the early embryo competitively inhibits Chk1 to prevent cell cycle slowing until a sufficient N/C ratio is achieved. We are currently working to determine if H3 inhibits Chk1 in other developmental contexts including in response to DNA-damage.
715C  Regulation of induced endocycling cells and their effects on tissue growth Hunter Herriage, Yi-Ting Huang, Savanna Brown, Brian Calvi Indiana University

Non-canonical variations of the cell cycle are important in development and disease. One of these variations is the endocycle, which results in large, polyploid cells that arise through inhibition of mitosis. In addition to these developmental endocycling cells (devECs), conditional signals can result in induced endocycling cells (iECs), such as during wound healing, aging, or cancer. We have previously shown in Drosophila that devECs and iECs repress the apoptotic response to DNA damage. With the Walczak lab, we have also shown that transient iECs in both flies and humans can undergo error-prone mitosis (RTM). These data are consistent with mounting evidence that cancer cells can transiently enter the endocycle in response to genotoxic therapies, and can then switch back to error-prone, polyploid mitoses that contribute to genome instability and tumor regrowth. Much remains unknown, however, about how the growth of transient iECs is regulated and how iECs affect tissue homeostasis and tumorigenesis.

We will report our progress using Drosophila as an in vivo model system to decipher how endocycles contribute to normal and abnormal tissue growth. Particularly, we will discuss our investigations into the possible synergies between pro-growth mutations and different variant cell cycles. Although iECs repress apoptosis, we will present evidence that a subset may senesce, depending on developmental context. This work will lead to a greater understanding of growth, proliferation, and checkpoint regulation in polyploid cells, and their effects on tissue homeostasis and cancer.

716A  Defining the Dynamics of Transcriptional Bursting in Developing Drosophila legs Rina Helt, Elizabeth Urban, Robert Johnston Johns Hopkins University

Transcription is a dynamic process during development. Much of our understanding of transcription comes from studies of snapshots of individual timepoints. Live imaging approaches based on MS2-MCP technologies have provided numerous insights into transcriptional dynamics during embryonic development, yet we still know very little about how these dynamics affect later developmental processes. Studying transcriptional bursting in genes that regulate numerous insights into transcriptional dynamics during embryonic development, yet we still know very little about how these dynamics affect later developmental processes. Studying transcriptional bursting in genes that regulate cell fate determination in different contexts.

I will determine how the transcriptional dynamics of a single gene can impact its regulation of cell fate determination in different contexts. By assessing the bursting patterns of different contexts. By assessing the bursting patterns of a subset may senesce, depending on developmental context. This work will lead to a greater understanding of growth, proliferation, and checkpoint regulation in polyploid cells, and their effects on tissue homeostasis and cancer.

717B  Molecular genetic analysis of the mutation I.3.2 by undergraduates participating in a Drosophila CURE Cory Evans¹, Veronica Casarez¹, Kayla Bieser², Jacob Kagey³ 1) Loyola Marymount University; 2) Nevada State College; 3) University of Detroit Mercy

The mutation I.3.2 was identified in an FLP/FRT-based genetic screen for novel regulators of cell growth located on the right arm of chromosome 2 (2R). Because abnormal growth phenotypes associated with new mutations may be suppressed by the parallel activation of apoptosis, EMS was used to mutagenize an FRT42D chromosome already carrying the Dark mutation, which strongly suppresses apoptosis. This screen successfully identified several new mutations affecting cell growth including I.3.2. Some of these mutations have been or are being characterized by undergraduate students participating in FlyCURE, a course-based undergraduate research experience (CURE) that is being implemented at several partnering colleges and universities across the country. Here we describe the FlyCURE analysis of mutation I.3.2, which behaves zygotically as a recessive lethal. I.3.2 is likely also a cell-lethal mutation, as flies with mitotic mutant clones created in the developing eye (using ey-FLP) are pupal lethal and exhibit an almost complete lack of developing head structures. Complementation mapping of the I.3.2 recessive-lethal phenotype using the Bloomington Drosophila Stock Center 2R Deficiency Kit identified two non-complementing deficiencies. These deficiencies overlap in the cytological interval 50A7-50A13, a small chromosomal region containing thirteen genes. A lethal allele of one of these genes, centromere identifier (cid), failed to complement I.3.2, suggesting that I.3.2 is an allele of cid. DNA sequence analysis of the cid locus in the I.3.2 genetic background is underway to confirm the identity of the I.3.2 gene and determine the molecular nature of the mutation.

718C  Probing the Temporal Regulation of Hatching in D. melanogaster Alexandra (Olenka) Jain¹,², Stanislav Shvartsman 1,² 1) Princeton University; 2) Lewis-Sigler Institute of Integrative Genomics

The ability to regulate the timing of irreversible transitions between sequential developmental states is one of the
most critical features of any developmental process. Understanding what controls the variation in the duration of a developmental stage is an important question in developmental biology. We address this question using Drosophila melanogaster embryogenesis because it has an extremely well-defined start and end. We describe a straightforward approach to characterize the inter-embryonic variability in the duration of embryogenesis by measuring the time of hatching events in cohorts of synchronized embryos. This approach is based on videos taken from oviposition until hatching, automated detection of hatching events, and nonparametric statistical analysis of the hatching time distributions. We demonstrate how the empirical distribution functions of hatching times can be used to study the effects of maternal age and genotype on the duration of embryogenesis. Specifically, this approach can be used to characterize the hatching distribution function of flies with a mutation in the amontillado gene, which has been shown to be important for the duration of embryogenesis (Siekhaus & Fuller, 1999). In addition to this, we revisit the connection between hatching and molecular and motor activities in the embryo.

719A Identification of Apoptosis and Junctional Tension as Pro-tumoral Factors in Drosophila Mariianne Montemurro, Bruno Monier, Magali Suzanne Centre de Biologie Integrative CBI - Molecular, Cellular and Developmental Biology Unit - UMR 5077

Cancer is a largely widespread pathology that corresponds to an overproliferation of cells that could finally invade other tissues. Tumors develop through three increasingly aggressive steps: (1) hyperplasia, which corresponds to cells overproliferation without any modification of epithelial properties; (2) neoplasia, during which cells can acquire a more mesenchymal phenotype, and finally (3) metastasis, when cells leave the primary tumor, migrate and form secondary tumors.

Tumor development can be influenced by mutations (or combination of mutations) but also by external factors, such as extracellular matrix rigidity. However, a comprehensive understanding of the factors driving tumor evolution is still lacking. My project aims to use Drosophila to identify unexpected factors that could influence tumor development, and more specifically the hyperplasia/neoplasia transition, a critical step in tumor aggressiveness. After an in-depth characterization of tumor progression at cellular level at successive timepoints, I selected two complementary genetic contexts for further analysis: a strictly hyperplastic tumor (“Yorkie overexpression” context) and an initially hyperplastic tumor that eventually evolves in neoplasia (“Avalanche loss-of-function” context).

Strikingly, I identified two new and unexpected factors involved in tumor progression and aggressiveness: apoptosis and tissue tension. Indeed, while tumor aggressiveness coincides with a high level of apoptotic cells, abolishing apoptosis in Avalanche tumors strongly decreases the hyperplasia/neoplasia shift, while introducing cell death in Yorkie tumors converts hyperplastic tissues into neoplastic ones. Hence, in those two different contexts, modulation of apoptosis surprisingly favors the hyperplasia/neoplasia transition. Moreover, I found that tumor aggressiveness is associated with high tissue tension and that increasing junctional tension (by modulating a Myosin II regulator) is sufficient to induce a hyperplasia/neoplasia transition in a tumor context normally strictly hyperplastic (Yorkie’s context).

I am now characterizing whether cell death and tissue mechanics are linked or whether they act independently of one another to drive tumor progression.

720B Examining the synthetic lethality between BRCA2 and methyl and ethyl Paraben Zainab Rizik, Nicole Banuelos, Blake Riggs, Lela Legesse San Francisco State University

There have been increases in incidences of breast cancer since 1970 due to increased exposure to chemicals in our daily lives. Parabens are found in our everyday lives and are disproportionately pushed on poor communities via personal products and food. Recent studies have found that parabens mimic estrogen and attach to the receptors of breast cancer cells and cause them to proliferate, however the pathway and additional targets are poorly understood. The BRCA2 gene plays a vital role in providing instructions for creating a tumor suppressor protein preventing cells from excessively multiplying. When there is a mutation in this gene, it could cause breast cancer due to the loss of function of the protein. This experiment analyzes the effect of the toxins methyl and ethyl paraben on organisms with BRCA2 gene mutations using Drosophila melanogaster. To study this, the survivability of homozygous BRCA2 mutant, we exposed larvae to different concentrations of parabens (300, 700, 1000 mg/l) and determined the survival rate. Overall, it was found that there was increased synthetic lethality in flies and at increased doses of parabens. These findings will pave the way in lowering the amount of future breast cancer cases caused by the use of methyl and ethyl paraben.

721C Candidate ion channel screen identifies modifiers of brain tumor size Isabella Maag, Ashley Bielawski, Hannah Christman, Beverly Piggott University of Montana, Missoula, MT

Ion channel function is critical for normal development and prevention of disease. These functions include maintaining membrane potential, controlling the flow of ions across the membrane, and regulating cell volume. Ion channels have well described roles in excitable cells where they are essential for establishing membrane potentials, the firing of action potentials and contraction of muscles among other functions, but their role in non-excitable cells is less defined. While ion channels are found in neural progenitors and upregulated in cancer, their roles, in these tissues, are not well understood. Previous work has found that reduction of voltage-gated sodium channel Para reduces brain tumor size. To
further define the bioelectric landscape of proliferation, we sought to identify ion channels that influence tumor size and potentially proliferation. Using a Deadpan overexpression (Dpn\textsuperscript{OE}) brain tumor model which displays hyper-proliferation, we conducted a RNAi candidate ion channel screen to identify modifiers of brain volume. This screen has revealed a number of ion channels whose knockdown, significantly alters brain volume. This suggests that these ion channels may regulate important aspects of tumorigenesis. As ion channels are druggable targets, identifying modifiers of tumor size may represent new therapeutic sites of intervention.

722A Yorkie dependent transcriptional network promotes tumor growth arushi rai\textsuperscript{1}, Indrayani Waghmare\textsuperscript{3}, Amit Singh\textsuperscript{1,2}, Madhuri Kango-Singh\textsuperscript{1,2} 1) Department of Biology, University of Dayton, Dayton, OH; 2) Integrative Science and Engineering Center, University of Dayton, Dayton, OH; 3) Department of Cell and Developmental Biology Vanderbilt University, Nashville, TN

Studies in Drosophila and other tumor models have revealed cancer promoting signaling interactions and transcriptional addictions in tumors cells. The Hippo pathway effector, Yorkie (Yki) is a key mediator of such interactions, and presents an attractive opportunity to study transcriptional dependencies in cancer cells. The Ras\textsuperscript{V12}, srib\textsuperscript{-/-} tumor mosaic model is well-established and shows activation of oncogenic Ras in the background of impaired apical-basal polarity. This model is widely used study molecular mechanisms and signaling events downstream of the oncogenic Ras and Ras-mediated Yorkie (Yki) activation in Ras\textsuperscript{V12}, srib\textsuperscript{-/-} tumor cells. Previously, we have shown that in Ras\textsuperscript{V12}, srib\textsuperscript{-/-} cells Wingless (Wg), Caspases (e.g., the initiator caspase Dronc) and JNK are activated to promote tumorigenesis through their non-apoptotic roles. Within these, Wg/Wnt pathway is known to act via canonical and non-canonical pathways during development and cancer, and interact with Yki to promote cancer growth. Genetic epistasis showed that Wg acts upstream of Caspases, JNK and Yki, and downregulation of Wg reduced tumor growth by downregulation of Caspases, JNK and Yki reporters. Our goal is to further understand how the two evolutionarily conserved signaling pathways i.e., Hippo and Wingless crosstalk and interact with each other to regulate tumor growth. To understand this intricate wiring of Wingless-Yorkie during tumor growth and invasion, we will use the Ras\textsuperscript{V12}, srib\textsuperscript{-/-} tumor model in Drosophila imaginal discs. Preliminary data showed that wg transcriptional reporters are upregulated in Ras\textsuperscript{V12}, srib\textsuperscript{-/-} cells, suggesting that increased accumulation of Wg may be due to increased transcription. In other contexts, wg is shown as a transcriptional target of Yki. Therefore, we will test for (a) the effects of Yorkie protein, the main effector molecule of Hippo pathway, on wg transcription and expression of other Wg pathway components by reporter assays, and qRT-PCR-based approaches, and (b) feedback interactions that promote tumorigenesis using genetic epistasis-, and immunohistochemistry-based approaches. Here, we present our progress on the organization of the molecular network involving Wingless and Yorkie.

723B Fmi-mediated cell polarity and adhesion are critical during cell competition and tumorigenesis in Drosophila Pablo Sanchez Bosch, Bomsoo Cho, Jeffrey Axelrod Stanford University

Cell competition is a process by which cells with a competitive advantage (winners) eliminate less fit neighboring cells (losers). This phenomenon was first seen when cells harboring ribosomal protein mutations were eliminated by their WT neighbors. Since that first observation in 1975, cell competition has been linked to mutations in many other genes, i.e. tumor suppressors, oncogenes, or cell polarity components. Even though the outcome is always the elimination of the less fit cells, very little is still known about the mechanisms leading to the selection of fitter cells and elimination of the losers. Tumors have been seen taking advantage of cell competition to outcompete the surrounding WT tissue and promote survival and local invasion. Therefore, understanding the molecular cues of cell competition will give us not only insight into a key developmental process but a deeper understanding of tumorigenesis.

We have found that Fmi, a key component of the planar cell polarity (PCP) pathway, plays a critical role in various cell competition scenarios and Ras\textsuperscript{V12}-dependent tumorigenesis. Intercellular Fmi homodimers scaffold the assembly of the other core PCP proteins and mediate asymmetric signaling between the core PCP proteins. PCP signaling polarizes cells within the plane of the epithelium to orient cellular structures, cell divisions, and cell movement during development and homeostasis. In addition to establishing planar polarity, numerous reports have linked PCP signaling to cancer.

Our experiments show that Fmi is critical for winner cells in cell competition and tumorigenesis. Tumors and winner cells lacking Fmi lose their ability to eliminate neighboring loser cells. Moreover, fmi-null winners undergo apoptosis. Removing Fmi from loser cells does not have any effect on cell competition outcomes, suggesting that Fmi is only critical by winners to outcompete losers. Our findings suggest that Fmi acts as a scaffolding and signaling hub, where Fmi molecules, via PCP or other signaling networks, allow for efficient cell communication, necessary for the elimination of loser cells.

Our results are striking in that Fmi is critical in all the study cases we’ve tackled, and establish the necessary background for comprehending the mechanisms underlying cell-cell communication during both developmental cell competition and tumorigenesis.
Proper regulation of cell proliferation and cell death is essential for tissue homeostasis and organ growth in developing animals. Multiple evolutionarily conserved signaling pathways regulate the growth of tissues and organs. Hippo signaling is a conserved mechanism for controlling organ growth. Increasing evidence suggests that Hippo signaling is modulated by various cellular factors for normal development and tumorigenesis. Hence, identification of these factors is pivotal for understanding the mechanism for the regulation of Hippo signaling. *Drosophila* Mnat9 is a putative N-acetyltransferase that is required for cell survival by affecting JNK signaling. Here we show that Mnat9 is involved in the negative regulation of Hippo signaling. RNAi knockdown of Mnat9 in the eye disc suppresses the rough eye phenotype of overexpressing Crumbs (Crb), an upstream factor of the Hippo pathway. Conversely, *Mnat9 RNAi* enhances the eye phenotype caused by overexpressing Expanded (Ex) or Warts (Wts) that acts downstream to Crb. Similar genetic interactions between *Mnat9* and Hippo pathway genes are found in the wing. The reduced wing phenotype of *Mnat9 RNAi* is suppressed by overexpression of Yorkie (Yki), while it is suppressed by knockdown of Hippo upstream factors like Ex, Merlin, or Kibra. Mnat9 co-immunoprecipitates with Mer, implying their function in a protein complex. Furthermore, Mnat9 overexpression together with Hpo knockdown causes tumorous overgrowth in the abdomen. Our data suggest that Mnat9 is required for organ growth and can induce tumorous growth by negatively regulating the Hippo signaling pathway. It has been suggested that Mnat9 functions in developing *Drosophila* are evolutionarily conserved since loss-of-function phenotypes of Mnat9 can be effectively rescued by expressing human homolog protein hNAT9. Our data also indicate that Mnat9 and hNAT9 share similar features in inducing tumor growth through Hippo signaling interaction. As the biological function of hNAT9 in humans are currently unknown, these results are expected to provide essential information on the role of *hNAT9* in humans and how it relates to the Hippo signaling pathway.

Mutation on the gene *brain tumour (brat)* results in malignant neoplasms that originate from the typell neuroblast lineage. To identify the key oncogenic processes involved in brat tumour development we have combined a large-scale genetic screen to identify genes that can be targeted to inhibit tumour growth (brat-SPRs) and RNAseq transcriptome profiling. We have identified 75 brat-SPRs among 4000 randomly selected genes. Highly enriched among them are proteins involved in translation and vesicle trafficking. We have also found a total of 282 genes that are significantly dysregulated in the tumour. Our results show that upregulation of gene expression correlates poorly with functional requirement: only 5% of the genes that are most highly upregulated in brat tumours are brat-SPRs. Likewise, only 35% of all brat-SPRs identified are significantly upregulated in brat tumours. The same conclusions were derived from an earlier similar study on tumours caused by mutation in lethal(3)malignant brain tumour (l(3)mbt)$^2$, that are equally malignant, but originate from the neuroepithelium and present notably distinct transcriptomic signatures. Notably, there are only 20 genes in common between the 75 brat-SPRs and the reported 131 mbt-SPRs$^2$ thus showing that most of the suppressors of tumor growth are tumour-type specific.

The relaxin-like secreted peptide, Drosophila insulin-like peptide 8 (Dilp8), was first identified as a regulator of the delayed puation induced by neoplastic or damaged imaginal discs. Through its receptor leucine-rich repeat-containing G protein-coupled receptor 3 (Lgr3) in a pair of bilateral central brain neurons, Dilp8 represses the activity of PTTH neurons, leading to a delay in the surge of the steroid hormone Ecdysone that stimulates pupariation. In addition, Dilp8 and Lgr3 are required for intra- and inter-organ growth regulation, in which damage to discs can slow the growth of undamaged regions and other discs.

To deconstruct how Dilp8 contributes to wing disc growth during normal disc development we examined its expression and that of *lgr3* in whole larvae and wing discs during the 2nd and 3rd instars. We find that relative to whole larvae, *dilp8* is very highly expressed in early 3rd instar wing discs. However, by late 3rd instar *dilp8* expression is considerably decreased in both, in a temporal pattern that is reciprocal to Ecdysone receptor activity in the wing disc. We compared wing discs from wildtype (WT) and *dilp8* mutant larvae across the 2nd and 3rd instars, and find that in *dilp8* mutants wing discs are initially smaller than WT discs at the early 3rd instar, but reach a similar size as WT discs by late 3rd instar. These data suggest that Dilp8 regulates the kinetics of wing disc growth during normal development. Dilp8 overexpression has been shown to modulate Ecdysone, Juvenile Hormone and insulin/insulin-like growth factor signaling, suggesting Dilp8 may regulate these pathways under physiological growth conditions. We aim to determine whether and how normal developmental changes in hormonal signaling mediate Dilp8’s role in tissue growth.
B chromosomes are supernumerary genetic elements found in hundreds of species across a wide range of taxa. Even though many B chromosomes are present in low copy numbers (1-2), the B chromosomes recently discovered in a stock of *Drosophila melanogaster* are carried at a relatively high copy number of 10-12 B chromosomes. Though these B chromosomes do not appear to carry active genes and are mostly heterochromatin, wild-type females with B chromosomes exhibit a significantly increased frequency of Chromosome 4 nondisjunction, leading us to speculate that the presence of B chromosomes may have an influence on other important meiotic processes. We set out to investigate the impact of these B chromosomes on transcription of protein coding genes during gametogenesis in both males and females by performing RNAseq on mRNA extracted from whole testes and the tips of ovaries (gerarium through stage 6). Unfortunately, the original B chromosome stock carries a null mutation of *matrimony* (*mtrm*), a critical regulator of Polo kinase during female meiosis, held over the balancer chromosome TM3, Sb Ser, making it difficult to assign significant transcriptional changes due solely to the B chromosomes since a genotypically comparable stock does not exist. To overcome this obstacle, we outcrossed the original B chromosome stock to a *Pr Dr/TM3, Sb Ser* stock that does not carry B chromosomes to create two new stocks, both with B chromosomes and either *Pr Dr or mtrm* held over the TM3, Sb Ser balancer. Together with the *Pr Dr/TM3, Sb Ser* stock without B chromosomes, these three genotypes allow us to triangulate genes that are differentially expressed in the germline due to the presence of these B chromosomes and not because of other confounding genetic factors. Our transcriptome data is currently being analyzed as part of a collaboration with the Cabral De Mello Lab from São Paulo University, and we hypothesize that the gene expression of mechanical and regulatory proteins involved in chromosome segregation, such as microtubules, kinetochore proteins, and Bub kinases, will be elevated in response to the presence of supernumerary B chromosomes. We anticipate that our RNAseq analysis will serve as the foundation for new investigations in the genetic pathways that are disrupted due to the presence of B chromosomes during *Drosophila* gametogenesis.

### 728V  Variation in genomic instability due to heat stress in early and late meiosis: Regulation of transcription and chromatin availability

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Changes in the environment affect organisms in several aspects including the cellular, population, and ecosystem levels. At higher levels, populations are affected by changes in fecundity and longevity, whereas at molecular levels organisms can be impacted at the genetic level leading to heritable variation if changes are in the germline. Genomic instability results from the increased stress response in multiple pathways, including during meiosis, regulation of chromatin, and DNA repair. In this study, the primary aim was to investigate mechanisms of genomic instability by assessing the relationships between chromatin availability, DNA damage, and differential gene expression. Two species were used to compare, the model system (*Drosophila melanogaster*, DMEL) to a species (*Drosophila pseudoobscura*, DPSE) that is more vulnerable to changes in environmental temperatures. Species-specific treatment crosses were set up at appropriate temperature ranges (standard vs high temperature), early and late meiotic stages were compared by assessing ovaries dissected from females before and after the first mature eggs developed. RNA from the ovaries was used to assay differential gene expression, pinpointing expression differences during early and late phases of meiotic events of pachytene checkpoint, double-strand break formation, and eggshell chorion assembly. Moreover, the ovaries were used to assay apoptosis to measure the DNA damage using DAPI staining to observe the effect of heat stress. Lastly, nuclei from ovaries were extracted to generate ATAC-seq libraries to investigate changes in chromatin availability in each treatment and time point. Results from this study allow for the comparison between species with different thermal tolerances and provide a mechanistic explanation of the relationship between stress response and genomic instability in the context of meiotic recombination.

### 729V  Active site phosphorylation of CDK11 is antagonised by PNUTS-PP1 and localised in the centrosomes

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Reversible protein phosphorylation is a post-translational modification (PTM) that plays a vital role in the regulation of cellular function and is controlled by protein kinases and phosphatases, which add and remove phosphate respectively. Serine/Threonine Protein Phosphatase 1 (PP1) has multiple and essential roles in cellular processes, including RNA splicing, cell division and metabolism of glycogen (Cohen, 2002). In vivo, PP1 catalytic activity is regulated by PP1 interacting proteins (PIPs), one of the most abundant of which is PNUTS/PP1. We found that PNUTS/PP1 associates with and regulates the phosphorylation state of the protein kinase Pitslre/CDK11. CDK11 becomes hyper-phosphorylated in its active site at S712 (or S538 in a shorter isoform) when binding of PP1 to PNUTS is disrupted by a mutation in the PP1-binding site of PNUTS (W726A). S712 in CDK11 is equivalent to a serine residue in CDK7 that modulates CDK7 activity and binding to...
cyclin H (Larochelle et al., 2001; Martinez et al., 1997) (Endicott & Noble, 2013). Our current focus is therefore to determine when and where CDK11 is phosphorylated. In *Drosophila* neuroblasts we found elevation of phosphorylated CDK11 (pCDK11) during mitosis, and enrichment of pCDK11 staining in centromeres. Overexpression of kinase dead forms of CDK11 resulted in short or aberrant mitotic spindles. To investigate in vivo function further, we generated non-phosphorylatable (S712/538A) and phosphomimic (S712/538E) CDK11 alleles by CRISPR-Cas9 mediated mutagenesis. We found that non-phosphorylatable CDK11 is recessive lethal in early larval stages, indicating that CDK11 phosphorylation is likely to play an essential role. Here we will describe the results of our findings and discuss the possibility that pCDK11 modulates CDK11 activity during the cell cycle to promote mitotic spindle formation or function.

730V  A non-cell-autonomous buffering mechanism protects cells from replication stress-driven DNA damage

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Replication stress is an important driver of genome instability in cancer. It can be triggered by imbalances in the dNTP pool, which is a common strategy of chemotherapeutic drugs. With the aim of understanding the effects of induced replication stress on stem and progenitor cells, we characterized the consequences of depletion of dNTPs in *Drosophila* adult intestinal stem cells and developing wing disc progenitors. Our findings suggest a previously unrecognized buffering mechanism of the dNTP pool from neighboring cells that depends on Gap Junctions. Specifically, the knock-down in intestinal stem cells of *RnRL*, a subunit of Ribonucleotide reductase - a rate-limiting enzyme essential for dNTP production, leads to high levels of DNA damage, S phase arrest, and ultimate depletion of stem cells from the intestine. In contrast, the clonal knockdown of *RnRL* in wing disc progenitor cells leads to little to no DNA damage. However, when Gap Junctions are inactivated along with *Rnrl*, high levels of DNA damage are now induced in the wing disc. This suggests that wing disc progenitor cells participate in dNTP-resource sharing through connections to their neighboring cells via Gap Junctions allowing buffering of induced replication stress. We find that in the intestine, Gap Junctions form between mature enterocytes, whereas intestinal stem cells lack them. Intestinal stem cells, therefore, cannot buffer their dNTP pool via neighboring cells, likely contributing to their exquisite sensitivity to induced replication stress. Overall, our data suggest that a shared dNTP pool can buffer replication stress and that differences in Gap Junction connectivity between cell types may have an important impact on DNA damage susceptibility. These findings have significant implications for understanding replication stress in healthy tissues and cancers.

731V  REV7 Acts Independently of Polymerase ζ to Maintain Genome Stability During Development

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Genome stability is of vital importance during an organism’s life. Without appropriate and timely DNA damage response, development is misregulated or entirely disrupted, leading to unhealthy or non-viable offspring. Recent mammalian work indicates REV7 plays a significant role in DNA damage response. REV7 is a component of Polymerase ζ, used in translesion DNA synthesis (TLS), as well as influences double-strand break (DSB) repair pathway choice as a member of the shieldin complex. REV7, a HORMA domain protein, employs its stereotypical safety belt region to interact with other proteins in these complexes.

As such, we wanted to investigate whether REV7 has functions beyond TLS in *Drosophila*. We used mutant stocks in which the REV7 coding sequence is interrupted by a piggyBac transposable element to characterize phenotypes during normal development and after treatment with exogenous DNA damaging agents. Our studies indicate REV7 loss impairs development to adulthood, decreases hatching rate, and produces abnormal eggshells. The sub-Mendelian ratio of homozygous flies indicates the importance of REV7 in early development. This conclusion is further bolstered by decreased hatching rates, reinforcing the significance of REV7 during development in response to innate endogenous damage. Imperative to successful development is the eggshell, a sophisticated proteinaceous structure composed of chorion, produced en masse via gene amplification. rev7 mutant females produce eggs with partially translucent eggshells. This phenotype has been observed in other DSB repair mutants and is attributed to persistent damage resulting from replication fork collisions during amplification at two chorion loci. Data collected from REV3, Pol ζ’s catalytic subunit, mutants do not echo REV7 loss, suggesting REV7 is acting independently of Pol ζ.

Our findings are consistent with REV7 acting in additional non-TLS roles. We are currently working to identify protein partners that interact with REV7 in support of genome stability. These investigations will provide additional insight into the mechanism(s) by which *Drosophila* REV7 promotes genome stability in the developmental context.

732V  Maintenance of genomic integrity in the male germline of *Drosophila melanogaster*

Kate Lemons, Kent Golic University of Utah
Successful transmission of genetic information from one generation to the next requires germline stem cells to maintain their genomic integrity. Germline cells with DNA damage, such as double strand breaks (DSBs), are largely eliminated. The pathway that works to eliminate cells containing DNA damage has been well studied in the soma. The upstream protein kinase Chk2 phosphorylates P53 which leads to apoptosis. However, in the germline, Chk2 is necessary for elimination of germline cells while P53 is not. This leads to the question: what is downstream of Chk2 in the germline? We used FLP/FRT to efficiently induce DSBs in the male germline. We find that mutation of any of Chk2’s domains, including its kinase function, eliminate its DSB checkpoint function in the germline. String (stg), a positive regulator of cell cycle progression, is a known target of Chk2 in some systems. If Chk2 negatively regulated Stg in response to a DSB, we reasoned that overexpression of stg might bypass the Chk2 effect and mimic the effect of chk2 mutants. We found this to be true, strongly suggesting that Chk2 acts through Stg in the germline to block cell cycle progression, ultimately leading to the elimination of cells with damaged DNA.

733V  **Shared functions of p53 and Xrp1 in DDR and cell competition** Chaitali Khan¹,², Nicholas E Baker¹ 1) Department of Genetics, Albert Einstein College of Medicine, Bronx, NY 10461; 2) National Heart, Lung, and Blood Institute, National Institutes of Health, Bethesda, MD 20892

Cell competition is a process of eliminating relatively unfit cells from chimeric tissues. The tumor suppressor gene p53 is implicated in many instances of cell competition in mammals. There is little evidence for this in *Drosophila*, despite its importance as a model for cell competition. Moreover, mammalian p53 is activated in response to ribosomal defects, whereas *Drosophila* p53 does not seem to be. We hypothesized in *Drosophila* a downstream target of p53 carries out some of its functions and additionally receives inputs from cellular stresses that *Drosophila* p53 does not recognize. This could be the transcription factor Xrp1, a known target of p53, which unlike p53 is activated in response to ribosomal defects and is a key effector of cell competition. Our gene expression analysis shows that Xrp1 is upregulated downstream of p53 in response to acute DNA damage and regulates the expression of molecular players of DNA repair pathway, oxidative response, Jun-kinase signaling and partly the pro-apoptotic gene *hid*. Further, we show that Xrp1 is necessary and sufficient for the expression of several of p53 target genes, but, we find that Xrp1 is only partly needed for DNA damage-induced cell death. Importantly, by activating p53 in a cell non-lethal manner, we show that differences in p53 levels can cause the elimination of cells with higher p53 when growing next to the cells with lower levels of p53. We perform the two-clone assay to confirm that cells with higher p53 are eliminated by cell competition and that this could occurs by activating Xrp1. Interestingly, removing Xrp1 function to show its role in eliminating cells with higher p53 revealed that Xrp1 not only mediate p53 dependent cell competition but is also needed to restrict the clonal expansion of p53 mutant cells. Which we explained by establishing that Xrp1 acts in a p53-independent manner to limit the passage of aneuploid cells created by DNA damage, via its role in cell competition. This further supports the idea that cell competition eliminates aneuploid cells. Overall, these studies establish a role of the p53 pathway in cell competition in *Drosophila* as well as mammals, with Xrp1 as a mediator of acute DDR function, cell competition, and genome stability.

734V  **Loss of rer1-mediated ER-stress drives cell competition in the developing Drosophila wing epithelium** Pranab Kumar Paul, Varun Chaudhary Indian Institute of Science Education and Research (IISER) Bhopal, India

Metazoan development is a robust, error-prone process that is tightly orchestrated by several intrinsic mechanisms. Cell competition is one such process where the defective cell is sensed and eventually eliminated from the tissue. The removal process of defective cells is crucial in the course of healthy tissue development, and several mechanisms of cell competition have been extensively studied to reveal the fact. Using *Drosophila* as a model, many genes have been identified which can provide both selective advantages and disadvantages to the suboptimal cells, but the sensing events caused due to the physiological changes inside an unfit cell before the initiation of elimination are not conserved and still poorly understood.

Here, using the wing imaginal disc of *Drosophila*, we establish that the loss of rer1, an essential regulator of ER homeostasis and Notch pathway activity in higher organisms, can trigger cell competition during the development of the polarised wing discs epithelium. In the absence of Rer1, cells are eliminated through cell death via the upregulation of Unfolded Protein Response (UPR) in ER. Moreover, the elimination of rer1 depleted tissue is blocked by overexpression of the growth arrest and DNA damage-inducible protein (Gadd34), a derepressor of global translation machinery, within the mutant tissue. Unexpectedly, we found that rer1 mutant tissue showed no significant reduction in the level of protein synthesis despite having high levels of P-eIF2α, an integrated stress reporter and repressor of protein synthesis machinery, indicating that P-eIF2α may play a role other than blocking translation machinery in the absence of Rer1. However, if the excessive ER stress or any unknown reason for increased P-eIF2α in rer1 depleted tissue to trigger the elimination of mutant cells is yet to be investigated further.

735V  **Regional differences in timing of apical cell area change associated with sex comb rotation during development** Nicolas Malagon, Anari-Mai Byfield, Eleanor Reimer, Stephen Ingram, David Nelson Canadian Mennonite University

A sex comb is a row of bristles present on the first leg of *Drosophila melanogaster* males. During development, these
bristles rotate from a perpendicular to a parallel position on the long axis of the leg. In previous work, we quantified the changes in apical cell area in 4 regions using 5-cell clusters surrounding the sex comb and found those 4 regions display irregular oscillations in size, increase and decreasing in apical cell area. To understand the cell dynamics underlying these observations we studied the entire region located above the comb (30 cells) during the period between 23 and 36 hours after pupation (AP). Using timelapse videos of confocal micrographs of the development of the forelegs of ubiDE-Cadherin::GFP flies, we visually determined the apical area by tracing the cells in ImageJ. We observed a reduction in variation in apical cell area associated with comb rotation. However, this uniformity in size is achieved following highly variable local dynamics in apical cell size. This work suggest that irregular oscillations of apical size may be due to local mechanical conditions which underly how overall change in tissue size can occur in a gradual manner.

736C  The effects of developmental ethanol exposure on markers of aging in Drosophila melanogaster Navneet Sanghera, Khaoula Belhorma, Rachael French San Jose State University

Fetal Alcohol Spectrum Disorder (FASD) results from consumption of alcohol during pregnancy, and can lead to a variety of effects including developmental delays, low body weight, and intellectual disabilities. We have established a Drosophila model for FASD, and have demonstrated that flies exposed to ethanol during larval development exhibit phenotypes very similar to those seen in mammals. In addition, we recently found that flies reared in ethanol-containing media showed reduced age-related decline in negative geotaxis. In addition, we found persistent ethanol-induced upregulation of the antioxidant genes Sod1 and Gss1 in adult flies, continuing for at least 5 days after removal from ethanol-containing food.

Because age-related declines in negative geotaxis are associated with increased levels of oxidative damage to the central nervous system, we hypothesize that increased expression of cellular antioxidants as a result of ethanol exposure during development leads to a decreased age of aging. To test whether this is a general effect on aging, rather than a specific effect on climbing ability, we are performing several sets of experiments to test the effects of developmental ethanol exposure on cellular and organismal aging. We will present the results of these experiments, including tests of fecundity over time, mortality and lifespan assays, and cellular measurements of the accumulation of oxidative damage.

737A  Live longer, climb further: Parabacteroides distasonis promotes healthy aging and gut barrier integrity in Drosophila melanogaster. Luana Machado1, Sofia Rosa1, Sarah Shnayder1, Jim Crott1, Mitch McVey1 1) Tufts University, Medford, MA; 2) Human Nutrition Research Center on Aging, Boston, MA

The term “inflammaging” has been accepted as an all-encompassing term for the age-related increase in inflammation that occurs in organisms. Inflammaging causes many negative impacts on health and lifespan and is associated with cancer progression. Interestingly, some bacteria have been shown to have anti-inflammatory properties. One example is the obligate anaerobe Parabacteroides distasonis (Pd), which was recently shown to decrease cancer incidence and progression in a mouse colon cancer model through the modulation of systemic inflammation.

Because inflammation has been linked to aging and decreased health, we explored the effects of Pd treatment on lifespan and healthspan in a Drosophila model system. We found that chronic administration of Pd in the food of adult flies extends the mean lifespan of two genetically distinct strains by 19% and 28% in males and females respectively. In addition, consumption of Pd promotes the maintenance of climbing ability in a negative geotaxis assay, suggesting that Pd also promotes healthy aging.

Having established the impact of Pd on lifespan and healthspan, we began our search for the molecular mechanism through which this bacteria is acting. We first explored its effect on gut membrane stability through Smurf assays, a qualitative approach to examining gut leakage by analyzing how much blue dye diffuses from the intestinal tract into the abdomen of the flies. By comparing dye leakage among treatment groups, we observed that Pd reduces gut barrier defects and increases gut stability in flies as they age.

Given these findings, we hypothesized that Pd could maintain gut membrane integrity in flies by impacting the expression and localization of smooth septate junction proteins (tight junction protein homologs) in the gut, decreasing systemic inflammation, or indirectly preventing hyperproliferation of intestinal stem cells. We conducted q-RT-PCR on candidate genes related to these processes. Our results suggest that expression of inflammatory markers and proteins required for smooth septate junction formation increases with age, but their levels are not affected by treatment with Pd. Analysis using RNA-Seq is ongoing. We are also using fluorescent imaging to visualize potential physical differences among gut tissue of varying treatment groups, specifically looking at stem cell proliferation patterns and smooth septate junction formation. Identification of conserved molecular pathways through which Pd exerts its effects could have therapeutic applications in humans.

738B  Developing a quantitative analysis of cysteine availability via iodoTMT-multiplex method using Drosophila S2 cells and w1118 eyes. Sarah Stanhope Purdue University
Reactive oxygen species (ROS) serve as intracellular signaling molecules; however, in excess ROS molecules are damaging to biomacromolecules such as DNA, lipids, and proteins. The excess accumulation of ROS leads to oxidative stress, disrupting redox homeostasis in the cell and has been linked with aging and neurodegenerative disease. The eye is particularly vulnerable to oxidative stress due to the tissue's high energy demand and generation of ROS by products. In proteins, the thiol group on cysteine residues is susceptible to oxidation. Cysteine residues have multiple oxidation states that can be classified into two categories—reversible and irreversible. Irreversible oxidation states include sulfenic and sulfonic acids, which may alter or impair the activity of the protein depending on the position of the cysteine residue. Understanding how the proteome is affected by increasing ROS levels may provide insights into proteins that are potential targets for oxidation. Here, we sought to evaluate how increasing amounts of oxidative stress altered the redox proteome landscape in Drosophila. To characterize the redox proteome, we developed an iodoTMT-multiplex isolation, labeling, and enrichment method to identify cysteine availability and potential oxidation in both S2 cells and w1118 fly eyes. To do this we exposed S2 cells to 20mM H2O2 and w1118 flies to prolonged blue light, followed by redox proteome profiling with iodoTMT-sixplex reagents. S2 cell studies revealed 127 significantly oxidized proteins including multiple well-known oxidative stress proteins such as Gluthionine S-transferase D1 (GstD1) and Superoxide dismutase [Cu-Zn] (Sod1). GO term analysis elucidated proteins involved in cellular metabolic processes, sulfur compound processes, and cofactor metabolic processes. Further studies in blue light exposed w1118 eyes revealed 42 significantly oxidized proteins involved in nucleoside metabolic processes, regulation of cytosolic calcium ion concentration, and glycosyl compound metabolic processes. Interestingly, the overlap of the identified proteins from the S2 cell and w1118 studies revealed oxidation of methionine metabolism enzymes suggesting that oxidative stress may perturb methionine metabolism affecting metabolic processes within the cell. As methionine metabolism is critical for lifespan across various organisms understanding how it can be affected by increasing oxidative stress and its role in the eye is critical.

739C Drosophila STING protein has a role in lipid metabolism Katarina Akhmetova, Maxim Balasov, Igor Chesnokov University of Alabama at Birmingham

STimulator of IINterferon Genes (STING) is an endoplasmic reticulum-associated transmembrane protein that plays an important role in innate immune response by controlling the transcription of many host defense genes. STING has been extensively studied in mammalian immune response, however, the role of STING in the innate immunity of insects have been just recently identified. Immune system is tightly linked with metabolic regulation in all animals, and proper re-distribution of the energy is crucial during immune challenges. Insect fat body integrates these two important systems: it serves as a major immune organ, and also stores excess nutrients (mostly in the form of lipids) and mobilizes them during metabolic shifts. In this work we describe a novel function of Drosophila STING (dSTING) in lipid metabolism. We discovered that flies with a deletion of dSTING are sensitive to the starvation and oxidative stress. Detailed analysis revealed that dSTING deletion resulted in a significant decrease in the main storage metabolites, such as TAG, trehalose and glycogen. We identified two fatty-acid biosynthesis enzymes - Acetyl-CoA carboxylase (ACC) and fatty acid synthase (FASN) - as the interacting partners for dSTING. Moreover, we found that FASN and ACC interacted with each other, indicating that all three proteins might be components of a large complex. Importantly, dSTING deletion leads to the decreased FASN activity and defects in ACC cellular localization suggesting a direct role of dSTING in lipid metabolism of fruit flies.

740A The steroid hormone ecdysone regulates growth rate in response to oxygen availability George Kapali1, Viviane Callier2, Samuel Gascoigne3, Jon Harrison4, Alexander Shingleton1,3 1) University of Illinois at Chicago, Chicago, IL; 2) Arizona State University, Tempe, AZ; 3) Lake Forest College, Lake Forest, IL

In almost all animals, physiologically low oxygen (hypoxia) during development slows growth and reduces adult body size. The developmental mechanisms that determine growth under hypoxic conditions are, however, poorly understood. Here we show that the growth and body size response to moderate hypoxia (10% O2) in Drosophila melanogaster is systemically regulated via the steroid hormone ecdysone. Levels of circulating ecdysone are elevated under hypoxia, and inhibition of ecdysone synthesis ameliorates the negative effect of low oxygen on growth rate. We also show that the effect of ecdysone on growth rate under hypoxic conditions is mediated by the insulin/IGF-signaling (IIS) pathway. Hypoxia reduces systemic IIS activity and the hypoxic growth-response is eliminated in larvae with suppressed IIS. Critically, hypoxia transiently increases the expression of Imp-L2, an ecdysone-response gene that suppresses systemic IIS. Inhibition of ecdysone synthesis eliminates this increase in Imp-L2 expression, while loss of Imp-L2 significantly reduces the negative effect of hypoxia on final body size. Collectively, these data indicate that growth suppression in hypoxic Drosophila larvae is accomplished by systemic endocrine mechanisms that overlap with mechanisms that regulate growth in response to low nutrition. This suggests the existence of growth-regulatory stress response mechanisms that respond to general environmental perturbation rather than individual environmental factors.

741B Beauty of adenosine and immune system metabolism Pavla Nedbalová, Lenka Chodáková, Nikol Kaislerová, Tomáš Doležal University of South Bohemia in České Budějovice

Immune system activation is an energy demanding process requiring metabolic changes on the organismal level.
to ensure enough energy for immune cells to successfully face the pathogen. Our laboratory is interested in immunometabolism and uses *Drosophila melanogaster* larvae infected with parasitoid wasp *Leptopilina boulardi* to induce immune system activation and examine both, systemic metabolic changes caused by activated immunity and also metabolism changes of the immune cells themselves. We have shown that larval immune cells actively produce extracellular adenosine as a systemic metabolism regulator upon infection. Extracellular adenosine is inhibiting glucose uptake by non-immune tissues such as imaginal discs. Thus, there is a high level of glucose available for activated hemocytes. The question is, what is the metabolic origin of this extracellular adenosine in hemocytes?

It is known in mammals that S-adenosylmethionine (SAM) cycle accelerates in activated immune cells. This metabolic pathway is responsible for the methylation of diverse biomolecules and this reaction produces S-adenosylhomocysteine (SAH), a potent inhibitor of this pathway, which is immediately converted to adenosine and homocysteine by S-adenosylhomocysteine hydrolase (Ahcy). Therefore, we have considered Ahcy and SAM cycle as a possible source of extracellular adenosine and we have already obtained data confirming this hypothesis and showing that SAM cycle activity increases also in activated *D. melanogaster* hemocytes.

Our data further imply that adenosine is not only released from hemocytes but is also highly used intracellularly. The first step of the SAM cycle is a unique reaction in which the adenosyl moiety of ATP is consumed along with methionine to synthesize SAM. We propose that a part of the adenosine pool produced by Ahcy is recycled back to ATP through the cooperation of adenosine kinase, adenylate kinase, and glycolysis, as losing all of the adenosine would be a huge energy wasting since all used ATP would have to be refilled by de novo purine synthesis. Our current effort is to examine this complex metabolic network showing adenosine as a crucial molecule affecting not only systemic energetic metabolism and supporting immune response but also participating in the intracellular energy balance of activated immune cells.

**742C  The Drosophila gene sima is an essential regulator of the larval glycolytic program** Yasaman Heidarian, Liam Mungcal, Jason Tennessen Indiana University

The rapid growth that occurs during *Drosophila* larvae requires a dramatic rewiring of central carbon metabolism to support biosynthesis. Previous studies in our lab demonstrated that this exponential growth phase is preceded by a dramatic metabolic switch, which results in the transcriptional up-regulation of genes involved in carbohydrate metabolism, including those that encode enzymes involved in glycolysis and the pentose phosphate pathway as well as Lactate Dehydrogenase. The resulting metabolic program exhibits the hallmark characteristics of aerobic glycolysis and establishes a physiological state that supports biomass accumulation similar to that seen in tumor cells. Studies in the fly have so far only discovered a single transcription factor involved in this process - the *Drosophila* Estrogen-Related Receptor (dERR), which is absolutely required to up-regulate carbohydrate metabolism in preparation for larval growth.

Here we describe our discovery that Sima, the sole *Drosophila* ortholog of the hypoxia inducible factor-1 alpha (Hif1-a), is also essential for promoting aerobic glycolysis in larvae. Using CRISPR/CAS9 generated sima alleles that disrupt an exon shared among all isoforms, we discovered that the classic sima p-element insertion mutation sima^{4070/07} represents a weak hypomorph that exhibits very mild metabolic defects. In contrast to this canonical allele, our novel null mutations exhibit a mid-L2 lethal phase and phenocopy the metabolic defects observed in dERR mutants. Moreover, transcriptomic analysis of sima mutants reveal a dramatic downregulation in glycolytic genes, with the overall transcriptional profile mimicking that of the dERR mutant. Subsequent metabolomics studies revealed that, when compared to control larvae, sima trans-heterozygous mutants exhibit >90% decrease in lactate and 2-HG and a two-fold increase in the circulating disaccharide trehalose. Altogether, these preliminary studies indicate that sima, like dERR, is an essential regulator of the larval metabolic program – a result supported by previous findings that Sima can physically bind dERR. Our future studies will examine the interaction between Sima and dERR to understand how these two evolutionarily conserved transcription factors control aerobic glycolysis in the context of both normal animal development and cancer metabolism.

**743A  Investigating the mechanism of the pro-aging effects of blue light in Drosophila** Jun Yang, Yujuan Song, Kelsey Shimoda, Benjamin Ramsell, David Hendrix, Jadwiga Giebultowicz Oregon State University

Blue light (BL) is increasingly present in the human environment since BL-enriched LEDs became one of the most common and inexpensive light sources in our homes and workspaces. Blue light is known to damage retina-related cells but also decrease the lifespan in the nematode, *C. elegans*. Our recent data showed that daily exposure of fruit fly (*Drosophila melanogaster*) to 12-h of blue light per day accelerates their aging process and shortens their lifespan. The lifespan of flies kept in constant blue light decreases even more significantly – a quarter of the lifespan for dark control (20 days maximum). To understand the mechanism of blue light damage, we transferred flies kept in constant blue light for 6-, 10-, 14-, and 16-days to a constant-dark environment. Mortality curves suggest the blue light damage could be reversed after 14- but not 16-days of BL-exposure. Next, we measured the gene expression with RNA-seq sampled from heads of 6-, 10-, and 14-day-BL-exposed and age-matched dark control flies. DESeq2 gene expression analysis revealed extensive transcriptome re-programming in response to blue light. Over thousands of genes were significantly altered with several genes strongly upregulated, including transcription factor *Xrp1*, small heat shock protein *I(2)"
and Arc1. Differentially expressed genes were enriched for functional groups such as neuropeptides, immunity, and metabolism. To further study the metabolic processes affected by blue light, we performed metabolomics using LC-MS and GC-MS. We detected 216 metabolites in the heads of 10- and 14-day-BL-exposed and control flies. MetaboAnalyst identified 42 compounds significantly altered by blue light. Pathway analysis showed that blue light most significantly impacted compounds in the tricarboxylic acid cycle (TCA), Riboflavin, Propanoate, and Butanoate metabolism. Levels of 4 metabolites in the TCA were reduced while levels of succinate were significantly increased in blue light. Moreover, we also determined that succinate dehydrogenase (SDH) activity was reduced in BL-exposed flies, which suggests the impairment of the electron transport chain in mitochondria. In addition, riboflavin, the precursor of SDH cofactor FAD, was significantly reduced by blue light. Lastly, ADP levels were significantly increased in BL-exposed flies while ATP was decreased. These results show that TCA and mitochondria are impaired in blue light. Another metabolite significantly reduced in BL-exposed flies is glutamate, which serves as an energy substrate and also functions as an excitatory neurotransmitter, and a precursor for the inhibitory neurotransmitter GABA. Taken together, our data suggest that blue light impairs energy metabolism and interferes with brain functions in flies.

**744B Nutrient-dependent acyl-CoA metabolism regulates tissue remodeling by adjusting stem cell quiescence and activation in Drosophila**

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Nutrient availability is a major selective force in the evolution of metazoa, and organisms thus need to carefully adapt to nutrient changes in order to promote homeostatic maintenance and reproduction. This adaptation has also shaped the cellular and molecular mechanisms that underly nutrient-dependent remodeling of tissue function and morphology. Here, we leveraged a meta-analysis of transcriptomics datasets and phylogenomics and discovered that Acbp6 (Acyl-CoA binding protein 6) is a critical regulator of nutrient-dependent tissue remodeling in Drosophila. Acbp6 arose by evolutionary duplication in the Drosophila order, and is uniquely expressed in functionally differentiated enterocytes of the regenerating fly midgut (intestine). We found that Acbp6, which binds Acyl-CoA with high-affinity and thus dictates cellular Acetyl-CoA metabolism, is required for intestinal regrowth after nutrient deprivation (fasting) through adjusting intestinal stem cell (ISC) activation from quiescence. Fasting induces intestine morphological recession, while refeeding promotes intestinal remodeling and growth in Drosophila. Acbp6 expression is induced by fasting, and normalizes after refeeding. However, attenuation of Acbp6 led to abnormal tissue remodeling after fasting, due to uncoordinated regulation of ISC activation and proliferation. Moreover, Acbp6 inhibition also drives aberrant ‘switching’ of metabolic cycles in enterocytes during nutrient adaptation, which overall impaired Acetyl-CoA production. Decreases in Acetyl-CoA led to attenuation of pan-acetylation of proteins in enterocytes during refeeding and intestinal regrowth. Re-expression of Acbp6 or acetate supplementation can restore pan-acetylation and promote ISC activation from quiescence during nutrient adaptation. Through genetic screening we identified Stat92e, which can be acetylated, as a key regulator of ISC activation from quiescence during refeeding. In enterocytes, Stat92e can govern Upd3 induction during nutrient adaptation, and Upd3 can dictate ISC proliferation and intestinal regrowth. Our data thus define a new regulatory mechanism, shaped by Acyl-CoA metabolism, that is required to adjust stem cell quiescence and activation, and therefore drive tissue remodeling during nutrient adaptation.

**745C Endocrine signals from the gut that regulate metabolism**

**Nadja Ahrentlov, Stanislav Nagy, Olga Kubrak, Alina Malita, Takashi Koyama, Michael Texada, Kim Rewitz**

Section for Cell and Neurobiology, Department of Biology, Copenhagen University, Denmark

Maintaining metabolic homeostasis requires interorgan coordination and thus communication, mediated by endocrine signals. Organs with specialized functions sense changes in internal and external nutrient availability and release hormonal factors that coordinate energy intake, storage, and expenditure. In both flies and mammals, the gut, the largest endocrine organ and the first that encounters food after ingestion, is a source of many secreted signals that regulate metabolism. Human gut-derived hormones such as glucagon-like peptide 1 (GLP-1), which regulates both insulin secretion and underlies the “incretin effect,” in which ingested glucose induces a larger insulin release than glucose injected directly into the bloodstream, thus bypassing the gut. Insulin action is opposed by the peptide hormone glucagon, but gut hormones that regulate glucagon release are poorly defined.

Many proteins are predicted to be secreted, but the functions of most of these potential hormones are largely unknown; these factors thus represent a significant unexploited resource for the discovery of new endocrine signals from the intestine that affect metabolic balance. Like its mammalian analog, the small intestine, the Drosophila midgut contains enteroendocrine cells (EECs) that release a variety of known endocrine factors in response to nutritional quantity and quality. We have performed a large-scale EEC-specific in-vivo RNAi-based screen of secreted factors in adult flies with the goal of identifying gut-derived factors that regulate food intake and metabolism. Our phenotypic readouts included measures of male and female flies’ ability to survive during nutritional deprivation and their storage of energy (as lipids and glycogen). Among our hits we have found several gut-derived hormonal factors that play key roles in regulating metabolism.
metabolism and feeding behavior, some of which act in a sexually dimorphic manner by modulating the glucagon analog Adipokinetic hormone. Taken together our results identify several gut hormones that may be of relevance to the treatment of diabetes and obesity.

746A  The loss of function mutation in the *Drosophila Neprilysin Like 15* changes expression of key enzymes involved in glycogen homeostasis, and effects longevity in sex specific manner, but exerts similar effects on motor activity in both sexes Nicolas Jones, Raghad Almotairy, Thanh Ha Vy Nguyen, Hannah Halmes, Erin Hatcher, Brianna Villines, Surya Banerjee Arkansas Tech University

Obesity predisposes a person to an array of life-threatening metabolic disorders including diabetes, high blood pressure, heart attack, and cancer. The long-term treatment for such diseases is often less effective in providing a complete cure, and the cost for it creates excessive economic burden. Thus, new and more effective therapeutic approaches are under review. In the last two decades, Neprilysins (Neps), the membrane bound metallo-ectopeptidases, have become targets to treat obesity and related disorders for their correlation with high fat and high sugar diet induced obesity and diabetes. Thus, the mechanisms by which Neps regulates obesity need to be explored. In our recent published work, we show that knock-out of the *Drosophila* specific gene *Neprilysin like 15 (Nepl15ko)*, which has overall higher expression in the wild type adult male compared to female flies, results in significant reduction of stored glycogen and lipids only in adult males, but increases glycogen in females. Although, the mutant flies show similar food intake compared to the controls. To further dissect the effects on the expression of the key enzymes in glycogen and lipid homeostasis pathway, we recently find that *Nepl15ko* adult males but not females have significant reduction in the expression of Glycogen Synthase and Glycogen Phosphorylase transcripts compared to controls. In addition, we have performed longevity and climbing assay to investigate the effect of Nepl15 loss of function on overall health and physiological motor activity. We reveal that the *Nepl15ko* adult males have similar life span, but *Nepl15ko* females show significantly extended life span compared to the controls. Both *Nepl15ko* adult males and females flies show enhanced ability of climbing in an age dependent manner compared to the controls. For example, whereas there is no difference in climbing ability between the mutant and control males and females up to 30 days of age, the mutant flies of both sexes exhibit significantly higher ability for climbing at the age of 40 days. Thus, our research reveals that mutation in the *Nepl15* provides health benefits in females, and keeps the physical activeness for longer in both sexes due to changes in nutrient homeostasis. In the future, we will investigate the lipid homeostasis pathway and energy metabolism in these flies. This project is funded by Arkansas NIH-INBRE grant.

747B  Two phases of ageing in mice, a mammal model for Smurfness. Celine Cansell1, Fanny Bain2, Vivien Goepp7, Nicolas Todd3, Veronique Douard4, Flaminia Zane5, Clara Sanchez6, Nicolas Pietrancosta6, Carole Rovere5, Raphael GP Denis5,9, Serge Luquet3, Michael Rera1 1) Center for Research and Interdisciplinarity (CRI), Inserm U1284, Université de Paris, F-75006 Paris, France; 2) MinesParisTech, CBIO – Centre for Computational Biology, PSL Research University, 75006, Paris, France & Institut Curie, PSL Research University, 75005, Paris, France & Inserm, U900, Paris; 3) Centre Roland Mousnier, CNRS, Sorbonne Université, 75005 Paris, France & Laboratory of Population Health, Max Planck Institute for Demographic Research, 18057 Rostock, Germany; 4) Université Paris-Saclay, INRAE, AgroParisTech, MICALIS Institute, 78350 Jouy-en-Josas, France; 5) Université Côte d’Azur, IPMC-CNRS UMR 7275, F-06560 Valbonne, France; 6) Laboratoire des Biomolécules, LBM, Département de chimie, École Normale Supérieure, PSL University, Sorbonne Université, CNRS, Paris 75005, France; 7) Neuroscience Paris Seine-Institut de Biologie Paris Seine (NPS-IBPS) INSERM, CNRS, Sorbonne Université, Paris 75005, France; 8) Université de Paris, Institut Cochin, INSERM, CNRS, F-75014 PARIS, France; 9) Université de Paris, BFA, UMR 8251, CNRS, F-75013 Paris, France

The “Smurf” model of ageing, first described in Drosophila, is characterized by two successive and necessary phases. A first phase A where individuals have no risk of mortality but an age-dependent increasing risk of entering phase B. A second phase B where individuals show a high risk of impending death from natural causes. Individuals in phase B show typical hallmarks of ageing such as decreased energy storage, locomotor activity and fertility, a deregulation of insulin signaling, and a strong increase in systemic inflammation. The discovery of this model was made possible by a singular characteristic of phase B, its increased intestinal permeability. To assess intestinal permeability in Drosophila, a non-absorbable and non-toxic blue food dye is added to the medium of adult individuals. In contrast to phase A, for individuals in phase B, the dye crosses the intestinal epithelium and reaches the hemolymph turning individuals blue. According to their color, blue individuals were hence dubbed “Smurfs”.

We propose here for the first time that the two phases of ageing separated by the “Smurf” transition are relevant to mice, thanks to a longitudinal longevity study using two different mouse strains, in both males and females. By integrating physiological, metabolic and molecular measurements with the life history, we assessed the predictive power of the model concerning the high risk of impending death in this model organism. This allows us to answer three research questions:

1. As described in Drosophila, is the end of life in mammals characterized by an increase in intestinal permeability?
2. Is there a specific physiological signature of the end of life in mammals?
3. Do identified biomarkers allow us to discriminate between two subpopulations characterized by different mortality
risk at any chronological age? The characterization of this “Smurf” two-phase ageing model in mammals is essential to study and to better understand ageing because it allows a primordial thing currently still impossible: to evaluate the physiological age of individuals. It also allows a better understanding of the underlying mechanisms of ageing by offering the possibility to exclude individuals presenting the classically accepted hallmarks of ageing in order to define the causes of the appearance of these hallmarks. Finally, it is a putative game changer in the domain of personalized medicine paving the way for a unifying and truly public model of ageing, while extending the Drosophila model to mammals.

**748C Investigating Flock House virus-mediated changes in bioenergetics in aged Drosophila melanogaster**

Dean Bunnell, Eli Hagedorn, Beate Henschel, Daniel L. Smith, Stephanie Dickinson, Andrew W. Brown, Maria De Luca, Ashley Turner, Stanislava Chtarbanova 1) University of Alabama; 2) Indiana University; 3) University of Alabama at Birmingham; 4) Jacksonville State University

We have previously shown that Flock House Virus (FHV) kills older flies more rapidly than young adults without higher virus titers. This indicates that survival to FHV infection depends on tolerance mechanisms, which are not well characterized. FHV triggers profound transcriptomic changes in older Drosophila at time points before significant mortality is observed in young and aged cohorts. The age-dependent impairment of disease tolerance to FHV in Drosophila is characterized by stronger regulation of genes whose products function in mitochondria and mitochondrial respiration. We hypothesized that the Drosophila bioenergetic profile will be deregulated following FHV infection. The first 3 days of infection, we performed whole-organism respirometry of 5- and 30-days old individual Drosophila receiving non-injected, Tris-injected (control) or FHV-injected treatments to determine if metabolism plays a role in infection outcomes. Pairwise comparisons were conducted on the Least Squares Mean Oxygen Consumption Rate (OCR) to detect statistically significant differences between treatment groups. Our results show that FHV infection significantly reduces OCR compared to non-injected (p=0.0264) and Tris-injected (p=0.0164) controls. Although the factor of «Age» and the interaction of «Age x Treatment» did not have a significant effect on OCR, the interaction of «Age x Time Post-Treatment» shows a significant change in OCR in aged flies compared to young individuals at 24-hours. The OCR signature at 24-hours varied in response to both treatment and an individual’s survival status at 48-hours. FHV-injected flies that died prior to 48-hour measurements had a significantly lower OCR (p=0.0438) compared to flies that were alive at 48-hours.

**749A Coordinated shifts in redox metabolites during quiescence are heritable factors that reprogram progeny metabolism**

Helin Hocaoglu, Lei Wang, Mengye Yang, Sibiao Yue, Matthew Sieber UT Southwestern Medical Center, Dallas, TX

Maternal diet and metabolic stress have a profound effect on health and disease susceptibility of progeny. Previous studies have shown this transgenerational effect; however, the mechanisms remain unknown because of the challenges in obtaining pure populations of quiescent oocytes. To overcome this challenge, we utilized the Drosophila oogenesis system to isolate large quantities of staged oocytes to study how the quiescent nature of oocytes impacts the effect of maternal metabolic stress on progeny metabolism. Using biochemical and systems-based approaches, we found that in the late stages of oogenesis, oocytes acquire a unique redox state due to suppression of mitochondrial oxidative metabolism in a process called mitochondria respiratory quiescence (MRQ). We have found that maternal metabolic stress triggers MRQ prematurely and induces the reprogramming of progeny metabolism and life-long changes in glucose and triglyceride homeostasis. In addition, we have found that maternal metabolic stress causes a disruption in the unique redox state of oocytes by impairing NAD biosynthesis. We have shown that reducing the levels of NAD in the oocyte are sufficient to induce progeny reprogramming, moreover maternal NAD precursor supplementation can suppress the progeny phenotypes caused by maternal metabolic stress. Compromised NAD levels inherited from the oocyte causes impaired methionine cycle activity and a 50% decrease in the production of the methyl-donor S-adenosyl methionine (SAM). Lower levels of SAM lead to a global 30% reduction in H3K27-me3, a corresponding increase in H3K27-ac, and de-repression of 550 genes during embryogenesis. Interestingly, 1/3 of these genes are specifically expressed in the intestine and are involved in processes such as lipid digestion, uptake, storage, protein digestion, nutrient transport, and calcium signaling. As a result of this intestinal metabolic shift, reprogrammed progeny were able to develop better on a low nutrient diet but have a shorter adult life span compared to control progeny. Taken together we believe that progeny metabolic reprogramming is an adaptive tradeoff that tunes progeny metabolism to promote development in a poor nutrient environment, at the expense of a shorter lifespan.

**750B Experimental Evolution to identify genes that contribute to fitness in high-sugar-fed Drosophila melanogaster**

Thomas Rundell, Azva Alvi, Melina Brunelli, Christina Capobianco, Gabrielle Safian, Laura Musselman Binghamton University

Evolve-and-resequence approaches have been shown to dramatically alter both phenotypic and genotypic characteristics of populations under laboratory selective pressures. In this project, an outbred population made from wild-caught Drosophila was subjected to a control or high-sugar (HS) feeding paradigm for many generations. HS feeding reduces
both the lifespan and healthspan in adult Drosophila. Sexes were separated and aged on either diet until mid-life, then mated to produce the next generation, allowing selection for protective alleles. Alleles that increase survival, metabolic homeostasis, and fecundity are hypothesized to be favored under the HS diet. We found that all selected populations increased their lifespan and healthspan over time. Four control and four HS-selected populations have been compared using pooled DNA sequencing coupled with RNA sequencing to identify specific, enriched loci that may have conferred protection against the negative sequelae of caloric excess. One cohort of genes identified across multiple HS-selected populations contained acetylcholine related genes including ChAT, CHT, and mAChR-m. We will use genetic approaches to test whether these genes mitigate overnutrition phenotypes, potentially revealing a novel link between the central nervous system and metabolism.

751C Lactate and glycerol-3-phosphate metabolism cooperatively regulate larval growth in a tissue nonautonomous manner Madhulika Rai, Hongde Li, Sarah Carter, Maria Sterrett, Geetanjali Chawla, Jason M. Tennessen Indiana University Bloomington

The dramatic growth that occurs during *Drosophila* larval development requires the rapid conversion of nutrients into biomass. In response to these biosynthetic demands, larval metabolism exhibits the hallmark features of aerobic glycolysis, a metabolic program ideally suited to synthesize macromolecules from carbohydrates. Central to the biosynthetic potential of aerobic glycolysis is lactate dehydrogenase (LDH), which promotes glycolytic flux by regenerating NAD\(^+\). We have seen that although Ldh mutants accumulate elevated NADH levels, larvae compensate for this metabolic insult by increasing glycerol-3-phosphate (G3P) production, which serves as a backup mechanism to regenerate NAD\(^+\), and the cooperative regulation of lactate and G3P metabolism imparts metabolic robustness on the larval glycolytic program. Further, lack of Ldh and Gpdh1 together, exhibit developmental delays, synthetic lethality, and aberrant carbohydrate metabolism. Although we understand the effect of the loss of Ldh and Gpdh1 in the whole body, tissue-specific roles of both enzymes remain unexplored. To address this deficiency, we used RNAi to understand how tissue-specific depletion of Ldh and Gpdh1 affects larval growth and metabolism. Our results demonstrate that while individual loss of either Ldh or Gpdh1 in fat body, muscle and neurons does not affect larval development, loss of both Ldh and Gpdh1 within either the fat body, muscle or neurons leads to systemic growth defects in larvae. Hence, Ldh and Gpdh1 can influence larval growth and metabolism in a cell nonautonomous manner, indicating that the cooperative activity of these two enzymes within individual tissues is capable of inducing systemic signals that coordinate intercellular metabolic states with growth of the entire organism. Finally, our preliminary findings indicate the cytokine Upd3 is a key signal in connecting loss of Ldh/Gpdh1 activity in individual tissues with systemic growth delays. Overall, our findings hint at a mechanism that coordinates larval growth with the rate of glycolytic flux in individual tissues.

752A Investigating the role of Glycerol-3-phosphate dehydrogenase 1 (GPDH1) in *Drosophila* growth and development Shefali Shefali, Madhulika Rai, Sarah Carter, Nader Mahmoudzadeh, Hongde Li, Maria Sterrett, Jason Tennessen Department of Biology, Indiana University Bloomington, Bloomington, IN, USA

As the fruit fly, *Drosophila* melanogaster, progresses from one life stage to the next, many of the enzymes that compose intermediary metabolism undergo substantial changes in both expression and activity. These predictable shifts in metabolic flux allows the fly to meet stage-specific requirements for energy production and biosynthesis. In this regard, the enzyme Glycerol-3-phosphate dehydrogenase (GPDH1) was the focus of biochemical genetics studies for several decades, and as a result, is one of the most well characterized enzymes within *Drosophila* metabolism. Among the findings of these earlier studies is that GPDH1 promotes mitochondrial energy production and triglyceride accumulation while also serving a key role in maintaining cellular redox balance. Moreover, GPDH1 activity is regulated in a stage-specific manner, suggesting that this enzyme plays an essential role in coordinating metabolism with developmental progression. Here we extend these earlier observations using metabolomics to expand upon these earlier observations. Our studies reveal that Gpdh1 mutants exhibit severe defects in amino acid metabolism, with both the maternal and zygotic GPDH1 enzyme pools influencing amino acid abundance. Moreover, our preliminary studies indicate that GPDH1 indirectly regulates growth signaling in larval tissues by controlling abundance of dihydroxyacetone phosphate. Overall, our results reiterate the importance of GPDH1 activity in larval development and reveal how this enzyme can control growth by previously unappreciated mechanisms.

753B Ribosomal profiling Reveals Changes in the Translatome of *kdm5*-Knockdown Neurons Matanel Yheskel, Julie Secombe Albert Einstein College of Medicine

KDM5 is a histone lysine demethylase responsible for the demethylation of H3K4me3, a histone modification associated with transcriptionally active promoters. Mutations in three of the four KDM5 paralogs in humans (KDM5A-C) are associated with intellectual disability but KDM5’s role in regulating neuronal function is unknown. RNA-Sequencing of neuron-specific *kdm5* knockout flies reveals that the majority of ribosome protein genes (RPG) are downregulated compared to control, however there is no dramatic change to bulk translation or translation initiation. Changes in RP stoichiometry has been shown to affect translation efficiency (TE) of different classes of mRNA; to that end, we performed ribosomal profiling in *kdm5* knockdown neurons. This analysis revealed that mRNAs encoding proteins
with roles in glycolysis, the tri-carboxylic acid (TCA) cycle, and fatty acid metabolism showed significantly reduced TE in kdm5 knockdown neurons. Interestingly, this included genes whose human homologs are implicated in neurological disorders. These data reveal a novel role of KDM5 in the translational regulation of key metabolic pathways necessary for proper neuronal homeostasis and function.

754C  Investigating the mechanisms that control glycolytic gene expression at the cessation of larval growth  Tess Fasteen, Jason Tennessen Indiana University

All growth during the *Drosophila* life cycle is restricted to larval development, when animals increase their body size ~200-fold over the course of four days. In order to support this exponential growth rate, larvae up-regulate glycolytic metabolism as a means to generate the necessary energy and biomass. The resulting metabolic program exhibits the hallmark characteristics of the Warburg effect, or aerobic glycolysis, which is also used by many types of cancer cells to fuel tumor growth. Thus, our ability to study the mechanism that controls larval glycolytic metabolism provides a powerful genetic model to explore how aerobic glycolysis is regulated *in vivo*. In this regard, the fly provides an unusual opportunity to identify the endogenous mechanisms that turn off aerobic glycolysis *in vivo* - unlike tumors, where the Warburg effect is indefinitely activated, fly larvae down-regulate glycolysis at the end of the growth phase in a predictable manner. Using a candidate gene approach, we are exploring two mechanisms that potentially control the global down-regulation of glycolytic gene expression at the onset of metamorphosis. First, we are exploring the possibility that the late-larval pulse of 20-hydroxyecdysone (20E) is necessary to downregulate glycolytic gene expression. Our preliminary studies support this hypothesis, as two metabolite markers of aerobic glycolysis, lactate and 2-hydroxyglutarate, fail to be downregulated in larvae lacking 20E signaling. In addition, we are examining the role of the *Drosophila* Estrogen Related Receptor (dERR) in this late-larval metabolic transition. Since dERR is required to activate glycolytic gene expression at the onset of larval development, and previous studies demonstrate that dERR activity is down-regulated during the late-L3 stage, we are testing the hypothesis that ectopic dERR activation would prevent down-regulation of glycolytic gene expression. Here too, our preliminary data indicate that ERR signaling may need to be attenuated to allow for the downregulation of glycolysis prior to metamorphosis. Overall, our findings suggest that nuclear receptor signaling plays a key role in the transcriptional down-regulation of glycolytic genes at the cessation of larval growth.

755A  Mutational characterization of phosphorylation sites suggests sex-specific regulation of the metabolic regulator Lipin  Michael Lehmann, Madeline Richards, Stephanie Hood University of Arkansas

Lipins constitute a conserved family of eukaryotic metabolic regulators. They are dual-functions proteins, acting as phosphatidate phosphatases in lipid synthesis in the cytoplasm and as transcriptional regulators of metabolism in the nucleus. The single *Drosophila* Lipin ortholog contains 26 serine and threonine phosphorylation sites that were identified by mass spectrometry, suggesting complex posttranslational control of Lipin functions. Indeed, in both flies and mammals, nuclear translocation of Lipin has been shown to be controlled by TORC1-dependent phosphorylation. To identify additional functions of phosphorylation, we have carried out a systematic mutational analysis of single phosphorylation sites and groups of sites, replacing serine and threonine residues with alanine residues or the phosphomimetic glutamic acid. CRISPR/Cas9-generated mutants were then characterized for Lipin expression, metabolic defects, and starvation resistance. Notably, we found that one of these mutants, carrying an amino acid substitution at serine residue 820 (S820), exhibited a significant difference in starvation resistance in the male and female sex. Interestingly, S820 is strategically located between the catalytic and transcriptional co-regulator motifs of the protein. Both males and females expressing Lipin S820E, in which the serine residue is replaced by phosphomimetic glutamate, showed reduced starvation resistance. In contrast, female flies expressing non-phosphorylatable Lipin S820A displayed significantly increased starvation resistance, whereas males showed reduced starvation resistance. These data suggest that the phosphorylation status at specific sites of the Lipin protein differentially affects functions of the protein in males and females. Thus, it may contribute to sex-specific differences in metabolism. The metabolic consequences of S820 phosphorylation in males and females are currently under investigation.

756B  Developmental Effects of Cactus on *Drosophila mettleri*  Lidane Noronha, Brian Lazzaro, Patrick O’Grady Cornell University

The cactophilic *Drosophila* system of the Sonoran Desert is an excellent model not only to study how organisms may thrive in toxic environments, but also how some taxa are associated with the microbes necessary to decompose plant tissue. Cacti contain toxic compounds that protect them from herbivory. When a cactus is injured, sap exudes from its wounds and microorganisms quickly colonize the sap and begin the decomposition process. Interestingly, several *Drosophila* species can exploit these substrates as feeding and breeding sites. While many members of the cactophilic *Drosophila repleta* species group have been able to transition seamlessly onto a standard cornmeal diet, *Drosophila mettleri*, is an exception. This is the only known species to breed exclusively in soil soaked with necrotic cactus sap and tissue, allowing it to take advantage of a more toxic and microbially rich niche. This taxon requires cactus-supplemented media to maintain healthy cultures under laboratory conditions. We hypothesized
that D. mettleri’s reliance on its host plant and associated microbiome resulted in the loss of its ability to adapt to a novel, more sterile environment.

Our understanding of the physiological processes required for Drosophila to utilize decomposing cactus as a host plant is limited. For example, we currently do not know whether D. mettleri derives nutritional benefit from the decomposing cactus, the saprophytic microorganisms responsible for the decomposition process, or a combination of the two. This study examines D. mettleri to understand how various life stages perform when reared on (1) standard media, (2) standard media + cactus powder, and (3) standard media + sterilized cactus powder. Development rates between each life stage and adult feeding behavior were examined. The results show that cactus has an essential role in ensuring that Drosophila survive the pupation and eclose successfully.

757C  Drosophila Undigested Metabolite Profiling - Uncovering age-related changes in amino acid absorption Abigail Monnem, Rachael Dack, Rebecca Clark Department of Biosciences, Durham University, UK

Unintentional weight loss is a common characteristic of ageing and frailty. While this may be partially explained by dietary changes and the metabolic cost of immune activation and inflammation that occur with age, other factors must contribute.

Nutrient absorption is a key role of the intestine which has not been studied in the context of ageing. Given the myriad changes occurring with age in the intestine, from increased bacterial load, to dysplasia and loss of intestinal barrier integrity, we feel that nutrient absorption is a likely contributing factor to this unintentional weight loss.

We have developed an assay to assess changes in the nutrient absorption/excretion profile in response to variables such as age or microbiota composition. Through this, we have found that more nutrients are egested with age, indicating a reduced absorptive capacity which may contribute to the unintentional weight loss observed with ageing and frailty. We are investigating changes in transporter expression with age as a possible mechanism. Additionally, by comparing the profile from axenic and conventionally reared flies, our assay confirms that the microbiota is a large source of protein and suggests that the microbiota further reduces the absorptive capacity of Drosophila. Thus potentially implicating the microbiome in age-related weight loss.

758A  Embryonic lipid transport works with TORC1 to ensure rapid and efficient development Marcus Kilwein, Michael Welte University of Rochester

Maternal nutrient stores are abundant and transient, providing embryos with energy that fuels development and carbon scaffolds for anabolism. Among this nutrient supply, the fat storage organelles, lipid droplets (LDs), occupy a unique position. At cellularization, most LDs are allocated to the incipient epithelium, while carbohydrate and protein reserves are confined to the yolk cell. Previous work has shown that two distinct mechanisms ensure this privileged localization of LDs. First, the LD protein Jabba keeps LDs and glycogen granules apart; LDs lacking Jabba protein embed into the surface of glycogen granules which results in them being mislocalized to the yolk cell. Second, during the blastoderm stages, LDs move bidirectionally along microtubules, driven by kinesin-1 and cytoplasmic dynein. In the absence of the motor regulator Klar, dynein activity predominates, also resulting in LD mislocalization to the yolk cell. To investigate why this LD distribution exists, we followed Jabba and klar embryos through subsequent embryogenesis and assessed triglyceride consumption, LD localization, and timing of embryogenesis. The yolk cell localized LDs in Jabba and klar persisted through hatching, demonstrating that proper LD localization is crucial to their consumption. This disruption in LD consumption appears to have detrimental consequences for the embryo. On the one hand, knocking down the triglyceride lipase Brummer in Jabba and klar mutants reduces embryo hatching success; under the same conditions, wild type embryos hatch normally. To test whether the hatching delay results from reduced LD consumption or excess LDs damaging the yolk cell, we generated embryos with a reduced maternal supply of LDs using mutants in the LD protein dPLIN2 or in the lipid uptake receptors LpR1 and LpR2. Both genotypes showed significant developmental delays in reciprocal crosses. We therefore conclude embryos respond to reduced access to LD stores with a developmental delay. To elucidate the genetic players in the delay, a candidate RNAi screen was done in reciprocal crosses. We therefore conclude embryos respond to reduced access to LD stores with a developmental delay. To elucidate the genetic players in the delay, a candidate RNAi screen was done in reciprocal crosses. To this end, we will be using Drosophila melanogaster to better understand how dysregulation of hormonal

759B  Consequences to Organismal Physiology upon Dysregulation of Hormonal Homeostasis using Drosophila melanogaster Cameron Dixon, Julia Ye, Kim McCall Boston University

Hormones orchestrate vital physiological pathways, such as metabolism, development, and reproduction. Beyond their influence on organismal homeostasis, hormones maintain integrity of stem cell niches and tissue integrity through molecular mechanisms. The presence of hormones is not restrictive to mammals and is evolutionarily conserved across species down to microorganisms. This conservation through evolution shows the importance in understanding how hormones interact with physiology and the negative implications of what happens to homeostasis post-hormonal disruption. To this end, we will be using Drosophila melanogaster to better understand how dysregulation of hormonal
homeostasis impacts physiology. *Drosophila* possess hormonal regulation that shares many similarities with mammalian hormonal homeostasis. While mammals are more complex organisms, many of the vital hormones essential to physiology are conserved. For example, the mammalian peptide hormone, insulin, is present in *Drosophila*. In *Drosophila*, it is known as *Drosophila* Insulin-like Peptide (DILP) and there are eight forms. Similar to how mammals have various insulin types, these eight forms serve specific functions across the organism to regulate glucose needs and metabolism. Additionally, DILPs are heavily involved in other physiological processes, including other hormonal pathways that regulate development and reproduction. The major hormones that are intimately connected to DILP are ecdysone and juvenile hormone (JH); which function similarly to that of mammalian sex steroids and growth hormones, respectively. All three of these major hormones are not only involved in their own canonical pathways but also influence each other by being necessary for each other to complete their function. To fully understand how these hormones effect physiology once homeostasis is dysregulated, we will be examining the effects of hormonal perturbations on physiology. Perturbations will be completed using the Gal4-UAS system and will be visualized using microscopy methods. Preliminary results have shown consequences to physiology through malformation of wings and increased wing venation, decreased survival, temperature sensitivity, and apoptotic phenotypes in reproductive tissues. Further investigation into these consequences are currently being completed to understand the effects of perturbing major hormones (DILP, ecdysone, JH) and their effects on physiological homeostasis and hormonal crosstalk.

**760C  Hormonal Effects of Glyphosate Based Herbicides on *Drosophila melanogaster*** **Maggie Santos**, Becky Tayln  
California State University San Bernardino

Glyphosate based herbicides are the most commonly used world-wide. The effects of glyphosate on reproductive and endocrine systems has not been fully explored. We used a JH inhibitor, precocene, and a JH analog, methoprene, to determine if glyphosate interacts with the JH pathway. So far, the lowest survivability to day three has been for Roundup Super Concentrate treated with precocene. Glyphosate, the active ingredient in Roundup, is a non-selective herbicide. It is the most widely used pesticide due to its indiscriminate ability to kill weeds. Originally believed to only affect plants because it affects the shikimate pathway, which animals lack, studies indicate that glyphosate based herbicides (GBHs) harm animals and microorganisms. GBHs can cause endocrine disrupting behaviors in non-target organisms (Gasnier et al. 2009), including oocyte production (Talyn et al. 2020). We seek to identify a mechanism by which GBHs reduce oocyte production and decrease reproduction, using the *Drosophila melanogaster* model system. We will use a JH inhibitor, precocene, and a JH analogue, methoprene, along with treatments of GBHs and glyphosate alone to determine whether GBHs act on reproductive output through a pathway involving JH, an important hormone in Drosophila reproduction (Wilson 1982).

**761A  Distinct dietary nutrients regulate circulating levels of Dilp2 and Dilp6 in *Drosophila* larvae** **Miyuki Suzawa**, Dalton Hilovsky, Kyle McPherson, Michelle Bland University of Virginia, Charlottesville, VA

The insulin/IGF1 signaling pathway regulates cell and organismal growth in animals ranging from fruit flies to humans. Insulin-like hormones are secreted in response to dietary nutrients and drive anabolic metabolism. In *Drosophila*, seven insulin-like peptides (Dilps) activate the insulin/IGF signaling pathway through a single known insulin receptor. We previously reported that Toll signaling in *Drosophila* larval fat body inhibited whole-animal growth by reducing circulating levels of fat body-derived Dilp6. In contrast, Dilp2 secretion from insulin producing cells was not affected by Toll pathway activation in larval fat body. To determine whether other physiological conditions differently regulated these two related hormones, we asked how Dilp2 and Dilp6 respond to starvation and specific dietary nutrients. We used dual-epitope tagged *Dilp2* and *Dilp6* alleles to measure hemolymph levels of each hormone in male and female larvae. Starvation of mid-third instar larvae on agar led to a 50% reduction in circulating Dilp2 after 6 hours and a 90% reduction in Dilp2 after 24 hours. In contrast, circulating Dilp6 levels were unchanged over the course of 24 hours of starvation on agar. Next, we asked whether Dilp2 and Dilp6 respond to specific dietary nutrients. We fed larvae sucrose, peptone or yeast extract in agar and compared hormone levels with fed and starved conditions. Hemolymph Dilp2 was partially recovered by feeding larvae peptone or yeast extract but not sucrose. Circulating levels of Dilp6 were modestly increased by a sucrose-only diet, but surprisingly, peptone or yeast extract feeding repressed Dilp6. The reduced circulating level of Dilp6 caused by the high protein, yeast extract diet was partially rescued when sucrose was added back. These data suggest that Dilp2 and Dilp6 secretion are regulated differently by environmental conditions ranging from infection to dietary composition. Two open questions are how Dilp2 and Dilp6 secretion are regulated by specific nutrients and how the single known insulin receptor integrates divergent signals from distinct Dilps to control growth and metabolism.

**762B  Dynamic expression of Lgr1 in the hindgut suggests a role in cold tolerance and acclimation** **Daniel Munteanu**, Sara Helms Cahan University of Vermont

Thermal tolerance of an organism depends on both its ability to dynamically adjust to a thermal stress and preparatory processes that enhance resistance to extreme temperatures (either cold or hot). However, our understanding of the molecular mechanisms that underpin these acclimatization processes is limited—especially for the cold. It has been previously shown that cold acclimated flies are better able to maintain ion homeostasis and fluid reabsorption across
the hindgut epithelium in extreme cold. We had previously conducted a GWAS analysis of the critical thermal minimum (CTmin) in the Drosophila Genetic Reference Panel (DGRP), which identified two single nucleotide polymorphisms (SNPs) in the first intron of Lgr1, a G-protein coupled receptor (GPCR) expressed predominantly in the hindgut. Baseline Lgr1 expression was higher in flies raised at colder temperatures as adults, suggesting that changes in Lgr1-mediated signaling may be involved in conferring cold stress resistance. To test this hypothesis, expression of Lgr1 was knocked down with Gal4 driving RNAi expression in hindgut. Lgr1 knockout significantly impaired baseline CTmin in both sexes, and in males, which express ~6X higher Lgr1 under baseline conditions, knockout reduced cold acclimation capacity. Altogether, this suggests that Lgr1 may play an important role in driving hindgut reabsorption during cold stress. Future studies will focus on elucidating Lgr1 interactors and identifying its part in the preparatory physiological processes changing to maintain homeostasis in the extreme cold.

763C Mechanisms of Action and Natural Variation within Fasting-induced Starvation Resistance in Drosophila Benedict Lenhart, Alan Bergland University of Virginia

Nutrient availability is a component of environment that impacts metabolism and organismal fitness. How nutrient-limitation affects organismal survival remains unclear, in part because we do not understand the extent of genetic variation in metabolic response to this environment. Research has established that limiting nutritional availability via a periodic fasting treatment produces health benefits including longevity and starvation resistance across a broad range of taxa, though this work has commonly avoided assaying natural populations. To expand our understanding of how clines such as space and time impact an organism’s physiological response to a nutrient-limiting environment, we are establishing a study system from environmentally-distinct populations of Drosophila melanogaster. We will sample and create lineages from latitudinally and seasonally varying populations, and leverage the genetic differences between these populations to explore variance in the ability of periodic fasting to improve starvation resistance. In the forthcoming assays: flies will be removed from nutrients for varying extents during early adult life, and their subsequent starvation resistance, fat storage levels, and fat storage dynamics measured throughout adult life. Early results lead us to hypothesize that while three weeks of periodic fasting can improve starvation resistance, this effect is line specific and subject to variation. Moreover, while measurements of fat concentrations will decrease post fasting, fasted-flies will be able to utilize existing reserves with higher efficiency during starvation. This work aims to indicate how schedules of nutrient limitations lead to changes to starvation resistance across natural populations, and how mechanisms of fat storage and lipophagy mediate this variation.

764A Time-restricted feeding improves striated muscle in genetic-induced obese Drosophila Yiming Guo1, Christopher Livelo1, Shweta Varshney2, Farah Abou Daya2, Hiep Le2, Satchidananda Panda2, Girish Melkani1,2 1) Department of Pathology, Division of Molecular and Cellular Pathology, School of Medicine, The University of Alabama at Birmingham, AL 35294, USA; 2) Department of Biology and Molecular Biology Institute, San Diego State University San Diego, CA 92182, USA; 3) Regulatory Biology Laboratory, Salk Institute for Biological Studies, La Jolla, CA 92037, USA.

Pathological obesity due to genetic predisposition affects countless people globally and leads to several ailments such as cardiovascular disease, metabolic syndrome and impaired muscle function. Our lab has developed a novel genetic-induced obesity model using Drosophila melanogaster which contains a sphingosine kinase 2 (Sk2) gene mutation. Sk2 mutation leads to accumulation of ceramide and eventually results in lipotoxicity which hampers muscle function and exhibits obese related metabolic parameters. Interestingly, by employing time-restricted feeding (TRF), an intervention which maintains isocaloric consumption that limits the daily feeding window from ZT0-ZT12, Drosophila melanogaster (fruit fly model) with genetic induced-obesity showed attenuated phenotypes. Flies under TRF improved skeletal muscle function, increased insulin sensitivity, reduced intramuscular lipid infiltration and preserved mitochondrial integrity compared to their ad libitum feeding (ALF) counterparts (Villanueva et al, 2019, Nature Communication). In this study, we explored the pathophysiological basis of TRF-mediated benefits utilizing transcriptomic data of indirect flight muscle (IFM). We investigated physiology using functional and genetic validations, cytological and biochemical approaches. We found flies under TRF intervention displayed significant upregulation of Gnm, Sardh and CG5955, key players of S-adenosylmethionine regulation, and downregulation of Dgat2, a key gene for triglyceride synthesis. IFM-specific knock-down (KD) of Gnm, Sardh and CG5955 led to progressive muscle dysfunction. Further, IFM-specific KD of Dgat2 retained muscle function during aging. In addition, genes associated with AMP kinase (AMPK) signaling, glycogen metabolism, glycolysis, tricarboxylic acid (TCA) cycle and electron transport chain (ETC) signaling were specifically upregulated in GIO model under TRF. Altogether, we identify the GIO-specific pathways in the regulation of muscle function under TRF, which emphasized the role of feeding-fasting rhythms on combating comorbidities linked with genetic obesity. Validation of these genes in obesity-induced skeletal muscle health and the role of circadian clock in that process is currently underway.

765B Time-restricted feeding promotes skeletal muscle function in diet-induced obesity through purine related pathway in Drosophila Christopher Livelo1, Yiming Guo1, Shweta Varshney2, Farah Abou Daya2, Hiep Le2, Satchidananda Panda2, Girish Melkani1 University of Alabama at Birmingham School of Medicine; Salk Institute for Biological studies
 Millions globally are affected by obesity stemming from a lifestyle of calorie-dense diets. Consequently, individuals afflicted with diet-induced obesity (DIO) incur complications including metabolic syndrome, cardiovascular disease, and compromised muscle function. In a previous study, we employed a time-restricted feeding (TRF) intervention, where daily feeding was limited to only the first 12 hours of the day in a Drosophila melanogaster model. Upon administration of a high-fat diet while under TRF, we observed improved skeletal muscle function compared to ad libitum feeding (ALF) counterparts in addition to improved metabolic parameters such as insulin sensitivity, mitochondrial integrity and body mass (Villanueva et al 2019, Nature communication). In this study, we evaluate the mechanistic basis of TRF-mediated benefits in DIO by utilizing muscle transcriptomic data of indirect flight muscle (IFM) followed by genetic validations, cytological and biochemical evidence. Interestingly, we found significant upregulation of glycine N-methyltransferase (Gnmt), sarcosine dehydrogenase (Sardh) and CG5955, key players of one carbon-metabolism and S-adenosylmethionine regulation under TRF. We found downregulation of diacylglycerol o-acyltransferase 2 (Dgat2), a key gene for triglyceride synthesis. RNAi knockdown of the upregulated genes led to muscle dysfunction and abrogated TRF-mediated improvement of muscle performance. However, knockdown of Dgat2 preserved muscle performance during aging. Furthermore, de novo purine biosynthesis appeared to be upregulated and potentially led to increased ATP levels resulting in improved muscle performance. Folic acid, a key factor required in fueling the purine cycle was found to attenuate high-fat diet related muscle impairment. These findings contribute to potential mechanistic foundations that underlie TRF mediated improvement in muscle and provide a blueprint for pursuing therapy related research development relevant to obesity and muscle impairment.

766C General anesthetics are toxic to flies mutant for a mitochondrially-encoded subunit of the electron transport chain. Amanda Scharenbrock, Zachariah Olufs, David Wassarman, Misha Perouansky University of Wisconsin-Madison, Madison, WI

Animals with mutations in Complex I of the mitochondrial electron transport chain (mETC) exhibit behavioral sensitivity to volatile general anesthetics (VGAs) and may be at increased risk of VGA-induced deleterious collateral effects. To investigate the toxicity of VGAs in mitochondrial mutants, we studied flies carrying a mutation in ND2, a gene in the mitochondrial genome that encodes a protein in the intermembrane arm of Complex I. We examined two commonly used VGAs isoflurane (ISO) and sevoflurane (SEVO), halogenated ethers, as well as various oxygen (O₂) concentrations. ISO proved to be toxic in hypoxic (5% O₂), normoxic, and hyperoxic (75% O₂) conditions while SEVO was only toxic in hyperoxic conditions. Interestingly, our previous work has shown that flies carrying a mutation in a nuclearly-encoded Complex I subunit, ND23, die when exposed to ISO in normoxia and hyperoxia, but ISO in hypoxia and SEVO under any oxygen conditions were not toxic. We have also expanded our anesthetics to include halothane (HALO), an alkane anesthetic, which like SEVO is non-pungent. Unlike SEVO, we found that HALO is toxic to both ND2 and ND23 mutants, with greater toxicity in ND2 mutants. These data indicate that mutations in different subunits of Complex I of the mETC confer differential susceptibility to VGA-induced toxicity. To investigate the underlying mechanism, we have begun examining whether heterozygous mutations in candidate genes enhance or suppress VGA-induced toxicity in ND2 mutants. To date, we have found that mutation of sima, which encodes a subunit of Hypoxia Inducible Factor (HIF) that transcriptionally regulates the response to hypoxia, enhanced the lethality of ND2 mutants exposed to ISO under normoxic or hypoxic conditions, indicating that HIF functions to prevent VGA toxicity. This work is part of an ongoing project aimed at identifying modifiers of VGA toxicity in mitochondrial mutants.

767A What Ingredients are Contributing to the Toxicity of Glyphosate-Based Herbicides, in Drosophila melanogaster? Noelle Roddam, Kalinah Winston, Becky Talyn California State University- San Bernardino

Growing evidence shows that some herbicides, including those intended to target plants, can be toxic to non-target animals and even humans, and that toxic residues remain in our food and water and pass to us through consumption. Our previous work, focused on glyphosate-based Roundup® formulations, indicates that Drosophila melanogaster fruit flies are a good model species to study these toxic effects on animals, by investigating their behavior, reproduction, anatomy, development, and mortality. This present study compares the toxicity of commercial glyphosate-based herbicide formulations to the toxicity of their key ingredients, alone and in specific mixtures. We hope to gain a better understanding of the toxic effects that glyphosate and other key ingredients lend to the toxicity of the formulations they are contained within, as well as that produced by the combination and interaction of the ingredients themselves. Preliminary results indicate that Glyphosate alone is not as toxic as the Roundup® formulations tested, which suggests that other ingredients, such as the known or unknown surfactants, are contributing to the overall toxicity of these formulations.

768B Positive selection of senescence through increased evolvability: ageing is not a by-product of evolution. Tristan Roget1,2, Pierre Jolivet3, Sylvie Méléard1, Michael Rera1 1) École Polytechnique, Palaiseau, France; 2) Université de Montpellier, Montpellier, France; 3) IRIT-APO - Algorithmes Parallèles et Optimisation - Institut de recherche en informatique de Toulouse, France; 4) Centre de Recherche Interdisciplinaire, INSERM, Université de Paris

The possibility of ageing being directly selected through evolution has been discussed for the past hundred years.
As ageing is occurring, by definition, only late in life, many think that it cannot be actively selected as a process. In addition, by decreasing an individual’s fitness, it is thought unlikely to be selected for. In order to explain the observation of its broad presence in the realm of life, numerous theories have been proposed in the past 60 years. Here, building upon a simple two parameters life history trait model that we recently introduced and that summarizes the life of an organism to its two core abilities - reproduce and thrive -, we discuss the possibility of ageing being selected for through evolution.

Our model shows that senescence will appear and be positively selected for through evolution thanks to the higher evolvability it confers to organisms. In addition, it predicts that the Lansing effect should be conserved in a large proportion of organisms showing age-related senescence. This formal and numerical analysis of ageing’s evolution also provides new hints to test the validity of existing theories by proposing a simple mathematical interpretation of the long-discussed longevity-fertility tradeoff or mutations accumulation.

769C Smurfness helps deconvolving ageing transcriptional signature Flaminia Zane1,5, Hayet Bouzid1, Céline Cansell1, Jean-Michel Camadro2, Sylvère Durand1, Christophe Antoniewski4,5, Michael Rera1 1) Center for Research and Interdisciplinarity (CRI), Inserm U1284, Université de Paris, 75006 Paris, France.; 2) Plateforme Protéomique/Spectrométrie de masse, Institut Jacques-Monod, 75013, Paris, France.; 3) Metabolomics platform, Institute Gustave Roussy, 94 805 Villejuif, France.; 4) ARTbio bioinformatic analysis platform, Institute of Biology Paris Seine, Sorbonne Université, F-75005, Paris, France.; 5) Sorbonne Université, 75006, Paris, France

Ageing is a process affecting a broad range of living organisms. In humans and multiple model organisms, it is characterized by an age-dependent decrease in functional efficiency and increased vulnerability to death. The classical approach for studying ageing involves comparing individuals of different chronological ages. Even if deregulation in certain pathways and mechanisms have been identified as associated with ageing, the so-called hallmarks of ageing, a comprehensive picture of the process is still missing. We believe that the inability to look at the physiological age of individuals - at the moment of sampling - rather than their chronological age is an important limiting factor in ageing research.

In 2012, Rera and collaborators described a new age-dependent and death-related phenotype in Drosophila melanogaster. By looking at a physiological increase in the intestinal permeability to a blue food dye, they were able to identify individuals committed to death in a few days. Interestingly, these individuals, called “Smurfs”, are the only ones, among a population, to exhibit various age-related changes and high-risk of impending death whatever their chronological age; Smurfness has been therefore hypothesized to be a good proxy to follow physiological age of single individuals.

We are here presenting our work on the characterization of the Smurf transcriptome using D. melanogaster. Through whole-body RNA-sequencing on Smurf and age-matched non-Smurfs flies of different chronological ages, we demonstrate that Smurfs carry a stereotypical transcriptional signature independently of their age. Strikingly, this signature mostly overlaps the previously described transcriptional signature of ageing. By studying concomitantly time-related changes and smurf-related changes in gene expression, we were able to identify genes moving through time but not necessarily associated to the physiological collapse of the organism and death. Those results confirm that not only Smurfs are a valid model to look at the physiological age of D. melanogaster, but are a powerful tool for deconvolving the changes related to physiological and chronological time. In conclusion, they highlight the importance of considering smurfness when designing ageing studies.

770A The role of commensal microbes in the longevity effects of Aronia berry (Aronia melanocarpa) in Drosophila melanogaster Ji-Hyeon Lee, Hye-Yeon Lee, Subeen Shin, Kyung-Jin Min Department of Biological Sciences and Bioengineering, Inha University, Incheon, Republic of Korea

Aronia berry (Aronia melanocarpa) is rich in polyphenolic compounds with various physiological and pharmacological activities. Several biologically active substances such as flavonoids and phenolic acids, have been reported to increase the lifespan of model organisms. With the increasing interest in the study of gut microbiota, it has been found that gut microbiota plays an important role in host health and aging. Recent studies have shown that the gut microbiota composition is changed by environmental factors like diet and aronia berry can modulate the gut microbial flora and lead to maintain a healthy/balanced gut microbiota. Although many studies have shown that aronia berry has anti-aging and prebiotic effect, the relationship between commensal microbes and the anti-aging effect of aronia berry is not illustrated yet. To test whether the anti-aging effect of aronia berry is related with commensal microbes, we generated axenic flies and measured their lifespan with or without aronia berry supplementation. In addition, 454-pyrosequencing analysis of the 16S rRNA gene was performed to investigate the microbial flora alteration in flies. Our results will provide the fundamental knowledge to address the relationship between commensal microbes and anti-aging effect of aronia berry.

771B The fly Tumor Necrosis Factor Receptor (TNFR), Wengen, restricts cytoplasmic TRAF3 levels to control gut metabolism, immunity, and tissue homeostasis Rihab Loudhaief, Christian Christensen, Julien Colombani, Ditte

#Dros22 | 257
The family of tumor necrosis factors (TNFs) are implicated in diverse processes ranging from cell proliferation, differentiation and apoptosis to innate and adaptive immunity. While TNFs contribute to tissue homeostasis and immunity, they are also notorious for their pathological roles in promoting tumor growth and inflammatory diseases. In addition to their well-described pro-inflammatory functions, an increasing body of evidence suggest that TNF signaling plays an instrumental role in orchestrating the metabolic rewiring associated with disease conditions in both flies and mammals. While this suggests that TNF can act as a metabolic hormone, it is not known whether TNF-TNFR signaling regulates to metabolic processes in healthy individuals. Here we show that the fly TNF receptor (TNFR), Wengen (Wgn), localizes in intracellular vesicles in the adult fly gut and plays an important role in controlling lipid metabolism, tissue homeostasis and immunity. Knocking down Wgn in enterocytes (ECs) results in activation of autophagy, degradation of lipids, and increased sensitivity to starvation. Moreover, wgn mutant guts display ectopic activation of JNK/MAPK and IMD signaling causing accelerated epithelial turnover and increased immune activity. We further show that all these effects are ligand-independent and mediated by TNF associated factor (TRAF)3, as Wgn is required to restrict cytoplasmic levels of TRAF3. These findings suggest that in addition to previously reported pathological roles of TNF-TNFR signaling in promoting inflammation and disrupting energy homeostasis, TNFRs are crucial mediators of metabolism and tissue homeostasis and moderators of immunity in healthy organisms.

**772C Screening for the genetic polymorphism underlying aging-related muscle degeneration** *Christina Talley, Soobin An, Kaveh Kiani, Anton Bryantsev Kennesaw State University*

Aging-related muscle tissue decline negatively impacts the quality of life for elderly individuals and causes billions of dollars in healthcare costs. The genetics of this phenomenon is not well understood, although such information may be critical for preventive care. Notably, previous attempts to link genetic polymorphisms with aging-related muscle decline have been limited to muscle structural genes.

We used *Drosophila* TDT (jump) muscle to study aging-dependent loss of muscle fibers. The TDT consists of 20-30 muscle fibers and functions to power jumping in flies. With progressive age, individual TDT fibers undergo spontaneous degeneration that is evident by the loss of cytoplasmic structures and nuclei. The extent and frequency of TDT degeneration are quantifiable and appear to be line dependent.

We have conducted a small genome-wide association analysis using 30 inbred fly lines demonstrating various rates of aging-dependent TDT fiber loss. Our data suggest that genes associated with the functioning of the nervous system are the strongest modifiers of TDT degeneration rates. To validate this finding, we further analyzed TDT degeneration rates in flies with TDTs uncoupled from the nervous system as well as seizure-prone flies with chronically hyperactivated nervous systems.

Our data demonstrates that deviations in the normal functioning of the nervous system significantly influence spontaneous muscle fiber degeneration during aging. Our results can guide future identification of the genetic polymorphism underlying muscle tissue decline in humans.

**773A Identifying the regulatory basis of sex differences in reproductive senescence in *Drosophila melanogaster*. Ruksana Amin, Caroline Hawkins, Rita Graze Auburn University*

Senescence is the gradual decline of function with advancing age. For example, in humans aging increases risk of hearing and vision loss. Reproductive senescence, is specifically the deterioration in reproductive success with advancing age. Hormone signaling pathways such as the insulin and insulin-like growth factor-like signaling (IIS) have many roles in aging and other processes. The IIS pathway, for example, can modulate not only lifespan, but also reproductive output, growth, metabolism, and stress. It has also been suggested that this pathway is likely to play a role in reproductive senescence. However, most reproductive senescence studies have focused on only one sex, though both show reproductive senescence and there are important effects of male-female interactions. Similarly, while the role of the IIS network in aging is well-characterized, there are few studies that have investigated tissue and age specific differences in both sexes allowing a direct comparison of changes in IIS with age. In order to understand why reproductive aging commonality differs between the sexes, we investigated how aging impacts sexually dimorphic expression of reproductive senescence candidate genes in different tissues — including those in the IIS pathway and in the sex determination hierarchy. Somatic and gonadal tissues were collected from males and females in a time series spanning day 3 to day 42 (post eclosion). RNA seq analysis will reveal how males and females differ in expression changes with advancing age.

**774B dSmad2 MARCM clones reveal a requirement for dILP2 secretion in the adult brain** *Samuel Goldsmith, Stuart Newfeld School of Life Sciences, Arizona State University, Tempe AZ*

Previous studies in our lab of dILP2 in the adult female brain showed that dCORL (fuss in Flybase) was required for
dILP2 expression in a subset of insulin producing cells (IPCs). To test the hypothesis that dCORL is acting downstream of dSmad2[SG(1)] in IPCs, as was shown earlier in the mushroom body of third instar larvae, we generated dSmad2 mutant MARCM clones in IPCs. Additionally, a series of follow-up experiments in IPCs were completed and the results were consistent with the data from the dSmad2 mutant MARCM clones. Employing a dCORL.Gal4 driver expressed in IPCs, dSmad2 mutant MARCM clones displayed excess dILP2 expression in comparison to adjacent wild type siblings (n=5; p=0.057). The overexpression phenotype was also seen with dSmad2 RNAi in IPCs driven by dCORL.Gal4 (n=14; p=0.013). The overexpression phenotype was rescued when expressing UAS.dSmad2 in mutant MARCM clones (n=7; p=0.67). The dSmad2 mutant phenotype appears similar to dILP2 overexpression in mutants for unpaired-2 that prevent dILP2 secretion. Studies with TrpA1 to test the secretion hypothesis are in progress. Overall, our study shows that dCORL functions independently of dSmad2 in the regulation of dILP2 expression in adult IPCs.

775C Identification of transcription factors acting in larval fat body to regulate whole-animal growth Dalton Hilovsky, Kyle McPherson, Leila Jamali, Shivani Reddy, Michelle Bland University of Virginia, Charlottesville, VA

The Drosophila larval fat body regulates growth of the whole animal in response to dietary nutrients, hormonal cues, and physiological stressors such as infection. The allocation of resources to growth and nutrient storage and the secretion of endocrine growth factors such as Dilp6 underlie fat body-dependent growth regulation. Indeed, our previous work showed that innate immune signaling in larval fat body targets Dilp6 to inhibit peripheral growth. To identify novel regulators of growth, we performed a directed genetic screen targeting transcription factors expressed in larval fat body. We screened for transcription factors that regulate peripheral growth basally and in response to active innate immune signaling in fat body. In a related screen, we queried a set of transcription factors shown to physically interact with the Dilp6 promoter using data from the ENCODE consortium. These screens identify novel regulators of whole-animal growth such as the transcription factor pleihomeotic like (phol), which modifies growth inhibition resulting from Toll signaling in larval fat body. Additionally, by screening predicted transcriptional regulators of Dilp6, we identified that increased expression of hairy (h) in fat body from the mid-third instar through the white prepupal stage reduces Dilp6 transcript levels. In contrast, fat body-specific knockdown of daughterless (da) suppresses Dilp6 expression in mid-third instar larvae but not in white prepupae. These two screens have identified discrete regulators that participate in Toll-dependent growth inhibition and Dilp6 regulation. Indeed, previous data from our lab indicate that reduced Dilp6 secretion in animals with active fat body Toll signaling does not account for the full degree of growth inhibition in these animals. Furthermore, our data point to additional, unappreciated signaling pathways that contribute to developmentally-appropriate regulation of Dilp6.

776A Lgr1 Localization Reveals a Larval-to-Adult Developmental Switch in Hindgut Compartmentalization Luis Sullivan1, Robert Scott1, Rosario Vicidomini2, Mihaela Serpe2, Benjamin White1 1) Laboratory of Molecular Biology, National Institute of Mental Health, NIH, Bethesda, MD 20892, USA; 2) Section on Cellular Communication, National Institute of Child Health and Human Development, NIH, Bethesda, MD 20892, USA

An evolutionarily ancient hormonal signaling system consisting of two glycoprotein hormone-like subunits, gpa2 and gpb5, and their leucine-rich repeat containing G-protein coupled receptor, called Lgr1 in Drosophila, are found throughout bilaterian genomes from flies to humans. The function of this conserved hormonal system remains obscure, but has been best characterized in the mosquito and the fruit fly, where Lgr1 is strongly expressed in the hindgut and has been implicated in maintenance of hydromineral balance. Using the Trojan exon method to create an Lgr1-specific Gal4 driver, we have confirmed robust expression of Lgr1 in both larval and adult hindgut, as well as in numerous other structures that support chloride transport such as the stellate cells of the Malpighian tubules and larval anal papillae. Interestingly, expression in the larval hindgut is limited to a ventral compartment, whereas in adult hindgut it is expressed throughout the hindgut epithelium. Using scRNAseq analysis combined with gene-specific Trojan Gal4 driver lines, we find that the larval epithelium is compartmentalized into ventral and dorsal cell-types conforming to those that express Delta and engrailed in the embryonic hindgut. Ventral cells express several genes implicated in ion transport and antidiuresis, while dorsal cells selectively express receptors for two known diuretic hormones, Dh44 and Leucokinin (Lk). These gene expression patterns suggest that half of the larval hindgut may provide water uptake from the hemolymph while the other may promote ion retrieval from the hindgut into the hemolymph. We speculate that this division of labor allows the larva to survive in its quasi-aqueous environment, permitting it to shed absorbed water, but retain ions. Consistent with this, hindgut compartmentalization is lost in the adult hindgut, together with Lk receptor expression, suggesting a purely antidiuretic function in the adult, which has a terrestrial lifestyle in which water retention is paramount.

777B fruitless Controls the Timing of Steroid Hormone Pulses in Drosophila Somatic Cell Jie Sun, Calder Ellsworth, Wu-min Deng Tulane University School of Medicine

Approximately 50% of all cancer patients develop cachexia, a devastating wasting syndrome. This figure increases up to 80% in patients with advanced cancers. In recent years, a number of Drosophila cachexia models have been established, however, steroid hormone level changes, a frequent symptom of cachexia in patient, has remained insufficiently
understood. We demonstrate here that the BTB zinc-finger transcription factor fruitless (fru) is a key player of steroid hormone signals and negatively regulates the synthesis of ecdysone in prothoracic gland (PG) cells by responding to steroid hormone levels in the body. Specifically, we show that Fru oscillates within the PG cells: a high concentration of Fru in the nuclei is concomitant with low-titer ecdysone levels in the body, and the protein is absent from PG nuclei at developmental stages when high-titer ecdysone pulses occur. Depletion of Fru in the PG arrests larval development by causing precocious ecdysone signaling and a failure to repress ecdysone pulses. In contrast, excessive Fru expression in the PG arrests development that can be rescued by administering an active ecysteroid, and we show that Fru inhibits the ecdysteroid biosynthesis pathway by regulating Halloween genes. Notably, we show that this regulation has implications for the cachexia model. Fru in PG nuclei of cachexic larvae is maintained at a high level, and the periodicity of Fru expression associated with normal development is abolished, thereby maintaining a low level of circulating steroids in cachexic larvae. These findings suggest that aberrant expression of Fru in PG is a major reason for hormone imbalance in cachexic larvae and is partially responsible for the cachexia phenotype, thus provide a theoretical basis for hormone treatment of cachexia.

### 778C  Studying the effect of Methotrexate on DNA damage and repair during ageing: drug treatments and models of JAK/STAT pathway-related blood cancers

Adel Alqarni, Martin Zeidler University of Sheffield

Originally developed as a chemotherapy drug, MTX acts as an anti-folate by binding to and inhibiting di-hydrofolate reductase (DHFR) that is involved in the synthesis of nucleotides required for DNA damage repair and cellular proliferation. In the 1980’s, MTX was repurposed as a treatment for auto-inflammatory and auto-immune diseases such as rheumatoid arthritis at much lower doses. However, the mechanism of action of MTX as an anti-inflammatory agent has been unclear until 2015 when the Zeidler lab identified MTX as a JAK/STAT pathway inhibitor. While our insight suggests that MTX could be repurposed again to treat JAK/STAT associated diseases, concerns about potential mutagenic toxicities associated with its inhibition of DHFR have been raised.

My project uses Drosophila melanogaster to study the effects of low-dose MTX therapy, its potential mutagenic activity and its effects on JAK/STAT signaling in vivo. More broadly, I aim to examine the impact of MTX and to better understand the processes that modulate DNA damage and DNA-damage responses. We have established an in vivo assay for DNA damage using w++; GMR-Gal4, UAS-white RNAi/+ flies to report the frequency of DNA damage in cells of the developing fly eye. Using this assay, along with measures of lifespan, fecundity, chromatinization, and JAK/STAT pathway activity we have tested a range of MTX doses added into Drosophila food. The effects on survival, development and DNA damage phenotypes will be shown. My results indicate that that high drug concentrations result in low survival rates, fecundity and developmental defects as well as a significant increase in the rate of DNA damage. However, low doses of MTX do not affect survival or fecundity and surprisingly, show significant increases in lifespan and may even reduce the rate of DNA damage. Suggesting that reduced JAK/STAT pathway activity may be providing a benefit under these conditions. In conclusion, while high doses of MTX lead to low survival and are mutagenic, low-doses may confer a survival and DNA-damage advantage. It will be interesting to establish whether these low-dose benefits are indeed linked to changes in JAK/STAT pathway activity in vivo. Latest results will be presented.

### 779A  Determining Critical Period of Herbicide Sensitivity in the Fruit Fly, Drosophila melanogaster

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Herbicides are widely used in agriculture, but many harm animals in addition to weeds. The most widely used brand of herbicide, Roundup®, occurs in multiple formulations, with proprietary ingredients, but currently all Roundup® formulations sold in the U.S. contain the active ingredient glyphosate. In this study, we determine the critical period for Drosophila melanogaster in terms of mortality and harm to the reproductive system after exposure to a Roundup® formulation with only glyphosate as an active ingredient, a Roundup® formulation that contains glyphosate and pelargonic acid, or Scythe®, an herbicide that contains only pelargonic acid as an active ingredient. For all herbicide treatments and exposure at all life stages, fewer larvae survived to adulthood than in control treatments at concentrations above 0.5 g/L. Preliminary data suggests that first instar larvae are most susceptible to mortality, regardless of herbicide exposure. Reproductive activity decreased with higher concentrations of herbicide exposure, regardless of formulation or exposure stage, possibly because survival was so low for those exposed starting at their 1st instar. Overall, 1st instar larvae are more susceptible to the harmful effects of exposure to both glyphosate-based and pelargonic acid-based herbicides than those exposed at later stages, and larvae of all stages are more sensitive than adults tested in previous studies with the same formulations and range of concentrations.

### 780B  Retrotransposons: a major driving force of aging

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Across all metazoans, aging correlates with functional decline. One potential contributor this is the expression and mobilization of retrotransposons (RTs), which are mobile genetic elements capable of self-replication and insertion into new genomic locations. RT expression and copy number have been shown to increase with age within multiple model organisms. Our goal is to determine de novo RT insertion number and genomic location within somatic cells of *Drosophila*. By carrying out whole genome sequencing from individual indirect flight muscle (IFM) nuclei from young and old female flies, RT insertions increase with age, with one of the most age-associated new insertions being from the element 412. However, unlike many RTs, the expression of 412 mRNA does not increase with age but remains constant throughout life. Additionally, this increase in RT insertion with age may not occur in all tissues, as when we purified nuclei from whole female thoraces that reflect a range of distinct cell types, we do not observe an age-associated increase in RT insertions. To determine the biological impact of 412 expression and/or insertion, we ubiquitously knocked down the expression of 412 using an inducible shRNA transgene and CRISPRi. This significantly extended female median and maximum lifespan. However, performance in several assays of health was not improved when 412 was knocked down, indicating that longevity is being targeted irrespective of changes to health. Since, we observe increase in 412 insertions in IFM, we also knocked down 412 specifically in this tissue and saw an increase in female lifespan, consistent with the importance of TE insertions in this cell type. In contrast, lifespan was unaltered by knocking down 412 in fat body. To understand the basis for the increased lifespan caused by reducing 412 expression we carried out RNA-sequencing using thoraces of control and 412 knockdown animals. These data suggest that knockdown flies may have improved mitochondrial function as several components of the electron transport chain are significantly upregulated.

**781V**  Parkinson’s disease genes interact with ATP7 to regulate copper distribution and availability in *Drosophila melanogaster*  Brooke Allen, Alysia Vrailas-Mortimer  Illinois State University

Copper is an essential element for enzymes that catalyze oxygen-dependent reactions. When an organism is exposed to either excess copper or deprived of copper, this micronutrient becomes detrimental. A mechanism used to control copper distribution and availability involves the ATPase transporter, ATP7. This X-linked transmembrane protein is responsible for delivering copper into the lumen of the cell by utilizing both endocytic and exocytic mechanisms. Mutations in ATP7 have been shown to cause Menkes disease and Wilson’s disease, which both share the phenotype of neurodegeneration. These genetic disorders with ATP7 defects both lead to mechanisms of neurodegeneration that is likely shared with other, more common neurodegenerative diseases, such as Parkinson’s disease. A screening of possible candidate genes that interact with ATP7 was conducted by inhibiting Parkinson’s disease genes in a ATP7 loss of function or ATP7 overexpression background. We find that several of the Parkinson’s disease genes showed a genetic interaction with ATP7, indicating that the mechanisms of neurodegeneration caused by ATP7 mutations may be conserved in Parkinson’s disease. These interactions and their link to neurological disorders will further discussed.

**782V**  Age-related neuroprotection by dietary restriction requires OXR1-mediated retromer function  Kenneth Wilson, Sudipta Bar, Enrique Carrera, George Brownridge III, Jennifer Beck, Tyler Hillsbeck, Christopher Nelson, Geetanjali Chawla, Rachel Brem, Hugo Bellén, Lisa Ellerby, Pankaj Kapahi  1) Buck Institute for Research on Aging, Novato, CA; 2) Dominican University of California, San Rafael, CA; 3) Leonard Davis School of Gerontology, University of Southern California, Los Angeles, CA; 4) Regional Centre for Biotechnology, Faridabad, Haryana, India; 5) Department of Plant and Microbial Biology, University of California, Berkeley, Berkeley, CA; 6) Department of Molecular and Human Genetics, Baylor College of Medicine, Houston, TX

Dietary restriction (DR) is the most robust method to delay aging and the onset of neurodegeneration, though the mechanisms behind this phenomenon remain unclear. Further, it remains unknown which factors influence why different individuals will respond to dietary interventions to different degrees. We reared over 150 fully sequenced fly strains from the Drosophila Genetic Reference Panel under ad libitum feeding or diet-restricted conditions and measured lifespan as well as healthspan. Through genome-wide association study, we identified genetic variants associated with influencing these traits under each dietary condition. A variant in one gene, *mustard (mtd)*, called *Oxidation resistance 1, OXR1*, in humans, significantly associated with DR-specific lifespan. We demonstrate that mtd/OXR1 in neurons is necessary for DR-mediated lifespan extension and that neuronal overexpression of human OXR1 is sufficient to extend lifespan upon DR in flies. Neuronal knockdown of mtd also inhibits dietary restriction-associated slowing of age-related visual decline, arguing for a specific role of mtd/OXR1 in DR-mediated neuroprotection. We additionally identified that natural variants of mtd in the promoter region are associated with enhanced longevity upon DR and are regulated by the transcription factor Traffic jam (TJ). We further show that mtd is essential for stabilizing the retromer complex, which is necessary for trafficking transmembrane proteins for reuse. As a result of OXR1 deficiency, the retromer destabilizes and lysosomes become overused. Overexpression of retromer proteins or supplementation with chaperone compound R55 rescues the lifespan defects and neurodegeneration seen in mtd-deficient flies, and R55 is capable of rescuing lysosomal aggregation and OXR1-retromer co-localization in cells from humans with OXR1 deficiency. Thus, mtd/OXR1 enhances protein
Variations in rate of aging in genetically heterogenous populations support the hypothesis that aging is at least partially genetically regulated. However, genetically identical individuals also vary in their time of death even when maintained in the same environment. We have observed in *D. melanogaster* that this variation is genotype dependent, as specific genotypes have characteristic survival curve shapes that are largely reproducible. We hypothesize that there is an underlying genetic component to this seemingly stochastic nature of lifespan curves. Typical aging studies reduce a strain's lifespan down to a population-level value, such as mean or max lifespan. While these metrics can represent the trends in a population, they are unable to encapsulate the variation in the aging of individuals from the same distinct population. Instead, we used two values that characterize the logistic fit of a strain's hazard ratio over its lifespan: the risk of initial mortality (α) and the rate of aging (β). To identify regulators of the rate of aging and stochasticity of lifespan, we performed a Genome-Wide Association Study (GWAS) of β for 160 different fly strains from the DGRP collection on two different diets late in life. This approach identified the candidate gene *Diabetes and Obesity-Regulated (DOR)*, which has known roles in stress response, autophagy, and senescence, as having a role in the late-life mortality. *DOR* inhibition leads to a significant increase in late-life mortality that is preceded by a reduction in healthspan-related traits. Further, we’ve found that germline-specific inhibition is sufficient to shorten lifespan. We conclude that a decrease in expression of *DOR*, a conserved gene, compromises an organism’s resilience, leading to a decline in healthspan and increased mortality, providing a potential target for bolstering the decline seen in human aging.

**784V**  Inter-kingdom lipid transfer mediates *D. melanogaster* temperature-adaption  Claudia Espinoza, Itay Budin University of California San Diego

Animals must undergo physiological modifications to adapt to yearly climate fluctuations. At low temperatures, cell membranes undergo a phase transition from liquid to gel-like state, lowering membrane fluidity and disrupting the function of essential membrane-associated machinery. In organisms that overwinter, a class of lipids in their membranes, polyunsaturated fatty acids (PUFAs) with 2-6 double bonds, can act to maintain membrane fluidity and function at low temperatures. However, most animals, including *Drosophila*, lack the desaturase enzymes responsible for the synthesis of PUFAs, and must rely solely on dietary sources to obtain this lipid class. This lipid exchange underlying the interplay between foraging behavior, food composition and cell membrane organization, which is crucial for organisms’ seasonal adaptation, remains unexplored. Using *Drosophila melanogaster* and different yeast species as its dietary counterpart, we are investigating how PUFAs obtained through diet determine seasonal adaptation and what the mechanisms of PUFA detection and selection are. Analysis of *D. melanogaster* associated yeast collected at different times of the year allowed us to identify the increase in PUFA-producing species during the cold months. In the lab, we find that controlled dietary intake of yeast PUFAs dictates *Drosophila* activity at low temperatures. Lipidomics analysis shows how the presence of the key temperature-responsive PUFA linolenic acid is determined in specific lipid classes as a function of diet. We also find that temperature also determines both feeding and oviposition preferences, with flies favoring feeding from and laying eggs near PUFA-producing yeast at low temperatures. Using a combination of mass spectrometry, feeding assays, and yeast and *Drosophila* mutants, we are currently elucidating the yeast volatiles and *Drosophila* genetic pathways responsible for the detection and selection of dietary PUFAs at low temperature. This knowledge will help us understand the mechanisms by which lipid exchange influences seasonal adaptation in symbiotic organisms like yeast and *Drosophila*.

**785V**  The impacts of sex and genetic background on the response of *Drosophila melanogaster* to essential and non-essential metal toxicity  Mitchell Slobodian, Athena Wallis, Joshua Keeping, Allie Sutherland-Hutchings, Jesse Petahtegoose, Solomon Arthur, Danica Levesque, Thomas Merritt Laurentian University, Sudbury, Ontario

*Drosophila melanogaster* is an effective model of metal toxicity, but we currently have only a limited understanding of any sex or genetic background effects. Similarly, while essential metals are relatively well studied, many non-essential metals remain understudied despite their chemical similarities to essential metals and their relevance as environmental contaminants. Further, essential and non-essential metals are not independent. The two groups interact with multiple interconnected metal pathways in a complex web with many proteins and mechanisms yet to be elucidated. In the current study, we are quantifying the influence of sex and genetic background on the response to essential and non-essential metal toxicity in *D. melanogaster*. We are using copper (Cu) and nickel (Ni) as our essential and non-essential metals, both delivered through the fly food. The two are chemically similar metals and can bind to many of the same biological ligands. Our study also includes different homozygous fly lines from the *Drosophila* Genetic Reference Panel (DGRP) to examine background effects. To date, we have conducted multiple four-day exposure Lethal Concentration 50 (LC50) assays using a range of CuSO4 and NiSO4 concentrations. There are significant effects of sex and genetic background. The effects of sex and genetic background on the response to essential and non-essential metal toxicity are likely to be crucial for understanding the factors that influence the susceptibility to metal toxicity in *Drosophila*.
background for both metals. Background effects were particularly substantial, and interestingly, the same lines were sensitive or resistant to both metals. We also found similar sex effects and combined sex and line effects for the two metals. Currently, we are quantifying the amount the concentration of metals in the fly samples, not just the fly food using two different methods, one with the non-specific metal chelator 4-(2-pyridylazo)resorcinol (PAR) and the second with Inductively Couple Plasma Mass Spectrometry (ICP-MS). Future work will investigate the genetic and biochemical influences that cause the differences in response to metal toxicity and the effects of simultaneous exposure to both metals. Our results to date clearly demonstrate that sex and genetic background are major influences in the response to essential and non-essential metal toxicity. Understanding the biology of this system, and likely any other biological system, requires experiments with multiple genetic backgrounds and both sexes.

**786V** HIF-1α promotes hypoxia tolerance by restraining excess cytokine signaling Kate Ding¹,²,³, Danielle Polan¹,²,³, Byoungchun Lee¹,²,³, Tiffany Cheung¹, Elizabeth Barretto¹, Savraj Grewal¹,² ¹) University of Calgary, Calgary, AB; 2) Arnie Charbonneau Cancer Institute; 3) Department of Biochemistry and Molecular Biology

Our cells and organs need oxygen from the air we breathe in order to function normally. In conditions of low oxygen, also known as hypoxia, animals experience tissue damage and deregulation of metabolic homeostasis which are characteristic of diseases such as stroke and ischemia. Tumors are also associated with hypoxia, which is thought to trigger enhanced cell proliferation and metastasis. Furthermore, diseases that obstruct proper oxygen intake such as chronic lung disease and sleep apnea can result in intermittent, but chronic, whole-body hypoxia that can have long-term deleterious effects on health.

Although the air we breathe is made up of ~20% oxygen, our cells and tissues experience much lower levels than this, ranging from 1-10%. This aspect of normal physiology is often overlooked in cell culture experiments where the cells are typically maintained in 20% oxygen. Understanding how tissues and organs respond to low oxygen is therefore an important question in biology. *Drosophila* experience sustained conditions of hypoxia as they bury in rotting fruit they forage through during the larval stages. They have therefore evolved to tolerate hypoxia. Trachea deliver oxygen directly to tissues, hence by modulating environmental oxygen we can trigger hypoxia in cells and tissue. As such, *Drosophila* have emerged as an ideal in vivo model for studying how organisms tolerate hypoxia. Recent studies have revealed an adaptive role for Upd/JAK/STAT signaling in response to conditions of low oxygen that involve tissue-tissue crosstalk to promote differentiation of specific immune cells.

In our lab, we have demonstrated a role for cytokine signaling mechanisms - involving the fly interleukin-6 homolog upd3- which is important for whole-body hypoxia tolerance. We found that expression of upd3 is strongly upregulated (>20-fold) in hypoxia and that upd3 mutants show reduced survival in low oxygen. Interestingly, we also found that knockdown of HIF-1alpha, the classic hypoxia-induced factor required for survival in low oxygen, lead to a further enhancement of upd3 expression. Further, when we genetically mimicked this enhancement in upd3 levels, we found animals also had reduced survival in low oxygen. Previous studies have demonstrated a link between HIF-1α and immune system modulation, thereby already implicating a role of hypoxia in immunity. However, with our recent discovery that upd3 is modulated in hypoxia and may be regulated in part by HIF-1α, this leaves open the question of what role upd3 is playing and how it promotes hypoxia tolerance. Using the versatility of *Drosophila* genetics, we have begun to tease apart what tissues may be producing upd3 and where the cytokines may be acting.

**787V** mTORC2 protects heart from HFD induced-damage through promoting mitochondrial fission Peiduo Liu ¹ Iowa State University

High fat diet (HFD)-associated lipotoxicity is one of the major causes of cardiovascular diseases. The mechanistic target of rapamycin (mTOR) pathway, especially mTOR complex 1 (mTORC1), has been previously implicated in HFD-induced heart dysfunction. In the present study, we find that unlike mTORC1, mTOR complex 2 (mTORC2) protects hearts from HFD-induced cardiomyopathy and mitochondrial dysfunction in Drosophila. We show that HFD feeding induces contractile dysfunction along with altered mitochondrial morphology and function. Upon HFD feeding, the mitochondria of cardiomyocytes exhibit fragmentation, decrease of membrane potential. Interestingly, short-term HFD feeding induces the activity of cardiac mTORC2 which could be a protective response. In line with this finding, the flies with cardiac-specific knockdown of rictor, the key subunit of mTORC2, show cardiac and mitochondrial dysfunction similar to what is observed in HFD-fed wild-type flies. Also the mitochondrial fission as an adaptive response is blocked by rictor knockdown. Conversely, cardiac-specific activation of mTORC2 by overexpressing rictor attenuates HFD-induced mitochondrial and cardiac dysfunction. Thus, our findings suggest that mTORC2 is a cardioprotective factor and may through promoting the mitochondrial fission.

**788V** Role of Wnt signaling in regulating lipid homeostasis in *Drosophila* Rajitha Udakara Sampath Hambo-Waduge¹, Xiahe Huang², Mengmeng Liu¹, Xiao Li¹,², Yingchun Wang², Jun-yuan Ji¹ ¹) Tulane University School of Medicine, New Orleans, LA; 2) Chinese Academy of Sciences, Beijing, China; 3) Princeton University, Princeton, NJ
Dysregulated Wnt signaling causes aberrant development and diseases such as cancer, but the role of Wnt signaling in regulating lipid metabolism remains poorly understood. We have reported that hyperactive Wnt signaling in *Drosophila* larvae reduces lipid accumulation in the fat body, which can be strongly rescued by feeding the larvae with proteasome inhibiting peptide boronic acids, such as Bortezomib. Interestingly, hyperactive Wnt signaling in adipocytes increases the levels of different types of free fatty acids but reduces the levels of a variety of triglycerides. To elucidate the mechanisms of how Wnt signaling regulates lipid homeostasis in adipocytes, we have performed transcriptomic and quantitative proteomic analyses using larvae with hyperactive Wnt signaling with or without Bortezomib treatment. These analyses led us to identify genes involved in regulation of lipid mobilization, such as *PLIN1* (*Perilipin*1), *PLIN2*, and genes encoding factors that regulate fatty acid beta-oxidation. Both *PLIN1* and *PLIN2* are significantly reduced by hyperactive Wnt signaling, but these effects are strongly reversed by Bortezomib treatment. Our genetic analyses suggest that adipocyte defects caused by hyperactive Wnt signaling can be strongly enhanced by depleting *PLIN1*, but rescued by ectopic expression of *PLIN1*, *PLIN2*, or both. Taken together, these observations suggest that Wnt signaling may regulate lipid homeostasis by promoting lipid mobilization through regulating the expression of *PLIN1* and *PLIN2* in larval fat body.

789V Identification of direct targets of Bortezomib in *Drosophila* using a chemical proteomics approach Mengmeng Liu Tulane University

**Identification of direct targets of Bortezomib in *Drosophila* using a chemical proteomics approach**

Mengmeng Liu¹, Yadagiri Kurra², Rajitha-Udakara-Sampath Hemba-Waduge¹, Xiao Li¹,³, Wenshe Liu², and Jun-yuan Ji*,¹

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Wnt signaling plays critical roles in regulating diverse biological processes, and deregulated Wnt signaling causes aberrant development and diseases such as cancer. We have previously reported that hyperactive Wnt signaling disrupts lipid homeostasis in *Drosophila* larvae, which can be strongly rescued by feeding the larvae with Bortezomib (BTZ) in an a-Catenin-dependent manner. BTZ is the first proteasome inhibitor approved by the Food and Drug Administration to treat patients suffering from multiple myeloma and mantle cell lymphoma. We have found that BTZ and two additional peptide boronic acids (PBAs), Delanzomib and Ixazomib, can rescue adipocyte defects caused by elevated Wnt signaling, but three other proteasome inhibitors that have different chemical structures (Carfilzomib, Marizomib, and Oprozomib) cannot rescue the phenotypes. These observations indicate that PBAs may have additional targets besides proteasome. To elucidate how PBAs regulates lipid homeostasis and Wnt signaling, we synthesized a photoactivatable biotin-tagged BTZ, which displays similar effects on rescuing adipocyte defects and inhibiting Wnt activity to BTZ. Using this biotin-tagged BTZ, we performed a pull-down assay using streptavidin beads followed by mass spectrometry analysis to identify proteins bind to BTZ. We have identified 126 candidate targets of BTZ, including majority subunits of the proteasome complex as expected. Interestingly, we also identified some peptidases and proteinases. We are performing genetic and cell biological analyses to determine whether these peptidases and proteinases play any roles in regulating a-Catenin stability and Wnt signaling in *Drosophila*. This study may advance our understanding of the molecular mechanism of how BTZ regulates a-Catenin stability and Wnt signaling in vivo.

790V Odor mediated control of blood-progenitor redox homeostasis in *Drosophila* Manisha Goyal², Ajay Tomar¹,², Sukanya Madhwal¹, Tina Mukherjee¹

1) Institute For Stem Cell Science and Regenerative Medicine (inStem); 2) The University of Trans-Disciplinary Health Sciences & Technology (TDU)

The importance of reactive oxygen species (ROS) in myeloid cell development and function is well established. However, any aberrant generation of ROS alters hematopoiesis. Thus, maintaining homeostatic levels of ROS is very crucial for progenitor development. While the autonomous mechanism of ROS homeostasis is explored but any physiological level interpretation that regulates this ROS homeostasis remains largely unclear. Here, we show a sensory regulation of redox homeostasis in blood-progenitor cells which is meditated by olfaction. We find that this long-range communication, to facilitate ROS homeostasis is central to the development of blood progenitor cells and successful immune response. This work shows yet another implication of environmental control of myeloid physiology at the axis of ROS homeostasis by odors and GABA. GABA metabolism in blood-progenitor cells regulate TCA cycle activity and antioxidants synthesis to control ROS balance in the lymph gland. We have identified the metabolic requirement of odor sensing and GABA in regulating redox homeostasis during *Drosophila* myeloid progenitor development, the relevance of which may be broadly conserved.

791V Effects of Ambient Temperature on Body Fat Jin Seo, Rachael Winfrey, Alexander Jochmans, Eunji Yoon Rogers State University

Temperature profoundly affects all living organisms on development, growth, longevity, and metabolism. Ambient temperature significantly alters lipid/carbohydrate storages in fruit flies; conversely, food intake alters flies’ preferred
temperature. The natural thermal range of fruit flies is between 13 and 33°C. When adult flies are incubated at a higher natural thermal temperature, body fat contents are strikingly reduced. Remarkably, fat loss does not appear to be due to decreased food intake and/or increased metabolic rate suggesting that temperature-mediated fat storage is regulated through active mechanisms rather than passive processes. The rates of biochemical reactions show a bell curve pattern in which the reaction rates increase up until the optimum temperature ($T_{\text{opt}}$) and drop sharply beyond the $T_{\text{opt}}$. To better understand mechanisms of temperature-mediated energy storage and identify thermosensing molecules, we have investigated enzyme activities in multiple signaling pathways, which are linked to lipid/carbohydrate metabolism, at suboptimum temperatures.

**792V** Lifestyles and metabolism of *Drosophila lutzii*, a floridosa group of species, and sympatric *D. simulans*, a generalist specie Juan Manuel Murillo-Maldonado, Juan R. Riesgo-Escovar Universidad Nacional Autónoma de México

The *Drosophila* genus of the family Drosophilidae comprises around 1600 described species. These species differ in their geographic distribution and ecologies, and consequently, in ecological niches and lifestyles. As they evolved in different environments, species may differentially regulate their metabolism and behavior as they adapt to these local conditions. Here, we characterize *Drosophila lutzii*, a Neotropical Floridosa group of species of *Drosophila*. As its group of species implies, they are not saprophytic, but rather feed on flowers. We made a comparative study between *D. lutzii*, a specialist, and sympatric *D. simulans*, a generalist. We analyzed metabolic and behavior parameters. *Drosophila simulans* is a saprophytic specialist, with feeding based on rotting plants and fruits, while *Drosophila lutzii* is a phytophagous specialist. We have found *D. lutzii* eggs, larvae, pupae and adults inside *Ipomoea sp.* flowers. This suggested a restricted diet, and thus, an interesting avenue for research in metabolism, in comparison to generalist species of flies. We found that freshly caught *D. lutzii* from the wild have higher carbohydrates levels, but similar lipid content, as compared to sympatric freshly caught *D. simulans*. Consistent with a restricted diet and specialist lifestyle, *D. lutzii* flies are less capable of surviving in culture in diets that differ in the amounts of carbohydrates, and when fed diets with high sugar concentrations, contrary to *D. simulans*, they significantly accumulate them. Triglycerides levels also were differentially affected in both species when fed with diets that varied in sugar content. *D. lutzii* flies are significantly and dramatically less motile, but possess a circadian activity rhythm akin to *D. melanogaster* or *D. simulans*. These three species showed a differential feeding behavior when exposed to food with different amounts of sugar in 30 minutes periods. Taken together, our results show that, in contrast to generalists, this specialist species, with more restricted habitat and feeding, is less capable of metabolic adjustments.

**793V** Genetic analysis of Juvenile hormone epoxide hydrolases in *Drosophila* Felipe Rogalski, Toshiro Aigaki Tokyo Metropolitan University, Tokyo, Japan

Juvenile hormone epoxide hydrolases (JHEHs) are enzymes that inactivate juvenile hormone (JH) in insects. The inactivation occurs through the hydrolysis of the epoxide group within the hormone into diol. JH’s main function is to ensure larval growth while preventing metamorphosis, but it also affects adult processes, such as egg production, fertility, and behavior. The genome of the fruit fly, *Drosophila melanogaster*, encodes three JHEHs: *Jheh1, Jheh2*, and *Jheh3*. It is not known whether those genes play a major role in JH related phenotypes. *Jheh1* and *Jheh2* have been repeatedly identified as oxidative stress response genes since they are induced by several xenobiotics, like parquat and caffeine. Recently, JHEH1 and JHEH2 have been shown to interact with a lipid droplet associated protein, suggesting a role in lipid metabolism. *Jheh3*’s function is completely unknown. To understand the roles of JHEHs in the fruit fly, we investigated the impact of genetic disruption of *Jheh*. We focused on *Jheh1* and *Jheh2* since they might be involved in resistance against stresses. Using the CRISPR/Cas-9 system, we have generated a mutant deleted for *Jheh1* and *Jheh2*. The mutant flies were viable, fertile, showed a normal lifespan and there was no difference in the resistance against parquat or caffeine between wild-type and Jheh1-Jheh2 KO flies. *Jheh1-Jheh2 KO* flies showed decreased body weight and reduced amount of triacylglycerol (TAG). Conversely, overexpression of *Jheh1* using the Gal4-UAS system increased TAG storage and body weight. JH is known to control developmental speed. *Jheh1-Jheh2 KO* flies had a slight delay in development, however, the expression level of *krüppel-homolog 1* (*kr-h1*), a JH-responsive gene, was not different between wild-type and mutant flies, suggesting that JH signaling was not involved in the mutant phenotype. To obtain insights into the metabolic processes of *Jheh* mutant flies, we performed metabolomic analysis using liquid chromatography-mass spectrometry (LC-MS). All glycolytic metabolites were decreased in *Jheh1-Jheh2 KO* flies, whereas TCA cycle metabolites remain normal. These results indicate that glycolytic activity is low in the mutants. In conclusion, our research shows that *Jheh1* and *Jheh2* are involved in the regulation of growth rate, glycolysis, and lipid metabolism, demonstrating that they are essential for energy homeostasis in *Drosophila*.

**794V** Developmental Exposure to the PFAS molecule, PFOA, alters Lipid Homeostasis in *Drosophila* Eric Kilbourn, Jason Tennessen Indiana University Bloomington

Perfluoroalkyl substances (PFAS) are man-made environmental contaminants that are now found on all seven continents. This family of compounds has been linked to heath issues such as liver cancers, decreased efficiency in vaccines, and obesity. The most prevalent of these compounds, perfluorooctanoic acid (PFOA), has a predicted environmental half life
of over 1000 years, meaning dozens of future generations be exposed to this molecule and the broad range of related PFAS chemicals. Despite the serious health concerns with PFAS, and PFOA in particular, relatively few mechanistic studies have examined the effects of PFAS in vivo. Here we address this deficiency by examining the effects of PFOA on Drosophila melanogaster. Our preliminary results reveal that flies readily absorb PFOA from dietary sources and accumulate this molecule to detectable levels. Moreover, our preliminary results reveal that developmental exposure of PFOA induces abnormal lipid droplet accumulation in the larval fat body. We are now examining the effects of PFOA exposure on both triglyceride metabolism as well as the metabolism of other lipid species. Overall, our data shows that D. melanogaster is a promising model for the investigating the effects of PFAS exposure on human health.

**795V Exploring pathophysiology in long-lived fly populations reared on two diets** Utsav Nyachhyon, Thomas Rundell, Laura Musselman Binghamton University

Overnutrition in Drosophila melanogaster and in humans causes pathophysiology including increased infection susceptibility, heart failure, insulin resistance, hyperglycemia, and reduced lifespan that are associated with obesity and other markers of metabolic disease. We used Drosophila as a model organism to identify genes that contribute to fitness and longevity during overnutrition. A large outbred population was selected for the ability of flies to survive chronic overnutrition conditions of high-sugar feeding. Every generation, half of the population that survived were bred to produce the next generation. Over time, the lifespan of high-calorie-adapted flies was extended, compared to those fed control diets. We hypothesize that improved heart function and/or immunity of the high-calorie-adapted flies might contribute to their increased survival. To test this hypothesis, these adapted flies are being subjected to cardiac stress and infection studies. In cardiac pacing, the flies are electrically stimulated to quantify the rate of heart failure and recovery. In infection studies, the flies are inoculated with Pseudomonas aeruginosa to quantify their infection susceptibility. Heart and immune function are being examined in control and adapted populations to search for mechanisms by which increases in the lifespan and healthspan may have occurred during selection. Significant changes in pathophysiology of adapted populations in comparison to control populations will be explored further by the ID of genes associated with these changes. Future studies will test the role of these genes which may be novel, conserved regulators of fitness during overnutrition in animals.

**796V Optimisation of macro- to micronutrient balance for larval growth on a holidic diet** Sebastian Sorge, Panayotis Pachnis, Vanessa Tixier, Alex Gould The Francis Crick Institute, London

The contributions of individual macro- and micronutrients to Drosophila melanogaster phenotypes can be untangled using holidic diets. Currently available holidic diets are useful for studying adult nutrition and lifespan but larval growth and/or development are substantially slower than on standard yeast-based diets. We previously published a chemically-defined diet (CDD) related to other published holidic diets and with a similarly poor larval performance. We now describe semi-systematic improvements to our CDD that substantially speed up larval growth and development, close to values obtained with a yeast-based diet. Our results confirm that amino-acid composition influences developmental speed but we also find that the precise macro- to micro-nutrient balance is of similar importance. Several non-essential micronutrients contribute to larval developmental progression to surprising degrees and even small changes in the nutrient balance can substantially perturb development. Together, these CDD studies help to define a narrow window of macro-to-micronutrient balance that is optimal for larval development. As a proof-of-principle, we apply the optimised CDD to larval stable isotope tracing experiments, determining the sugar carbon contribution to hemolymph polar metabolites and to de-novo synthesized fatty acids.

**797V The conquest of a new habitat: A study of the nutritional and sensory adaptations of the D. suzukii larvae.** Diego Galagovsky, Ana Depetris-Chauvin, Markus Knaden, Bill Hansson Max Planck Institute for Chemical Ecology

Animals need to feed to incorporate nutrients –protein, carbohydrates, and lipids, to use as source of energy and as building blocks to maintain their bodies. However, feeding substrates contain nutrients in different proportions; therefore, depending on the specific needs of an organism, food sources will have varying nutritional quality. To assess their needs and feed accordingly, animals have both mechanisms to evaluate their internal needs, and mechanisms to sense for chemical cues in foods that hint at their nutrient content. However, for some organisms, it is not their decision where to feed, but that of their parents.

In the life cycle of Drosophila flies, we can distinguish two main feeding stages—the larva and the adult, with different objectives and nutritional needs. The larva feeds mainly to grow and accumulate resources for metamorphosis while the adult needs nutrients to survive and achieve reproduction. Adult D. melanogaster flies prefer to feed and lay eggs on protein-rich overripe fruits. These make a good feeding substrate for the larvae. Conversely, while D. suzukii adult flies also feed on overripe fruit, they are physically and behaviorally adapted to lay eggs on ripening fruits, which are harder and have a high sugar-protein ratio. D. suzukii larvae have to face the harsh nutritional landscape that has resulted from this novel adult behavior. How do they do it? Are the larvae physically adapted to this medium? How have their nutritional needs, their behavior and their physiology adapted to the shift in adult ecology?
I am addressing these questions at different levels; the nutritional composition of host fruits, the larval behavior, and the larval physiology.

Our results indicate that the environment in which D. suzukii larvae develop, the ripening fruit picked by their mother, is not a temporally homogeneous environment. It starts out as a sugar-rich, protein-poor environment, in which D. melanogaster larvae cannot survive. As the D. suzukii larvae process it, the protein content of the fruit increases. Their presence and activity also stimulate the fermentation of the fruit. By the time the larvae reach the late stages, the feeding substrate has become rich in proteins, enough to sustain D. melanogaster development. We have also observed that, compared to D. melanogaster, the 1st instar larvae of D. suzukii have massive mouth hooks, which aid them in processing the hard ripening fruit. Do D. suzuki larvae have special behaviors associated to the use of their physical adaptations? How does their physiology deal with the composition of the medium before they convert it? We are currently exploring the behavioral and physiological characteristics of the early larvae that allow them to develop in the ripening fruit.

798V  Oxidative stress resistance in insulin-signaling impaired male and female Drosophila melanogaster Jessica Alvarez, Juan Rafael Riesgo-Escovar UNAM, Queretaro, Mx

Diabetes encompasses a group of metabolic disorders characterized by disglycemia, caused by insufficient insulin production or its inefficient signaling, and entails detrimental consequences for the patient’s quality of life. Drosophila melanogaster has an evolutionarily conserved insulin signaling pathway. We set to understand how an organismal deficiency in insulin signaling in adult male and female flies affects different aspects of the fly’s life, particularly the response to oxidative stress.

We exposed InR (the insulin receptor fly homolog) and dS6K (the ribosomal protein S6 kinase beta-1 fly homolog, under the control of TORC1, also regulated by insulin signaling) heteroallelic mutants, as well as wildtype controls with the same genetic background, to different prooxidant compounds. We found that females have higher lipid and carbohydrate accumulation and show better resistance to oxidative stress (H₂O₂) than males. A heterozygous mutation in Keap1 (a negative regulator of CncC, a master regulator of the antioxidant response) only rescued the phenotype of mutant InR males. We conclude that the type of oxidant fed to the fly and the genetic manipulation of the antioxidant response affected differentially the survival of males and females.

799V  The Role of Copper in Parkinson’s Disease Jessica Burkhart, Alysia Vrailas-Mortimer Illinois State University

Parkinson’s Disease (PD) is a neurodegenerative disease caused by the death of dopaminergic neurons in the substantia nigra region of the brain. PD is characterized by the presence of dysfunctional mitochondria and increased levels of oxidative stress. Though a handful of genes, such as parkin and PINK1, have been identified in familial forms of PD, most cases are sporadic. Therefore, it is thought that environmental factors may act on genetic risk factors to promote disease onset. Therefore, we are exploring the relationship between copper toxicity, which has been linked to other neurological disorders, and parkin and PINK1. We are testing the effect of environmental exposure to copper as well as altering copper levels genetically by manipulating the copper transporter ATP7, which is mutated in the neurodegenerative disorder, Menkes disease. Preliminary findings have shown that increased extracellular copper, from over-expression of ATP7, in conjunction with knockdown of parkin and PINK1 exasperate Parkinson’s symptoms.

800V  Determining the mechanism of anesthetic-induced neurotoxicity in a Drosophila model of mitochondrial disease Zachariah Olufs, David Wassarman, Misha Perouansky University of Wisconsin - Madison, Madison, WI

General anesthetics are broadly used and are safe in healthy individuals. However, adverse sequelae collectively referred to as anesthetic-induced neurotoxicity (AiN) have been reported in pre-clinical animal models and clinical studies of mitochondrial disease. Leigh Syndrome (LS) is the most common pediatric manifestation of mitochondrial disease. Over 75 distinct genes underlie LS, including mutations in Complex I of the mitochondrial electron transport chain. To investigate the mechanism of AiN, we have characterized flies carrying a mutation in ND23, which encodes a core subunit of Complex I and is orthologous to mammalian NDUFS8. We previously found that ND23 mutants are behaviorally sensitive to two volatile general anesthetics (VGAs), isoflurane (Iso) and sevoflurane (Sevo). Furthermore, Iso, but not Sevo, causes mortality within 24 hours of exposure, the extent of which is dependent on oxygen concentration ([O₂]) during exposure; low [O₂] (hypoxia) reduces mortality and high [O₂] (hyperoxia) increases mortality. To identify genes involved in the differential response of ND23 mutants to different anesthetics and to different oxygen concentrations, we performed RNA-seq analysis of fly heads. We found that Iso and Sevo differentially affected expression of chaperone and antioxidant genes, but high and low [O₂] did not appreciably influence gene expression. To test whether differential expression of chaperones or antioxidant genes is responsible for the difference in mortality between Iso- and Sevo-exposed flies, we used orthologous approaches to induce the transcriptional responses prior to anesthesia exposure, since exposure to a non-injurious stress might initiate protective mechanisms that would result in a state of tolerance to the subsequent injurious stress. We found that pre-exposure to Sevo or heat shock did not reduce mortality from a subsequent exposure to Iso. These data argue against a role for transcriptional activation of chaperone and antioxidant genes in the mortality mechanism. We are currently testing other non-injurious stressors as well as using loss-of-function
801V Role of peroxisome in mitochondrial dynamics during aging in Drosophila melanogaster Ankur Kumar, Pham Vo, Peiduo Liu, Hua Bai Iowa State University

Damaged mitochondria are repaired and recycled through the mechanisms of mitochondrial dynamics in response to stress; this helps in restoring cellular homeostasis. Mitochondrial dynamics have emerged as a novel regulator of aging in recent years. During aging, alterations in mitochondrial morphology and structure have been observed. Researchers have performed genetic manipulations of genes involved in the fission and fusion of mitochondria, which extended the lifespan. However, the causes of the age-dependent alteration in mitochondrial dynamics remain unanswered. Our focus is to explore the involvement of the peroxisome in maintaining mitochondrial homeostasis during animal aging. Recent studies in our lab have shown mitochondrial morphology and function alteration due to impaired peroxisomal protein import in aging oenocytes (hepatocytes) of fruit flies. We found an increase in mitochondrial size in oenocytes during fly aging. Similarly, we have found that the knockdown of Pex5, a peroxisomal import protein, alters mitochondrial morphology. Interestingly, we also have found that peroxisomal plasmalogen level decreases in aging flies, and knocking down the genes involved in peroxisomal plasmalogen synthesis, such as GNPAT (Glyceronephosphate O-acyltransferase) resulted in enlarged mitochondria. Alterations in other peroxisomal genes also have changed the mitochondrial morphology. Thus, our future goal is to understand how peroxisomes contribute to age-related alterations of mitochondrial dynamics and functions.

802V Transcriptional regulator of DR responsive genes extends lifespan and regulate Tau pathology in Drosophila Sudipta Bar, Patrick Wai-Lun Li, Tyler Hilsabeck, Guiping Du, Simon Melov, Pankaj Kapahi Buck Institute for research on aging

Dietary restriction (DR) is a robust intervention that slows aging in multiple species and staves off age-related diseases, including Alzheimer’s disease and related dementias (ADRDs). Despite identifying some of the genetic risk factors for Alzheimer’s disease (AD), therapeutics to treat ADRDs have been unsuccessful. However, the underlying mechanisms by which DR protects against AD are unknown. Identifying the mechanism behind the DR-mediated neuroprotection and lifespan extension can lead us to new therapeutic targets for AD. Using Translating Ribosome affinity purification (TRAP) method we have identified tissue-specific gene expression profiles of Drosophila in Ad libitum (AL) and DR conditions. We have identified a relatively large proportion of the unique DR-responsive genes to be altered in neurons. Utilizing bioinformatics approaches and publicly available chromatin immunoprecipitation sequencing (ChIP-seq) data from the ENCODE consortium we identified transcription factors (TFs) that are significantly enriched at promoter regions of genes that are upregulated or downregulated in fly brains on DR, compared to flies reared on AL diet. Screening of those TFs in lifespan extension revealed downregulation of Jarid2 extends lifespan and rescues tau pathology in a fly model of AD. We determined there are 359 downstream target genes of Jarid2 among which homolog of 16 genes have been reported to influence AD risk. Gene ontology (GO) enrichment analysis of the Jarid2 downstream targets identified enrichment of several biological processes including system development, nervous system development, neurogenesis, axon development, etc. We identified the downregulation of a neurogenesis factor NetB, a downstream target of Jarid2, which abrogates the DR effect. Hence we are working to understand the role of Jarid2 and NetB in neuroprotection to establish as potential targets for AD.

803V The bestrophin-1 chloride channel is required in the Malpighian tubules and hindgut for osmoregulation in response to high salt diet John Pleinis, Sima Jonusaitė, Katherine Beebe, Daryl Morrison, Forest Streeter, Jacob Hudac, Austin Goodwin, Ashlee Roberts, Jeffrey Schellinger, Adrian Rothenfluh, Aylin Rodan University of Utah

The mechanisms allowing Drosophila melanogaster to tolerate a high-salt diet are poorly understood. Here, we study flies carrying a mutation in the chloride channel, Bestrophin-1 (Best1). Flies with a loss-of-function mutation in Best1, Best1<sup>Δ340 Δ66</sup>, have increased lethality when fed a high-salt (0.3 M NaCl) diet. High salt lethality is rescued in mutant flies carrying a genomic rescue construct, or with expression of wild-type Best1 in the Malpighian tubules and the rectal pads of the hindgut, the ionic- and osmoregulatory epithelia of insects. Malpighian tubules from Best1 mutants have impaired fluid secretion and ion flux. This is likely due to impaired ability to recycle chloride entering through the sodium-potassium-2-chloride cotransporter (NKCC), as there is no effect of Best1 knockdown in flies carrying a loss-of-function mutation in the NKCC-encoding Ncc69. Best1 mutants have lower hemolymph osmolality when fed a normal diet, and fail to increase osmolality as much as controls when fed a high-salt diet. This phenotype is also rescued by the genomic rescue construct, as well as by expression of Best1 in the Malpighian tubule and rectal pads. High salt diet leads to dehydration in control flies, and this is further exacerbated in the Best1 mutants, likely due to greater osmotic loss of water from the hemolymph into the high osmolar gut lumen, as suggested by increased excretion and excreta osmolality in Best1 mutants. Analysis using specific GAL4 lines indicates that coordinated action of the Malpighian tubules and the hindgut is required to modulate excretion and dehydration. Metabolic analysis indicates that urea is the most upregulated solute in control flies, whereas Best1 mutants fail to upregulate urea. Accordingly, feeding Best1 mutants
urea partially rescues high salt lethality, while having no effect on survival in control flies. Best1 mutants have seizures, as occurs in patients with tonicity disorders. Best1 expression in the Malpighian tubules and rectal pads rescues the seizure phenotype, indicating that it is due to the osmoregulatory defects, whereas Best1 knockdown in neurons or glia does not result in seizures. Finally, Best1 mutants sleep less, likely due to searching for water, as water supplementation restores sleep. Thus, Best1 plays an essential role in the Malpighian tubules and hindgut to allow Drosophila to cope with salt stress.

804V The epicuticular lipid barrier is highly dynamic across the life course in Drosophila Lena Lampe1, Clare L Newell1, Ian Gilmore2, Alex P Gould1 1) Francis Crick Institute; 2) National Physical Laboratory

An epicutaneous lipid barrier is found in all terrestrial organisms and protects them from dehydration and other potentially harmful stresses in the environment. The high surface area to volume ratio of insects means that an effective epicuticular lipid barrier to dessication is particularly important. The composition of the barrier lipid blend is highly complex and, although hydrocarbons have been extensively analysed, other cuticular lipid species are less well investigated. Here we use a new surface-specific technique for mass spectrometry, OrbiSIMS, to characterize the cuticular lipid repertoire of Drosophila melanogaster during ageing. We identify epicuticular ceramides, wax esters, phospholipids and free fatty acids and show that they have specific temporal profiles that change substantially during ageing. Using stable isotope tracing, we measure the turnover rates of epicuticular lipids and show that larval nutritional resources contribute greatly to some but not all molecular species. This study shows how larval nutrition contributes to the adult epicuticular lipid blend and the quality of the dessication barrier.

805V Endogenous degradation of hormones by two distinct classes of enzymes uniquely impact coordinated animal growth and development Rebecca Spokony1, Lacy Barton2, Ruth Lehmann3 1) Baruch College, CUNY, New York, NY; 2) Skirball Institute at NYU School of Medicine, New York, NY; 3) The Whitehead Institute at MIT, Boston, MA

Juvenile hormones (JHs) are central coordinators of insect development. These potent small molecules regulate developmental progression through different tissues as well as respond to changes in external conditions. JH titers are dynamic throughout development and the bioavailability of JH is presumed to be tightly controlled by dynamic expression of enzymes responsible for its synthesis and degradation. Two major classes of JH degradation enzymes are Juvenile hormone esterase (Jhe) and Juvenile hormone epoxide hydrolase (Jheh). Due to the importance of JH regulation, insecticides have been designed to block their functions to inhibit the maturation of insect pests and disease vectors.

The endogenous roles of these two distinct classes of JH degradation enzymes in Drosophila are not understood. This knowledge gap may arise from the fact that the D. melanogaster genome encodes two Jhe genes and three Jheh genes. Using CRISPR, we generated Jhe, JheDup null double mutant animals and jheh1,2,3 null mutant animals to understand the functions of these two classes of enzymes. We find that while both jhe,jheDup and jheh1,2,3 larvae exhibit delayed pupariation, the delayed maturation in jheh1,2,3 triple mutant animals is much more dramatic, pupariating three days later than matched wildtype larvae. Interestingly, jhe,jheDup pupae and adults are significantly larger than matched wildtype animals, while jheh1,2,3 animals are smaller. These findings suggest that while all these enzymes can degrade JHs, each class has tissue-specific functions that impact coordinated development and growth. We are currently investigating this exciting postulate, as well as compensatory feedback mechanisms across JH enzymatic classes. Our findings are consistent with global knockdown studies in Bombyx mori2, Leptinotarsa decemlineata3, and Nilaparvata lugens4. Thus, the novel genetic tools we generated in Drosophila will allow us to harness the powerful of this model organism to understand the spatiotemporal and tissue-specific functions of these important enzymes.

References:

806V Impacts of Intestinal Occluding Junction Modulation on Non-Cell Autonomous Hallmarks of Aging Anna Salazar, Cody Sessions, Kayla Pauley, Sierra Boyce Christopher Newport University, Newport News, Virginia

Intestinal barrier function is tightly linked to longevity in Drosophila melanogaster and other organisms. We have previously shown that altered expression of occluding junctions in the guts of fruit flies can lead to various hallmarks of aging, including modulation of intestinal homeostasis, variations in microbial dynamics, changes in immune activity, and alterations in lifespan. Loss of a specific occluding junction, Snakeskin (Ssk), leads to rapid and reversible intestinal
barrier dysfunction, altered gut morphology, dysbiosis, and a dramatically reduced lifespan. Remarkably, restoration of Ssk expression in flies showing intestinal barrier dysfunction rescues each of these phenotypes previously linked to aging. Intestinal up-regulation of Ssk protects against microbial translocation following oral infection with pathogenic bacteria. Furthermore, intestinal up-regulation of Ssk improves intestinal barrier function during aging, limits dysbiosis, and extends lifespan. Additionally, perturbing barrier function in the gut has non-cell-autonomous impacts, including alterations in the brain and muscle. Moreover, these analyses add more information about the impact of the gut on tissue outside the gut and begin to address communication between the gut and the brain and muscles in disease models. These findings indicate that intestinal occluding junctions may represent prolongevity targets in mammals, in addition to their possible roles in intestinal dysfunction, aging, and disease. They also attempt to gain a better understanding of the impact of intestinal dysfunction on cells outside of the gut.

807V Single-nucleus RNA-seq of Drosophila Thorax Post Exercise Treatment: Pilot Study Bre Minniefield, Nicole Riddle University of Alabama at Birmingham

Physical activity in any form can significantly decrease the incidence of various diseases and benefit overall health. However, we are unable to recommend optimized exercise treatments for a specific individual’s health needs. Despite significant progress, we lack a thorough understanding of the biological pathways that are activated or impacted by exercise. For example, little is known about how individual tissues are affected by exercise treatments and which cell- and tissue-level changes in response to exercise. Additionally, little is known about the pathway(s) that mediate short- and long-term exercise responses. Identifying molecular mediators of exercise response has the potential to provide trackable factors to better predict the outcomes of exercise for an individual, thus, allowing for optimized and personalized exercise treatments.

Here, we use a Drosophila exercise model to gain an understanding of the cell- and tissue-level changes precipitated by a short-term exercise treatment. Rotational stimulation by rotating the fly enclosures has been shown to increase animal activity in Drosophila, modeling exercise. Depending on genotype, animals treated with rotational stimulation show increases in activity up to 10-fold, and this increased activity alters gene expression, metabolite levels, and body condition. Using our established protocols with the TreadWheel exercise system, here, we carry out a single-nucleus transcriptome study focused on muscle to compare exercise-treated and control animals. Specifically, we use a wild-derived strain, DGRP line 304, which in previous studies showed a significant change in activity level with rotation and a change in weight as response to exercise in both sexes. Exercise-treated animals experienced five consecutive days of rotational stimulation for 2hrs, while control animals do not. Post exercise, thorax dissections were performed, a body part which consists of ~95% muscle. Using 10xGenomics technology, we collect single-nucleus transcriptome profiles from this tissue. We use these profiles to identify expression patterns, key cell types, and molecular pathways that are 1) differing between exercised and controlled flies; and 2) significantly affected by exercise. The analysis is in progress, and results will be reported. Discovering the key elements of exercise response will provide the foundational data to develop effective, personalized exercise plans, including alternatives for individuals unable to exercise.

808V dFNDC5 Regulates Exercise Performance and Adaptations in Drosophila Tyler Cobb1, Jun Hee Lee2, Myungjin Kim3, Robert Wessells1 1) Wayne State University; 2) University of Michigan

Endurance exercise has profound benefits on health and mobility, but the mechanisms underlying these benefits are not fully understood. Identification of targets that mediate the benefits of exercise can be useful for the development of therapies to improve health in populations that cannot exercise or display exercise intolerance. We have developed a model to exercise train Drosophila and have shown that many adaptations in mobility, metabolism, and gene expression are conserved. Using this model, we have identified dFNDC5 as a key mediator of endurance exercise performance and adaptations. dFNDC5 is a homolog of mammalian FNDC5, which encodes a transmembrane protein expressed in muscle that is cleaved during exercise to give rise to a peptide called Irisin that regulates systemic adaptations in mammals. However, its role in exercise performance and endurance adaptation has not been studied in depth. We show that dFNDC5 mutants have impaired baseline endurance and do not adapt to chronic exercise training. Tissue-specific experiments reveal that dFNDC5 is required in muscle for normal baseline running endurance and for the ability to adapt to exercise. We also show muscle-specific dFNDC5 overexpression extends baseline endurance and is sufficient to promote mobility adaptations in unexercised flies. These findings suggest a conserved role of dFNDC5 in the exercise response.


Serotonergic neurons produce extensively branched axons that fill most of the central nervous system, where they modulate behaviors such as mood, sleep, appetite, locomotion, and cognition. Proper behavioral output therefore depends on the precise outgrowth and targeting of serotonergic axons during development. To assist in this process, serotonergic neurons utilize serotonin as a trophic signaling molecule prior to it assuming its canonical role as a
neurotransmitter. This process, termed autoregulation, plays a negative role in *Drosophila* by limiting axon outgrowth, branching, and varicosity development. Yet the underlying mechanism of serotonin autoregulation remains unknown. In non-serotonergic neurons, activation of serotonin receptors causes downstream signaling events that are linked with F-actin depolymerization, growth cone collapse, and reduced neurite outgrowth. We therefore hypothesized that serotonin autoreceptors initiate autoregulation in serotonergic neurons. To test this, we adapted a primary neuron culture system in which *Drosophila* serotonergic neurons could be grown and unambiguously identified using Gal4-dependent expression of the fluorescent protein tdTomato. Next, we initiated autoregulation by applying exogenous serotonin to the culture. We found reduced axon outgrowth and branching, suggesting serotonergic neurons in culture respond similarly as previously reported *in vivo*. Lastly, we used pharmacological activation of serotonin autoreceptors to test their role in autoregulation. Axons treated with receptor agonists showed reduced outgrowth and branching, thereby mimicking the effects of serotonin application. This suggests that serotonin initiates autoregulation through activation of autoreceptors.

**810B** Investigating mechanisms of Frazzled/Dcc signaling in axon guidance *Sarah Gagnon*, Yixin Zang, Greg J. Bashaw University of Pennsylvania, Philadelphia, PA, USA

Precise wiring of neural circuits is essential for the execution of vital functions, from breathing to complex motor movements. This connectivity is established during development as the axons of newborn neurons extend through the embryonic environment, sensing both attractive and repulsive cues via guidance receptors. In the central nervous system (CNS), commissural neurons send their axons across the midline to connect to contralateral targets. This is essential for left-right coordination, yet the underlying mechanisms are not fully understood. In flies, the guidance receptor Frazzled (Fra), like its human ortholog Deleted in Colorectal Cancer (Dcc), promotes axon growth toward the midline. Despite its critical role in axon guidance, our understanding of Fra/Dcc signaling remains incomplete. We performed an affinity purification-mass spectrometry screen in fly embryonic neurons to identify novel Fra interactors. We selected three candidates, the Toll-like receptor family member Toll-7, the iRhom pseudoprotease family member rhomboid-5 (rho-5), and the exportin embargoed (emb), all expressed in the embryonic CNS, for further investigation. Toll-7 regulates neuronal connectivity in the fly olfactory system, but its involvement in Fra signaling has not been investigated. Human iRhoms promote ADAM17 metalloprotease activity, which was shown by our lab to cleave and positively regulate the functions of both Fra and Dcc, making rho-5 a promising candidate regulator of Fra signaling. Finally, as nuclear export antagonizes the transcriptional activity of Fra, emb may also modulate Fra signaling. Our goal is to characterize the roles of Toll-7, rho-5, and emb in Fra and Dcc signaling. We show that Toll-7 and emb biochemically interact with Fra in S2R+ cells. Importantly, we find that all three candidates modulate midline crossing: Toll-7 and rho-5 promote crossing, while emb antagonizes this process. Given their physical association with Fra, it is likely that the candidates regulate midline crossing via the Fra pathway. Future experiments will determine whether Toll-7, rho-5 and emb function in the Fra pathway and if these genes play conserved roles in vertebrate Dcc signaling. Our research will shed light on uncharacterized mechanisms of Fra/Dcc signaling and axon guidance in *Drosophila* and vertebrates. These findings will provide insight into the mechanisms driving assembly of neural circuits, which may inform interventions to promote neuronal repair.

**811C** Developmental axon guidance cues are critical for adult neuronal survival and function *Aarya Vaikakkara Chithran*, Douglas Allan, Timothy O’Connor Department of Cellular and Physiological Sciences, University of British Columbia, Vancouver, BC, Canada.

One of the most extraordinary characteristics of the nervous system is the complexity of its neural circuits. Although there are many mechanisms involved in shaping the pattern of these connections, one of the most important events is the guidance of axons to their specific targets. Model organism research in the past several decades has advanced our understanding of the function of numerous guidance cues, their receptors and their underlying signaling cascades. More recently it has become clear that after functional circuits have been established, many neurons continue to express developmental guidance cues. The expression of these genes in the adult indicates that there are likely additional roles for them beyond the initial phase of neuronal process outgrowth, growth cone navigation and target innervation.

To test this hypothesis, we performed an RNAi screen against axon guidance genes expressed in the adult nervous system. Using available databases, 151 axon guidance genes were identified that are expressed in the adult *Drosophila* nervous system. 44 genes were prioritized based on their higher expression profiles and previously known roles in neuronal pathfinding. We knocked down the expression of each of these genes using spatial and temporal control of the GAL4-UAS system and identified 15 genes that are required for adult survival. Knockdown of these 15 guidance cues also caused motility defects.

To understand the impact of axon guidance gene knockdown at a cellular level, we examined well-established circuits such as the olfactory system and motoneuron circuits in adult *Drosophila*. Knocking down Fasciclin-3, an Ig containing homophilic cell adhesion molecule, led to neuronal death in the adult olfactory system. This phenotype was rescued
by overexpressing p35, an anti-apoptotic protein. Knocking down Semaphorins and Plexins, members of the canonical repulsive guidance signalling pathway, resulted in loss of motoneurons. Currently, we are examining the impact of Semaphorin/ Plexin double knockdowns to understand the cellular and molecular mechanism responsible for these phenotypes. We are also examining the impact of Semaphorin/ Plexin knockdown on the olfactory circuit and neuromuscular junctions.

Taken together, we believe that the continued expression of axon guidance genes in the adult is critical for the maintenance of neural circuits in the mature nervous system.

812A  Target-independent visual map formation Egemen Agi1, Charlotte B. Wit1, Eric Reifenstein2,3, Max von Kleist2,3, P. Robin Hiesinger1 1) Division of Neurobiology, Institute for Biology, Freie Universitaet Berlin, Germany; 2) Systems Medicine of Infectious Disease (PS), Robert Koch Institute, Berlin, Germany; 3) Bioinformatics (MF1), Robert Koch Institute, Berlin, Germany

The fly visual map represents ~800 neighboring points in space as synaptic ensembles (cartridges) in the lamina of the fly brain. These 800 visual axes are represented by six photoreceptor axons (R1-R6) in each cartridge, where R1-R6 provide pooled presynaptic input onto their main postsynaptic partner lamina cells L1-L3. Due to the arrangement of separate light-sensing elements that receive input from different visual axes in each single ommatidium of the fly eye, R1-R6 axons from six different ommatidia have to grow towards a shared cartridge in the lamina, a principle called ‘neural superposition’. In effect, six times 800 R1-R6 axons synchronously grow away from their original arrival points in the lamina by elongating their growth cones orthogonally to their axons in a sorting plane. At the time of growth cone sorting, the postsynaptic L cells are present in the lamina in an apparent target grid. Surprisingly, genetic ablation of L cells does not affect early pattern formation, growth cone elongation nor the initial formation of the neural superposition pattern. Only after superposition patterning through R1-6 growth cones alone, the fronts of all elongated growth cones are stabilized by the ‘target grid’ in an N-Cadherin-dependent manner. To understand how initial neural superposition pattern formation is established by six times 800 growth cones in the absence of any other cell type, we performed non-invasive two-color two-photon live imaging and developed a computational model of the process based on photoreceptor filopodial dynamics. Both data and model reveal a dynamic pattern formed by the ‘heels’ (back ends) of the elongating growth cones with shifting membrane densities throughout the sorting process. Front filopodia predominantly sample the space between the highest heel densities, i.e. the valleys of the heel density landscape. In turn, the weighted directions and lifetimes of front filopodia predict the subsequent growth cone elongation vectors. We conclude that six times 800 photoreceptor growth cones self-organize into the neural superposition pattern by separating growth cone front and heel patterns, with the former appearing to chase the latter. This self-organization process can be explained and modeled based on a minimal number of guidance cues by largely relying on intrinsic and dynamic cell and tissue properties.

813B  Temporal regulation of nicotinic acetylcholine receptor subunits supports central cholinergic synapse development in Drosophila Justin Rosenthal1,2, Jun Yin1, Jimmy Lei1, Anupama Sathyamurthy1, Jacob Short1, Caixia Long1, Emma Spillman1, Chengyu Sheng1, Quan Yuan1 1) National Institutes of Health, Bethesda, MD; 2) University of Maryland-College Park, College Park, MD; 3) University of California-San Diego, San Diego, CA

The construction and maturation of the postsynaptic apparatus are crucial for synapse and dendrite development. The fundamental mechanisms underlying these processes are most often studied in glutamatergic central synapses in vertebrates. Whether the same principles apply to excitatory cholinergic synapses, such as those found in the insect central nervous system, is not known. To address this question, we investigated a group of projection neurons in the Drosophila larval visual system, the ventral lateral neurons (LNvs), and identified Da1 and Da6 as the main functional nicotinic acetylcholine receptor (nAChR) subunits in the larval LNvs. Using morphological analyses and calcium imaging studies, we demonstrated critical roles of these two subunits in supporting dendrite morphogenesis and synaptic transmission. Furthermore, our developmental expression profiling and endogenous tagging approaches identified distinct transcriptional controls over the two subunits resulting in the up-regulation of Da1, which is suppressed by elevated presynaptic activity, and down-regulation of Da6 during larval development. Additional functional analyses of synapse formation and dendrite dynamics further revealed a close association between the temporal regulation of these two nAChR subunits and their sequential requirements during cholinergic synapse maturation. Finally, preliminary screens have identified candidate regulatory factors coordinating nAChR expression and postsynaptic activity, including the transcription factor and activity-regulated gene Hr38 as well as the nAChR-specific molecular chaperone NACHO. Together, our findings highlight how transcriptional control of nAChR subunits is a core element of developmental and activity-dependent regulation of central cholinergic synapses, which is likely dependent on both transcriptional and post-translational processes.

814C  Differential expression of the roundabout 3 (Robo3) guidance receptor regulates interneuron dendrite morphogenesis in Drosophila melanogaster somatosensory circuit development Jake Henderson, Ellie Heckscher University of Chicago

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For any animal to sense and interact with their environment, it is critical their neural circuits develop properly. The development of functioning neural circuits requires each neuron to correctly position its axonal and dendritic arbors and establish synaptic connections with their correct partners. Within neural circuits, interneurons are the most numerous cell type and are required for complex processing of sensory stimuli. Interneurons are an under investigated cell type due to the lack of tools and knowledge regarding neural circuit architecture within the CNS. In Drosophila, recent studies have clarified unique synaptic connections for many circuits, however how these connections are determined for interneurons is unclear. Temporal transcription factors and guidance cues are known to regulate neuronal development, however the role of these molecular factors has not been deeply investigated in interneuron dendrite development, synaptic partner selection, and circuit formation. We hypothesize that differential guidance receptor expression establishes specific dendritic morphologies and synaptic connections within interneurons born from the same stem cell. We investigated this hypothesis using the Drosophila melanogaster Neuroblast 3-3 stem cell lineage as a model. The NB3-3 lineage produces two cohorts of interneurons that create synaptic connections with two different sensory circuits: vibrational and proprioceptive. During development, the sensory neurons position their axons first at different locations within the ventral nerve cord. The two cohorts of even-skipped(+/−) lateral (EI) interneurons are then required to position their dendrites appropriately to receive the sensory input. We investigated the role of guidance receptors in interneuron dendrite morphogenesis by examining the expression pattern of Roundabout receptors within EI interneurons. We also examined the necessity and sufficiency of the Roundabout 3 (Robo3) receptor by manipulating its expression. These experiments revealed that Robo3 is involved in regulating EI interneuron dendrite position and therefore is a potential important factor in determining circuit wiring. This work will also enable future research to investigate the broad conservation of neuronal circuit specification and development of therapies for neurodevelopmental pathologies in which circuit wiring is disrupted.

815A  Promiscuous wiring via variable spatial sampling of an orderly array Emma Thornton-Kolbe, Maria Ahmed, E. Josie Clowney University of Michigan, Ann Arbor MI

The patterns of neuronal connectivity determined by development generate much of the computational power of the central nervous system. In brain regions devoted to sensory processing, sparse, combinatorial connectivity expands perceptual capacity. In the Drosophila mushroom body, sparse combinatorial connectivity between projection neurons carrying olfactory information from the periphery and Kenyon cells results in separable coding of many diverse scents and thus underlies successful associative learning. Each Kenyon cell (KC) receives input from a handful of projection neuron (PN) types through claw-like dendrites which each enwrap a single projection neuron bouton. Our lab has demonstrated that Kenyon cells determine connectivity density through the number of claws they produce, which are invariant when KC population is increased or reduced. Rather, PN bouton number changes to accommodate the changes in postsynaptic KCs. Here, we ask how the combinations of PN inputs to KCs are determined. Two possible sources of variation in projection neurons connections to Kenyon cells are molecular and spatial variation. By describing levels of transcriptional variation among projection neurons and Kenyon cells and the spatial relationships among these cells in connectomic data, we are testing the relative importance of these factors. Insights into the development of the mushroom body calyx will add to the existing knowledge of the diverse ways in which neuronal connectivity is controlled during development to produce functional circuits.

816B  Codes of cell surface proteins coordinate stochastic and deterministic cell fates during Drosophila color vision circuit assembly Yu-Chieh David Chen, Claude Desplan New York University, Department of Biology, New York, NY

How vast numbers of neurons are specified into correct cell fates and connected with their proper targets during development represents a fascinating area of developmental neuroscience. Mechanisms of stochastic and deterministic cell specification programs to achieve neuronal diversity have been extensively studied. A number of axon guidance molecules have also been identified. However, little is known about the coordination between neuronal specification and specific connectivity patterns, especially when two synaptic partners undergo two different modes of cell specification (stochastic vs. deterministic). In the fly retina, pale (p) and yellow (y) subtypes of color photoreceptors (R7 and R8) are stochastically specified, whereas their synaptic partners in the optic lobe are produced through highly deterministic programs. How do stochastically determined p vs. y R7 and R8 find their respective targets that are deterministically specified in the optic lobes?

By focusing on p/y R7 and their main downstream target p/y Dm8 neurons, we and others have recently identified a pair of DIPs and Dprs, members of an interacting network of immunoglobulin superfamily proteins, as critical regulators of the synaptic connection between yR7 that expresses Dpr11 and yDm8 that expresses DIPy. We thus hypothesize that different pairs of cell adhesion molecules can mediate the matching of other synaptic partners. We aimed to uncover such molecular code for the pairing of pR7 and pDm8. It is unlikely to be achieved via another DIP-Dpr pair since our recent single-cell RNA sequencing (scRNAseq) data showed no other DIPs expressed in Dm8s. To identify candidate molecule(s) that mediate pR7-pDm8 connectivity, we bioinformatically split our recent scRNAseq transcriptomic profile of the Dm8 cluster into DIPγ-(yDm8s) and DIPγ-(pDm8s) neurons during development. We found that Beat-IIIC, a
member of Beat family proteins, is specifically expressed in pDm8s cells during the developmental stage of synaptic partner pairing. Members of the Beat family and Side proteins form a receptor-ligand system to control axonal targeting. Since there are 14 Beat and 8 Side members, we aim to survey the expression pattern of all the Sides and look for their expression in pR7s. Preliminary experiments suggest Side-V is a promising candidate as it is only expressed in a subset (presumably p) of R7 terminals. We have generated CRISPR mutant alleles of these candidates and will be presenting our functional analyses of these candidate molecules in regulating synaptic partner matching. Overall, our work has uncovered novel molecular mechanisms regulating synaptic pairing and probes the fundamental principles underlying the propagation of stochastic cell fate choices during circuit assembly.

817C Investigation of the tRNA modifying enzyme, TRMT1, in neurodevelopment Sara Ríos Méndez1, Kimberly Rose Madhwanii, Ambar Delgado1, Jennifer Droumouchel9, Kate O’Connor-Giles3,5 1) NIH Post-Baccalaureate Research Education Program (PREP), Brown University, Providence RI; 2) Neuroscience Graduate Training Program, Brown University, Providence, RI; 3) Department of Neuroscience, Brown University, Providence, RI; 4) Molecular Pharmacology & Physiology Graduate Training Program, Brown University, Providence, RI; 5) Carney Institute for Brain Science, Brown University, Providence, RI

The dynamic regulation of global gene expression is required for proper development and function of the nervous system. Emerging work reveals that dysregulation of global gene expression can result in a myriad of neurodevelopmental and degenerative disorders. Specifically, transfer RNAs (tRNAs), which act as adaptors to decode messenger RNA and deliver the correct amino acid to growing polypeptides, undergo a variety of post-transcriptional modifications that are crucial for tRNA folding and stability, thereby ensuring accurate protein synthesis. tRNA methyltransferase 1 (TRMT1) is one of two mammalian orthologs of yeast tRNA methyltransferase, Trm1. TRMT1 dimethylates the guanosine nucleotide at position 26 of most tRNAs to yield the N2,N2-dimethylguanosine (m2,2G) modification. Frameshift mutations in human TRMT1 underlie autosomal recessive intellectual disability in multiple families, suggesting that guanosine-26 demethylation of tRNAs plays an important role in proper neurodevelopment and function. Potentially overlapping roles for TRMT1 and TRMT1-L complicate mammalian studies so we generated a model of TRMT1 in Drosophila, which encodes a single Trm1 homolog. Our preliminary studies suggest a role for TRMT1 in promoting synapse formation. We will report our latest findings on TRMT1’s in vivo role in nervous system development and function. Further study into TRMT1 will provide insight into the important role of tRNA modification in regulating gene expression in the nervous system.

818A Neurodevelopmental role of a tRNA methyltransferase implicated in intellectual disability Kimberly Rose Madhwanii, Sara Ríos Méndez1, Caley Hogan2, Jennifer L. Droumouchel, Jenna Lentini3, Dragony Fu4, Kate M. O’Connor-Giles5 1) Neuroscience Graduate Training Program, Brown University, Providence, RI; 2) NIH Post-Baccalaureate Research Education Program (PREP), Brown University, Providence RI; 3) Laboratory of Genetics, University of Wisconsin-Madison, Madison, WI; 4) Molecular Pharmacology & Physiology Graduate Training Program, Brown University, Providence, RI; 5) Department of Biology, University of Rochester, Rochester, NY; 6) Department of Neuroscience, Brown University, Providence, RI; 7) Carney Institute for Brain Science, Providence, RI

Gene regulation at multiple levels is critical for nervous system development and function. A number of mutations leading to global misregulation of gene expression disproportionately affect the nervous system, resulting in both neurodevelopmental and degenerative disorders. Emerging work demonstrates that post-transcriptional modification of transfer RNAs (tRNAs) regulates tRNA stability and codon-anticodon pairing and, thus, translation rate and fidelity. ALKBH8 is one of two metazoan homologs of the yeast tRNA methyltransferase TRM9. Mutations in the highly conserved human ALKBH8 were recently found to cause intellectual disability in three families. However, ALKBH8’s role in the nervous system is unknown. In yeast, TRM9 methylates uridines in the wobble position of the anticodon loop to reinforce cognate codon-anticodon pairings, resulting in increased translation of mRNAs enriched for cognate codons. We generated ALKBH8 null mutants and analyzed tRNA post-transcriptional modifications in the brain by mass spectrometry. We observed a complete loss of wobble uridine methylation in Drosophila ALKBH8 null mutants, consistent with prior findings in mammals. We next investigated a role for ALKBH8 in nervous system development and found that ALKBH8 attenuates synaptic growth. In addition, we found that ALKBH8 mutants exhibit increased levels of and sensitivity to reactive oxygen species (ROS). ALKBH8 is required to methylate tRNA-selenocysteine, which enables the incorporation of selenocysteine into selenoproteins. Selenoproteins are potent regulators of oxidative stress, and we have found that blocking selenoprotein synthesis by other means also results in synaptic overgrowth. Interestingly, oxidative stress has been shown to manifest in synaptic overgrowth. To determine if increased ROS and synaptic overgrowth in ALKBH8 mutants are causally linked, we treated ALKBH8 null animals with the antioxidant N-acetylcysteine amide and found that synaptic growth is partially rescued. These findings support a model in which ALKBH8 regulates synaptic growth through its role in regulating ROS via methylation of tRNA-selenocysteine, and reveals antioxidants as a potential therapeutic treatment for individuals with ALKBH8-linked intellectual disability.

819B Long-range temporal patterning of neuroblasts in the developing Drosophila medulla couples neurogenesis to circuit assembly Teddy Erclik, Urfa Arain, Maliha Islam, Priscilla Valentino University of Toronto - Mississauga
Cascades of transcription factors (TF) that are transiently expressed in neural stem cells are responsible for generating the enormous diversity of cell types in nervous systems. However, the gene-regulatory mechanisms that establish and maintain these cell fates in postmitotic neurons and instruct their specific morphology, connectivity and physiology remain largely unclear. Using a large scRNA-seq atlas of the *Drosophila* optic lobes (Özel et al. 2020, *Nature*), we tracked ~200 neuronal types across 6 stages and found that each one stably expresses a unique combination of 95 TFs (~10 per cell type) that are maintained throughout development to adulthood. We hypothesized that these function as “selector” TFs that are activated in each neuron immediately after their birth and function as top-level regulators of all type-specific gene expression thereafter. Through genetic gain- and loss-of-function experiments, we show that modification of these selector TF codes is sufficient to induce predictable switches of identity between various optic lobe neurons. For instance, *pdm3* is necessary and sufficient to determine the fate choice between Tm2 and Tm4 neurons. In addition to their morphological identity, scRNA-seq of perturbed neurons revealed that such conversions are also transcriptomically complete. Similarly, ectopic expression of *Vsx* genes in Mi15 neurons not only leads their complete morphological conversion to Dm2, but also a loss of their aminergic identity. Thus, continuously maintained TF codes instruct both the adult terminal features and the type-specific development of optic lobe neurons.

To understand the mechanisms that control brain wiring downstream of selector TFs, we combined scRNA and scATAC-seq (chromatin accessibility) datasets to build computational models of gene regulatory networks using Inferelator 3.0 (Gibbs et al. 2021, *BioRxiv*). Networks inferred at mid-pupal stages suggest that selector TFs interact with hormone-responsive TFs to activate a large repertoire of cell-surface proteins in each neuron prior to synapse formation. For instance, we show that Netrin receptor Frazzled, downstream of *pdm3*, specifically mediates the unique ‘forking’ of Tm2 dendrites: overexpression of *fra* in Tm4 neurons is sufficient to produce this feature.

Overall, our results provide a unified framework of how specific fates are maintained in postmitotic neurons, and open up new avenues to understand synaptic specificity through gene regulatory networks.
The number of neurons formed during development need to be tightly regulated to form functional neural circuits. Known strategies include regulating the number of neurons that are formed or that survive during development. Here, we study how neuronal numbers are regulated in the lamina neuropil of the visual system. The lamina is comprised of ~800 units or columns. Each column consists of six precursors, of which only five differentiate into the five lamina neuron types (L1-L5). The extra precursor is eliminated by apoptosis. This process is highly stereotyped with neuronal differentiation and apoptosis occurring invariably in the same cell positions in each column. We asked how exactly five lamina neurons (and neuron types) differentiate from a pool of six precursors. Previously we showed that L1-L4 neurons are induced to differentiate by wrapping glia. Therefore, we focused on the extra precursor and the differentiating L5s. We uncovered that differential Hedgehog signaling levels along the length of columns pattern lamina precursors, such that the 2 precursors experiencing the lowest levels of signaling are specified as L5s. Thus, twice as many precursors are specified as L5s than undergo differentiation normally. We showed that in response to photoreceptor-derived Spitz, a glial population, the outer chiasm giant glia (XgO), secrete multiple ligands (Spitz and Collagen IV alpha 1) to induce L5 differentiation. Moreover, we found that the newly-induced L5 neurons antagonize differentiation signals from reaching the extra precursors, which as a result undergo apoptosis. Therefore, our results indicate that newly induced neurons limit the availability of glial differentiation signals which sets the number of L5 neurons in each lamina column. Furthermore, our work highlights how stereotyped patterns of programmed cell death in the lamina arise from extrinsic signals which reliably pattern the development of the nervous system.

In the *Drosophila* visual system, photoreceptors in the compound eye project visual information to neurons in the underlying optic lobe for processing. The largest neuropil of the optic lobe is the medulla, which mediates motion and colour processing, and is comprised of over 90 cell types. The medulla develops from a larval crescent of neuroepithelial (NE) cells termed the outer proliferation centre (OPC). In the 3rd instar larva, the NE cells of the OPC are converted into neuroblasts, which then asymmetrically divide to generate the 90 types of medulla neurons. Previous studies have shown that two axes of positional information are required for neuronal specification. In the temporal axis, 5 transcription factors are sequentially expressed in the medulla neuroblasts as they age. In the spatial axis, the OPC is patterned into 5 spatial compartments by the non-overlapping expression of 3 homeobox transcription factors; Vsx1 in the center of the crescent, Optix in the adjacent arms and Rx in the tips. Thus, distinct neuronal types can be assigned unique spatio-temporal birth addresses based on the spatial compartment and temporal window from which they were born. Surprisingly, it has also been observed that neurons with identical spatio-temporal birth addresses can assume different fates based on whether they are born in the dorsal or ventral half of the OPC, suggesting that the dorsal and ventral halves of the OPC crescent are also spatially patterned. Here, using a technique involving micropipette-based cell isolation and RNA-seq, we identify 4 transcription factors that are differentially expressed in the dorsal and ventral halves of the OPC NE; salm and salr are expressed in all dorsal OPC cells, whereas disco and disco-r are expressed in all ventral cells. We show that the expression patterns of these genes meet to form a sharp dorsal-ventral boundary in the Vsx1 region of the OPC. We further demonstrate that salm/salr and disco/disco-r negatively cross-regulate each other in the OPC NE to maintain this dorsal-ventral compartment boundary. Loss- and gain-of-function experiments are currently underway to determine whether these genes contribute to medulla neuronal specification. It is anticipated that investigating the neurogenic role of these genes in the medulla will further our understanding of how neural diversity is generated at the intersection of temporal and spatial inputs.
Membrane trafficking pathways are essential to sculpting neuronal morphology. The closely related GARP (Golgi-Associated Retrograde Protein) and EARp (Endosome-Associated Recycling Protein) complexes are conserved membrane tethers that function in the secretory and endolysosomal pathways. These complexes share the proteins Vps51, Vps52, and Vps53, and each contain a complex-specific protein: Vps54 in GARP and Vps50 in EARp. Several mutations in the core components (Vps51 and Vps53) are associated with neurodevelopmental disorders, but a precise role for these complexes in neurodevelopment is still unclear. We generated CRISPR knockout Drosophila of the shared component Vps53, and the complex-specific components Vps50 and Vps54 (also known as scattered). We find that both complexes are required for dendrite morphogenesis at a specific time during development. Larval neurons are largely spared, but alterations in endolysosomes and dendrite morphology emerge during remodeling of peripheral sensory neurons during pupation. We also observe accumulation of sterol specifically in GARP-, but not EARp-deficient neurons. Previous studies of GARP-deficient cells found that sterol accumulates in lysosomes. To our surprise, we find that sterols accumulate at the trans-Golgi network (TGN), not in lysosomes, in Vps54<sup>KO/KO</sup> Drosophila neurons. As sterol auxotrophs, Drosophila obtain sterol from dietary sources only and therefore may utilize additional mechanisms to transfer sterol from the endolysosomal to the secretory pathway after uptake. Targeting genes that regulate sterols and related lipids at the TGN can modulate the Vps54<sup>KO/KO</sup> phenotype. Specifically, overexpressing the oxysterol binding protein (Osbp), which transfers sterol between the endolysosomal and secretory pathways at multiple membrane contact sites, exacerbates the dendrite regrowth defect of Vps54<sup>KO/KO</sup> neurons, while a null allele of Osbp partially rescues it. This suggests that the activity of Osbp may be upregulated in Vps54<sup>KO/KO</sup> neurons and that the accumulation of excess sterol at the TGN is, at least in part, responsible for inhibiting dendrite regrowth. These studies begin to functionally distinguish the GARP and EARp complexes in neurodevelopment and indicate that both membrane trafficking and lipid transfer pathways are essential for proper dendrite growth.

**826C  Genetic mechanisms underlying the development and distribution of Dm4 neurons in the Drosophila medulla Urfa Arain<sup>1</sup>, Ted Erclik<sup>1,2</sup>, 1) Cell & Systems Biology, University of Toronto, Toronto, Ontario, Canada ; 2) Biology, University of Toronto, Mississauga, Ontario, Canada**

The Drosophila compound eye is comprised of an array of photoreceptor units that send visual stimuli to the processing centers of the brain. Colour and motion processing occurs in the medulla, which is made up of repeating columnar units that are innervated by over 90 neuronal cell-types. During brain development, a subset of medulla neurons migrate from their spatially restricted birth locations to reach their final positions in the adult retinotopic circuit. To date, little is known about the mechanisms underlying this movement of medulla neurons. To elucidate the mechanisms that regulate this process, we have focused on the migration of a single neuronal cell-type, Dm4. There are approximately 40 Dm4 neurons per medulla, each of which cover ~20 columns across the antero-posterior and dorso-ventral axes of the adult retinotopic field. A single Dm4 neuron sends its axon into the M6 layer of the medulla neuropil, and forms dendritic arbors in the superficial M3 layer to make synaptic contacts with lamina monopolar cells (L3). To understand how these circuits are established, we generated a Dm4-specific split-Gal4 driver that allows us to track the relationship of Dm4 movement with morphological changes during development. We find that Dm4 neurons are born in the mid-third-instar larval brain as a cluster of cells in the medial and ventral region of the developing medulla cortex. Subsequently, during pupal development, Dm4 neurons move across the antero-posterior and dorso-ventral axes. We find that Dm4 neurons initiate their movement across the antero-posterior axis at ~25h APF, followed by their dorso-ventral migration 5 hours later at ~30h APF. We show that Dm4 axons project to their target regions in the medulla neuropil well before migration is initiated, whereas dendritogenesis occurs during migration and is completed by 48h APF. An RNAi-based screen for genes required for Dm4 migration identified the transcription factor Traffic Jam (Tj) as a critical regulator of the process. Loss of tj in Dm4 neurons results in a ventromedial cluster of cells in the adult medulla that failed to migrate across both the antero-posterior and dorso-ventral axes. tj-mutant Dm4 neurons project to the correct target layers in the medulla, but exhibit disorganized dendritic projections in the M3 layer. Strikingly, single-cell clones of tj-mutant Dm4s reveal regional differences in the severity of these defects; cells fated to move a larger distance from their birth location exhibit greater dendritic defects compared to those with shorter migration trajectories. Taken together, our data support a model in which neuronal migration is required for neurons to form proper dendritic connections within their target columns.

**827A  Investigating the role of VAPB in axonal ER and motorneuron development and degeneration Elizabeth Anderson Case Western Reserve University**

transcription factors toy and acj6. Our results indicate that they belong to three LCN subtypes previously identified to encode looming and the onset of edge. The high specificity of expression in this line provides an ideal tool for further developmental characterization of these neurons, as well as to understand how their highly precise patterns of axon-target matching occurs during development.
Smooth ER (endoplasmic reticulum) forms a continuous network inside of each motorneuron, dominating the intracellular landscape and extending through each axon from root to tip. Smooth ER acts as a source and sink of Ca\(^{2+}\) for organelles and tethers itself to other organelles, all lending to the hypothesis that it could be dynamically communicating and trafficking intracellular information, like a “neuron within a neuron”. Recent findings suggest that VAPB (vesicle-associated membrane protein [VAMP]-associated protein B) regulates smooth ER morphology and function in cultured cells. A highly conserved portion of VAPB can also be cleaved to act as an extracellular ligand. Intriguingly, this domain can harbor a point mutation (P58S) causative of amyotrophic lateral sclerosis (ALS). The complete molecular mechanisms linking VAPB to ER and motorneuron development and degeneration remain elusive. We will investigate these interactions using a simple Drosophila Crispr-mediated knockout model that allows us to knock back in both GFP-tagged rescue and P58S VAPB constructs. Excitingly, the VAPB knockout I engineered seems viable. In conjunction with new adult neuromuscular junction (NMJ) dissection techniques, this uniquely poises us to track synaptic development and VAPB localization at the NMJ in vivo from larvae to adult. We will observe gross ER morphology and function by crossing these mutants with existing lines that mark ER, calcium signaling, and autophagosomal trafficking. Flight behavior assays will measure motorneuron activity. We hypothesize, due to prior in vitro studies, that VAPB knockout and VAPB P58S mutant Drosophila will exhibit deficiencies in VAPB localization and ER morphology and function at the NMJ. We further hypothesize that this will correlate with deficits in motorneuron development, and ultimately degeneration and altered flight behaviors.

828B  The Role of Thrombospondin in Neuromuscular Junction Development and Function Grace Woods\(^1\), Karli Corey\(^1\), Sonya Lee\(^1\), Eve Lowenstein\(^2\), Asia Wooten\(^3\), Allie Osgood\(^4\), Luke Rotello\(^5\), Daniela Mendoza Ortiz\(^6\), Norma Velazquez Ullola\(^1\) 1) Department of Biology, Lewis & Clark College, Portland, OR; 2) PMCB Graduate Program, OHSU, Portland, OR; 3) Medical School Program, OHSU, Portland, OR; 4) National Human Genome Research Institute, NIH, Bethesda, MD; 5) Biochemistry and Molecular Biology Program, Lewis & Clark College, Portland, OR; 6) University of Turku, Turku, Finland

Thrombospondin (TSP) is an extracellular matrix glycoprotein that plays a role in synaptogenesis at glutamatergic synapses in the mammalian brain. While there are 5 TSP genes in humans, there is a single homologous gene in Drosophila melanogaster (D-TSP) and there is conservation in the protein domains involved in TSP’s function in synaptogenesis. It has not been investigated if D-TSP plays a role in synaptogenesis in Drosophila melanogaster. Here we determined if D-TSP modulates synaptogenesis and locomotor behavior in the D. melanogaster third instar larval NMJ. We hypothesized that D-TSP would be necessary for normal NMJ formation and locomotor behavior. We used the GAL4-UAS system to knock down D-TSP in neurons with two different RNAi lines and quantified features of the NMJ structure and locomotor behavior in larvae with normal or decreased D-TSP expression. Both RNAi experiments showed no significant survival differences between the parental lines and the cross. For the anatomical analysis, we took images of NMJs from muscle 4 of segments A3-4 in female third instar larvae from both RNAi crosses. We manually counted the number of boutons, number of branches, number of branch points, and number of islands of each NMJ, and generated a complexity index comprising the number of branches, number of branch points, and number of islands. Our preliminary results from experiments with one of the RNAi lines suggest an increase in NMJ complexity when D-TSP is knocked down in neurons. We also calculated the polygon area for each NMJ, and found no differences in the D-TSP knockdowns. We are currently normalizing our morphological data by muscle area, increasing the sample size of NMJs for the first RNAi line, and completing our anatomical analysis of the second RNAi line. To investigate locomotor behavior, we took videos of male and female larvae moving in a gridded arena, and analyzed the larval trajectories for distance, velocity, curvature, and zones (a measure of how far larvae move from their point of origin). We found that TSP knockdown larvae from the second RNAi line exhibited a significantly higher distance travelled compared to control larvae. We are in the process of validating the knockdowns of D-TSP by RT-qPCR. In the future, we will analyze additional behaviors, such as head-turning and body contractions.

829C  Investigating roles of conserved domains in the calcium channel subunit \(\alpha_\delta-3\) during synapse development Marina Bostelman, Heather Broihier Case Western Reserve University, Cleveland, OH

Synapses are highly specialized for neurotransmitter signaling, yet activity-dependent growth factor release also plays critical roles at synapses. While efficient neurotransmitter signaling relies on precise apposition of release sites and neurotransmitter receptors, molecular mechanisms enabling high-fidelity growth factor signaling within the synaptic micro-environment remain obscure. At the Drosophila neuromuscular junction (NMJ), the highly evolutionarily conserved auxiliary calcium channel subunit \(\alpha_\delta-3\) promotes an activity-dependent, autocrine Bone Morphogenetic Protein (BMP) signal by functioning as an extracellular scaffold, thereby regulating synapse density, structure, and function. Furthermore, \(\alpha_\delta-3\)’s promotion of BMP signaling is independent of its canonical role in the localization of the pore-forming \(\alpha_0\) calcium channel subunit, Cacophony. How these functions are separated, and what roles several evolutionarily conserved protein domains within \(\alpha_\delta-3\)’s structure play in each of these functions, is currently unknown. This question is being addressed by utilizing the CRISPR-Cas9 gene editing system to generate Drosophila lines containing deletions or modifications of highly conserved key motifs within \(\alpha_\delta-3\). Assessing NMJ morphology and function in these lines will exhibit deficiencies in VAPB localization and ER morphology and function at the NMJ. We further hypothesize that this will correlate with deficits in motorneuron development, and ultimately degeneration and altered flight behaviors.
mutants at embryonic and larval stages will provide insight into the role of these motifs in synapse formation throughout early *Drosophila* development.

**830A** **Inhibitors of BMP signaling during synapse development in *Drosophila melanogaster*** Kendall Cook¹, Cameron Rodriguez², Joseph Peters¹, Polina Yagusevich¹, Emily Januck¹, Kaitlin Tortorette², Amin Ghabrial², Pam Vanderzalm¹ 1) John Carroll University, Cleveland, OH; 2) Columbia University, New York City, NY

The canonical Bone Morphogenic Protein (BMP) pathway helps coordinate the growth and development of synapses in both vertebrates and invertebrates. The *Drosophila melanogaster* neuromuscular junction (NMJ) is a glutamatergic synapse used as a model for the AMPA-type excitatory synapses of the mammalian central nervous system, which develop in a structurally similar way. Retrograde BMP signaling from postsynaptic muscle to presynaptic neuron is critical for the scaling growth of the synaptic termini proportional to larval muscle growth. In the absence of any BMP signaling, NMJs severely undergrow and do not release neurotransmitter normally.

Though nearly twenty years have passed since the identification of the core BMP signaling components regulating synaptic growth and function, we still have an incomplete understanding of how the core pathway itself is regulated. We identified Tao, a conserved serine/threonine kinase implicated in autism, as an inhibitor of retrograde BMP signaling during the scaling growth of the larval NMJ. Loss of neuronal Tao results in overgrowth of synaptic termini without affecting muscle size, suggesting that Tao normally inhibits NMJ growth. Evoked release of neurotransmitter is also impaired in Tao loss-of-function, resulting in larval locomotion defects. Previous studies showed that Tao inhibits growth of epithelial tissues via activation of the conserved Hippo pathway, as a kinase functioning upstream of Hippo itself. In NMJ development, however, Tao functions independently of Hippo signaling, instead inhibiting the transcription of BMP target genes through a yet-unknown mechanism. We will present our recent progress made in identifying Tao's neuronal signaling partners and understanding the mechanism by which Tao and its partners inhibit BMP signaling during synapse development.

**831B** **TRMT9B regulates synaptic function and motor behavior** Ambar Delgado¹, Kimberly Rose Madhwanii, Caley Hogan³, Jennifer Dumouchel⁴, Kate O’Connor-Giles¹,⁵ 1) Department of Neuroscience, Brown University, Providence, RI; 2) Neuroscience Graduate Training Program, Brown University, Providence, RI; 3) Laboratory of Genetics, University of Wisconsin-Madison, Madison, WI; 4) Therapeutic Sciences Graduate Training Program, Brown University, Providence, RI; 5) Carney Institute for Brain Science, Brown University, Providence, RI

The nervous system is highly dependent on the dynamic regulation of translation. Posttranscriptional modification of RNAs is emerging as a key regulator of protein synthesis. tRNAs are extensively post-transcriptionally modified to regulate their structure, stability, decoding speed, and accuracy. Through a genetic screen in *Drosophila* to identify new, conserved synaptic genes, we discovered that tRNA methyltransferase family member TRMT9B, also known as fid, negatively regulates synaptic growth. Here, we investigated TRMT9B’s role in regulating neurotransmission under basal conditions and during homeostatic plasticity, and in locomotor behavior. Current clamp electrophysiology at the neuromuscular junction (NMJ) shows a significant decrease in neurotransmitter release in TRMT9B mutants. Considering the presence of ectopic synapses in TRMT9B mutants, this suggests diminished function of individual synapses at the NMJ and a role for TRMT9B in establishing precise synaptic communication.

Synapses modulate their strength during plasticity and to homeostatically maintain function at physiological levels in response to perturbation. At the Drosophila NMJ, disruption of postsynaptic glutamate receptors triggers a compensatory increase in presynaptic neurotransmitter release that precisely offsets the deficit to maintain optimal synaptic communication. Chronic presynaptic homeostatic potentiation (PHP) depends on the dynamic regulation of gene expression, so we investigated the role of TRMT9B in chronic PHP. Interestingly, we found that loss of glutamate receptor IIAI resulted in lethality in the absence of TRMT9B. While this lethality precludes electrophysiological analysis, it suggests that long-term homeostatic compensation of neurotransmission may require TRMT9B. Finally, we investigated TRMT9B’s role in locomotion through larval crawling and climbing assays, and observed significant impairment at both developmental stages. Overall, our findings indicate critical roles for TRMT9B in motor function.

**832C** **Na+/H+ exchanger (Nhe) regulates neuronal morphology at the neuromuscular junction** Ashley Bielawski², Isabella Maag³, Beverly Piggott¹,² 1) University of Montana, Missoula, MT; 2) Center for Biomolecular Structure and Dynamics, University of Montana, Missoula, MT

Emerging evidence points to a link between pH imbalances, and psychiatric conditions such as schizophrenia, autism spectrum disorder, and bipolar disorder. Due to its wide-ranging potential for new treatment approaches, it is imperative to establish whether altered pH may influence neural development and drive disease pathology. Intracellular pH can influence critical developmental pathways including cell migration, ion channel activity and abundance, and protein trafficking. However, it’s not clear which of these processes act as pH-dependent modifiers of neuronal development. Na+/H+ exchangers (Nhe) are major regulators of pH and electrolyte balance, essential for brain development and neural
function. Disruption of Nhes in early life is associated with developmental disorders including microcephaly, intellectual disabilities, autism, and febrile seizures, but the specific role of Nhes in disease manifestation is not well understood. Compared to mammalian genomes which typically encode 9 Nh proteins, Drosophila melanogaster encodes 3, making them a more straightforward model to elucidate Nhe function. Development of the Drosophila neuromuscular junction (NMJ) is an activity-dependent process, which precisely controls the number of synaptic boutons. It is known that pH-sensitive, presynaptic ion channels play a role in regulating neuronal excitability and bouton number at the NMJ. In addition, transient changes in pH at the synaptic cleft have been observed following an action potential, but how pH influences neuronal development and morphology is unclear. We find that altered Nhe2 expression affects bouton number at the NMJ. This work defines a new role for Nhe proteins in neuronal morphology and development.

833A Ion channel trafficking is coordinated with dendrite morphogenesis in sensory neurons Josephine Mitchell1, Ipek Midillioglu2, Jill Wildonger3,4, Kathy Wang1, Chun Han1 1) Biochemistry Department, University of Wisconsin-Madison, Madison, WI; 2) Pediatrics Department, University of California, San Diego, La Jolla, CA; 3) Weill Institute for Cell and Molecular Biology and Department of Molecular Biology and Genetics, Cornell University, Ithaca, NY; 4) Division of Biological Sciences, Cell & Developmental Biology Section, University of California, San Diego, La Jolla, CA

An organism’s ability to perceive stimuli, such as pain, from its surrounding environment is essential to survival. The perception of an external stimuli and its transformation into a cellular response depends on ion channels that are distributed throughout the dendritic arbor of peripheral sensory neurons. However, it is not well understood how trafficking of ion channels to their sites of function in the dendritic membrane is regulated and whether or how their localization is coordinated with dendritic morphogenesis. We investigated how ion channel localization is regulated using the Drosophila melanogaster class IV dendritic arborization neurons. In larvae, painful mechanical stimuli are sensed in part by pickpocket 1 (ppk1) and pickpocket 26 (ppk26), which are members of the degenerin/epithelial Na+ channel/acid sensing ion channel (DEG/ENaC/ASIC) family. Ppk1 and ppk26 form a heterotrimeric channel in the class IV sensory dendrites and are dependent on each other for membrane insertion. To study the regulation of pickpocket ion channel trafficking, we used CRISPR-Cas9 genome engineering to fluorescently tag endogenous ppk1. We found that ppk1 localizes robustly and uniformly throughout the dendritic membrane and is present in actively growing dendrite tips, indicating that sensory function is established during dendrite morphogenesis. To delineate mechanisms that regulate ppk1 trafficking we investigated the endocytic network. We hypothesized that Rab11 may be involved in forward trafficking of ppk channels to the membrane and/or through local regulation via recycling pathways. We found that ppk1 and ppk26 localization to the dendritic membrane depends on the recycling endosome GTPase Rab11. Additionally, we found that ppk1 and ppk26 colocalize with the early endosome GTPase Rab5. When ppk1 is absent, ppk26 is not membrane expressed and instead forms puncta throughout the dendrites. Over half of these ppk26 puncta colocalize with Rab5, suggesting involvement of early endosomes in quality control of pickpocket channel trafficking. Together, our results suggest that pickpocket channel trafficking is coordinated with sensory dendrite morphogenesis and that its membrane expression relies on the endosomal network.

834B Na+/H+ Exchangers play essential roles in neurogenesis Ashley Bielawski1, Isabella Maag1, Beverly Piggott1,2 1) University of Montana, Missoula MT; 2) Center for Biomolecular Structure and Dynamics, University of Montana, Missoula, MT

Electrolyte balance and the maintenance of physiological pH are important in all cells across the animal kingdom. Sodium (Na+) Proton (H+) Exchangers (Nh) are powerful proteins with the capacity to shape their cellular environment by acting as major modifiers of electrolyte balance and pH. They have emerged as important regulators of cell size, cellular metabolism, and neural excitability. Despite their potential to influence many key processes in brain development, their role in neurogenesis is not well defined. Human mutations in Nh proteins are found to cause microcephaly, autism, and epileptic seizures, but how they drive disease pathology is not well understood. Defining the role of Nh proteins in neurogenesis is not well defined. Human mutations in Nh proteins are found to cause microcephaly, autism, and febrile seizures, but the specific role of Nhps in disease manifestation is not well understood. Defining the role of Nh proteins in neurogenesis is not well defined. Human mutations in Nh proteins are found to cause microcephaly, autism, and febrile seizures, but the specific role of Nhps in disease manifestation is not well understood.

835C Glia-dependent regulation of synapses in the Drosophila antennal lobe Dan Jindal1, Isabella Bacon1, Elizabeth Seitz2, Sugapradha Saravanan1, Alexis Bell1, Rhea Mahajan2, Heather Broihier1 1) Case Western Reserve University School of Medicine, Cleveland, OH; 2) Hathaway Brown School, Shaker Heights, OH

Developing nervous systems initially form with an overabundance of synapses. The conserved process by which excess synapses are eliminated to yield mature nervous system circuitry is known as synaptic pruning. Beyond traditional supportive roles in neuronal homeostasis and function, glia are the primary effectors of synaptic pruning and clear

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synapses in part by phagocytosis. The mechanisms that tune glia-mediated synaptic pruning over development and adulthood are largely unclear.

Draper, an ortholog of Ced-1/MEGF10/Jedi, is the conserved surface receptor on glia that mediates engulfment and clearance in Drosophila. Glia require Draper to restrict presynapse number to normal levels in the adult antennal lobe. Directed screens are ongoing to reveal putative genes upstream of Draper that control the glial phagocytic program to allow for normal presynapse morphology. In tandem with this screen, it is important to assess (1) the effect of adult age on observed differences in presynapses between wild-type and candidate gene knockdown flies and (2) the split in pruning responsibility over time between two glial subtypes in direct contact with synaptic neuropil: astrocytes and ensheathing glia.

After this screen, it is crucial to characterize the effect of positive hits on the postsynaptic side of antennal lobe synapses. Beyond development, synaptic pruning is utilized by processes in the adult CNS like memory formation and maintenance. And so, the when and where of identified genes in regulating synapses is important; the developmental timing and the specific glia in which these genes act. Lastly, it is important to determine the functional impact of screened genes on adult olfaction and memory. Given that dysfunctional signaling between glia and synapses contribute to a host of synaptopathies, understanding the signaling pathways that direct the balance of synapse formation and removal is of significant therapeutic interest.

836A Exploring the role of glial Syndecan on neuroepithelium expansion in the Drosophila optic lobe Duo Cheng, Jaimy Coates, Vanessa Auld University of British Columbia

In order to establish a functional brain, neural stem cells (NSC) need to generate an abundant number of neurons and glia under tight temporal and spatial regulations. The Drosophila melanogaster’s optic lobe is a popular model to study neurogenesis as each optic lobe is generated by neuroblasts (NB) that differentiate from the columnar neuroepithelium (NE). Glia provide essential modulation to the NSC niche to ensure appropriate NSC differentiation. During the 3rd larval larval stage, the OPC+NE are surrounded by two distinct classes of glia, the subperineurial glia (SPG) and the cortex glia (CG). Both are known to modulate the NSC niche and play a role in NSC homeostasis. Yet the tools by which glia utilize to interpret and control the NSC niche are far from being fully characterized. Here, we explored the role of a transmembrane heparan sulfate proteoglycan (HSPG), Syndecan (Sdc) in regulating the NSC niche during development. Using super-resolution microscopy, we revealed Sdc is widely expressed across the brain lobe, though we notice Sdc is particularly enriched around the NE. Pan-glial knockdown of Sdc revealed a dramatic decrease in brain lobe volume, accompanied by a reduction of the NB population size within the OPC. Moreover, we observed the lengthening of the ventral nerve cord, indicative of disruption between glia and the overlying extracellular matrix that coats the entire nervous system. While the loss of Sdc in glia did not significantly alter the level of apoptosis, it did lead to a reduction in mitotic events in the OPC. Upon further experiments, we observed pan-glial Sdc knockdown perturbed the NE morphology, and reduced Notch activity within the NE. Notch signaling is known to prevent premature differentiation of NE into medulla neuroblasts, ensuring the expansion of the NE. Therefore we are investigating the role of Sdc in mediating Notch signaling cross-talk between the subperineurial glia and the NE. We are also exploring the potential influence of cortex glia on OPC proliferation and NE morphology. Overall, our results support a novel aspect of Sdc’s role in regulating the NSC niche through glial regulation of the NE in the developing brain lobe.

837B Glia-derived lipid binding protein confers resistance to oxidative stress in the Drosophila brain Jun Yin, Hsueh-Ling Cheng, Jingce Lei, Anna Grigsby-Brown, Mary Gibbs, Aidan Dermady, Quan Yun National Institute of Neurological Disorders and Stroke, NIH, Bethesda, MD

Lipid trafficking between neurons and glia contributes to the development, function, and stress responses of the nervous system. Due to the diversity of lipid transport proteins and lipoprotein receptors, cellular and molecular pathways involved in neuron-glial lipid shuttling have not been well characterized in vivo under physiological conditions. How neurons obtain their lipid supply during development and how synaptic activity regulates this process remain open questions. To identify the molecular network involved in this process, we performed genetic screens on glia-derived secretory factors and evaluated their distributions and functions in the Drosophila central nervous system (CNS). One of our top hits is Odorant-Binding-Protein 44a (Obp44a), a small secretory protein highly expressed in both larval and adult astrocytes. Structure homology modeling indicates OBP44a as a lipid binding protein that potentially interacts with both fatty acids and cholesterol. Using CRISPR-mediated genome editing, we generated knock-in and knock-out OBP44a alleles, validated its glial expression and revealed its specific functions in regulating the physiological properties of fly neurons. Notably, OBP44a level is upregulated in flies challenged with the H2O2 treatment, while the mutant shows a significant reduction in their resistance to oxidative stress, as well as deficits in locomotor activity and sleep maintenance. Based on these findings, we propose that Obp44a functions as an antioxidant that protects lipids from peroxidation within the fly brain and represents a new class of glia-derived molecules contributing to the unique CNS lipid environment.

838C Divergent signaling requirements of dSARM in injury-induced degeneration and developmental glial phagocytosis Yizhou Liu, Kelsey Herrmann, Arnau Llobet-Rosell, Colleen McLaughlin, Lukas Neukomm, Jaeda Ling Cheng, Jingce Lei, Anna Grigsby-Brown, Mary Gibbs, Aidan Dermady, Quan Yun National Institute of Neurological Disorders and Stroke, NIH, Bethesda, MD
Elucidating signal transduction mechanisms of innate immune pathways is essential to defining how they elicit distinct cellular responses. Toll-like receptors (TLR) signal through cytoplasmic TIR domains that bind other TIR domain-containing adaptors. dSARM/SARM1 is one such TIR domain adaptor best known for its role as the central axon degeneration trigger after injury. In degeneration, SARM1’s domains have been assigned unique functions: the ARM domain is auto-inhibitory, SAM-SAM domain interactions mediate multimerization, and the TIR domain has intrinsic NAD+ hydrolase activity that precipitates axonal demise. Whether and how these distinct functions contribute to TLR signaling is unknown. Here we show divergent signaling requirements for dSARM in injury-induced axon degeneration and TLR-mediated developmental glial phagocytosis through analysis of new knock-in domain and point mutations. We demonstrate intragenic complementation between reciprocal pairs of domain mutants during development, providing evidence for separability of dSARM functional domains in TLR signaling. Surprisingly, dSARM’s NAD+ hydrolase activity is strictly required for both degenerative and developmental signaling, demonstrating that TLR signal transduction requires dSARM’s enzymatic activity. In contrast, while SAM domain-mediated dSARM multimerization is important for axon degeneration, it is dispensable for TLR signaling. Finally, we find that dSARM functions with the MAP3K Ask1 during development but not in degenerating axons. Thus, we propose that dSARM exists in distinct signaling states in developmental and pathological contexts.

839A  Characterising the molecular basis of Drosophila glial diversity Inês Lago-Baldaia1, Austin Seroka2, Chintan Trivedi1, Maia Cooper1, Gareth Powell1, Stephen Wilson1, Sarah Ackerman2, Vilaiwan Fernandes1 1) Department of Cell and Developmental Biology, University College London, London, London, UK; 2) Institute of Neuroscience, Howard Hughes Medical Institute, University of Oregon, Eugene, OR, USA

Glial form a vital part of the nervous system and play pivotal roles in nervous system development and function. Nonetheless, they have been understudied and are poorly characterised. In Drosophila, glia are broadly classed as surface glia, cortex glia, ensheathing glia and astrocyte-like glia. Whether these categories can be further subdivided to define unique cell types is unclear. Recent advances in single cell transcriptomics have been invaluable for rapidly characterising and defining cell types in molecular terms. Here we used published single cell RNA sequencing (scRNA seq) datasets of the complex but spatially ordered Drosophila optic lobe to characterise glial diversity in molecular terms. The optic lobe contains ~14 morphologically distinct glia, making it an ideal system to study glial diversity. We first used the R Seurat package to integrate two independent glial scRNA seq datasets (Özel et al., 2020; Kurmangaliyev et al., 2020) for the adult optic lobe. This analysis generated 17 distinct clusters. We used in situ hybridisation chain reaction (HCR) and reporter lines to test in vivo expression of cluster markers and annotated 15 of the clusters. We determined that the majority of specialisation exists between the lamina neuropil and the rest of the optic lobe, with lamina-specific subtypes of cortex, astrocyte-like and ensheathing glia. While our bioinformatic analysis of non-lamina glia generated multiple clusters with either astrocyte-like, ensheathing or cortex glial molecular characteristics, these could not be explained by morphological or regional differences when we validated markers in vivo. We speculate that at least some of these clusters may correspond to distinct cell ‘states’ rather than ‘types’. On the other hand, our analysis also uncovered unexpected diversity within the chiasm glia. We have used annotated glial clusters from the adult dataset to identify glia from earlier developmental stages beginning with the third larval instar (Konstantinides et al., 2021). We have also compared glial transcriptomes of the embryonic central nervous system (unpublished) and adult optic lobe and found remarkable overlap in gene expression, suggesting a very strong maintenance of cell identity. Thus, by creating a Drosophila glial atlas, our work has begun to provide important insights into glial diversity and development.

840B  Regulation of Glial Septate Junction proteins by microRNA-184 Sravya Paluri, Vanessa Auld Life Sciences Institute, University of British Columbia, Vancouver, Canada

Glia form a vital part of the nervous system and play pivotal roles in nervous system development and function. Nonetheless, they have been understudied and are poorly characterised. In Drosophila, glia are broadly classed as surface glia, cortex glia, ensheathing glia and astrocyte-like glia. Whether these categories can be further subdivided to define unique cell types is unclear. Recent advances in single cell transcriptomics have been invaluable for rapidly characterising and defining cell types in molecular terms. Here we used published single cell RNA sequencing (scRNA seq) datasets of the complex but spatially ordered Drosophila optic lobe to characterise glial diversity in molecular terms. The optic lobe contains ~14 morphologically distinct glia, making it an ideal system to study glial diversity. We first used the R Seurat package to integrate two independent glial scRNA seq datasets (Özel et al., 2020; Kurmangaliyev et al., 2020) for the adult optic lobe. This analysis generated 17 distinct clusters. We used in situ hybridisation chain reaction (HCR) and reporter lines to test in vivo expression of cluster markers and annotated 15 of the clusters. We determined that the majority of specialisation exists between the lamina neuropil and the rest of the optic lobe, with lamina-specific subtypes of cortex, astrocyte-like and ensheathing glia. While our bioinformatic analysis of non-lamina glia generated multiple clusters with either astrocyte-like, ensheathing or cortex glial molecular characteristics, these could not be explained by morphological or regional differences when we validated markers in vivo. We speculate that at least some of these clusters may correspond to distinct cell ‘states’ rather than ‘types’. On the other hand, our analysis also uncovered unexpected diversity within the chiasm glia. We have used annotated glial clusters from the adult dataset to identify glia from earlier developmental stages beginning with the third larval instar (Konstantinides et al., 2021). We have also compared glial transcriptomes of the embryonic central nervous system (unpublished) and adult optic lobe and found remarkable overlap in gene expression, suggesting a very strong maintenance of cell identity. Thus, by creating a Drosophila glial atlas, our work has begun to provide important insights into glial diversity and development.
and decreased larval locomotion. In future, we will assess BBB integrity by dye penetration assays to understand the functional implication of SJ degradation. To determine which SJ mRNAs are affected by miR-184, we will quantify mRNA levels for each potential target using qRT-PCR. These results will reveal if miR-184 directly targets all SJ mRNAs predicted to be targets of miR-184, or just key SJ mRNAs which affect other SJ protein localization. A potential candidate for the latter possibility is NrxIV, which is a core protein that mediates SJ assembly. We will analyze the role played by NrxIV by testing if NrxIV mRNA lacking the miR-184 target sequence can rescue the SJ degradation phenotype. Our study will reveal the presence and regulation of key SJ proteins in SPG crucial for BBB maintenance. Degradation of permeability barriers in glia can result in a compromised BBB—a hallmark of many diseases across many species.

841C Investigating the localization and function of laminin and dystroglycan in Drosophila wrapping glia development Katherine Clayworth, Vanessa Auld University of British Columbia, Vancouver, BC

Peripheral nervous system (PNS) health is largely dependent on proper glial cell functioning during development. Myelinating and non-myelinating Schwann cells (MSCs and NMSCs, respectively) are glial cells in the PNS that ensheathe and protect axons. Communication between Schwann cells and the extracellular matrix (ECM) is essential for PNS development. The ECM protein laminin, and its receptor dystroglycan [Dg; part of the dystrophin-glycoprotein complex (DGC)], are important for MSC development, however very little is known about the mechanisms underlying the role of laminins, Dg, and the DGC in NMSC development. We use developing Drosophila wrapping glia (WG), which ensheathe axons similarly to NMSCs, as a model to study the role of laminin/Dg in NMSC development. We found strong expression of LanA (one of two laminin alpha subunits in Drosophila), around WG. Wing blister, the other laminin alpha subunit, is not strongly expressed in the peripheral nerve, indicating that the LanA-containing isoform is the primary laminin isoform expressed. Knockdown of LanA in WG eliminated LanA expression around WG and caused WG swellings, suggesting that LanA is expressed by WG. Preliminary data suggests that LanA is most often found at the adaxonal WG membrane (between WG and axons), rather than the abaxonal WG membrane (between WG and its adjacent glial layer, subperineurial glia). These results potentially indicate a form of WG polarization, a feature that has not been well understood in WG thus far. We found the laminin receptor Dg is also expressed on WG membranes, however there are three isoforms of Dg and their individual expression patterns in the PNS are unknown. Therefore, we are in the process of tagging alternatively spliced exons of Dg, which will allow us to follow their localization in vivo. Knockdown of Dg and dystrophin (a component of the DGC), leads to WG ensheathment failure, indicating that Dg and dystrophin are important for WG development. We are investigating other DGC components (e.g., dystrobrevin, syntrophins) to determine the composition of the DGC in WG. Due to the highly conserved nature of laminins and DGC proteins, our results have implications for NMSC development—thus improving our understanding of the factors underlying PNS development in all animals.

842A Identifying subperineurial glia-specific dlg1 isoforms required for septate junction function Mary Gilbert, Vanessa Auld University of British Columbia

Septate junctions (SJ) within the subperineurial glia (SPG) form permeability barriers in Drosophila that protect central and peripheral neurons from exposure to the hemolymph; the blood-brain and blood-nerve barriers. The ladder-like structure (or septa) of SJs forms “rails” running parallel to two adjacent cell membranes between neighboring cells or autotypically in the same cell along the length of the glia, and these create the physical permeability barrier. Core components of SJs (such as NeurexinIV, Neuroglian and the Na/K ATPase pump) are defined as proteins that create the physical junction and are essential for SJ integrity, where the absence of any one leads to a compromised permeability barrier. Another critical component of the SJ is discs large (dlg1), a polarity protein necessary for SJ scaffolding. Though how dlg1 functions to ensure SJ development and maturation is not known. Dlg1 plays wide range of cellular roles including neuromuscular junction formation, establishing polarity in epithelial cells and neuroblasts, and scaffolding various protein complexes to the cytoskeleton. The dlg1 gene organization is complex with two start sites and alternatively spliced exons capable of expressing at least 21 transcripts with a variety of functional domains including S97, 3 PDZs, SH3 and GUK domains. I have found that existing loss of function dlg1 alleles or RNAi-mediated knockdown disrupts the morphology and integrity of the glial SJs. However it is unclear which dlg1 isoforms and which domains are required in glial SJ formation. I am using a two-part strategy to investigate the which dlg1 isoforms are present in SPG and which dlg1 isoforms are necessary for SPG SJ function. I am using RNAseq approaches to identify dlg1 transcripts from isolated and purified SPG mRNA. I am using CRISPRR based approaches to generate complete deletion alleles that remove the two transcriptional start sites (S97 start site or the dlgA start site). I plan to test the requirement of each SPG dlg1 isoform by expressing dlg1 transgenes in these null backgrounds to determine which subset is able to rescue the SJ defects. In this way I will determine which domains ofdlg1 are necessary for the formation of the Drosophila blood-nerve and blood-brain barriers.

843B Exploring molecular mechanisms of Abnormal spindle function in brain growth and development Shalini Chakraborty, Todd Schoborg University of Wyoming

Autosomal recessive primary microcephaly (MCPH) is a congenital condition which is characterized by smaller brain size,
intellectual disabilities, and life span reduction. Human MCPH is most commonly caused by homozygous mutations in the abnormal spindle like microcephaly associated (aspm) gene, which has a Drosophila ortholog called abnormal spindle (asp). Recent structure-function studies from our lab have identified a 600 amino acid ‘minimal fragment’ (AspMF) of Asp’s N-terminus that is sufficient to rescue brain size in asp mutant flies. Although the MCPH phenotype is clinically well-characterized, the cellular basis of this disorder remains unknown. As an approach to uncover the cellular and molecular basis of MCPH, we employed a high-definition yeast two hybrid (Y2H) screen to identify potential interactors of AspMF. Using a third-instar larval brain cDNA library, we screened over 100 million potential domain-level interactions with AspMF. Gene ontology analysis of the candidate interactors revealed molecular functions including microtubule and actin cytoskeleton related functions in addition to signaling pathways involved in cell differentiation. Genetic rescue experiments suggest that these interactions are specifically required in neuroepithelial and lamina precursor cells of the larval brain during the neurogenic time window of development in order to promote proper brain size and morphology. We will discuss these ongoing studies to identify the molecular basis of MCPH and Asp’s ability to promote proper brain growth and development.

844C The neurodevelopmental transcriptional landscape of a fly model for human microcephaly Constanza Mannino, Mercedes Bartels-Cassidy, Todd Schoborg University of Wyoming

Autosomal recessive primary microcephaly (MCPH) is a congenital disease characterized by reduced brain size which affects 2-12 cases per 10,000 live births in U.S. There are currently 27 genes related to the MCPH phenotype. Mutations in the Abnormal Spindle-like, Microcephaly Associated (ASPM) gene account for the majority of human MCPH cases; this gene has an ortholog in Drosophila, abnormal spindle (asp), that displays the same small brain phenotype. Although mitotic spindle defects have been proposed to be the major driver of asp mutant brains, previous work from our lab and others have shown that the etiology of MCPH is complex and that multiple cellular pathways likely contribute to this disorder. This likely reflects additional roles that Asp has during neurogenesis. To uncover these roles and identify new pathways involved in MCPH, we analyzed the transcriptional landscape of asp mutant brains across three key neurodevelopmental time points: neurogenesis (3rd instar larvae), neuronal tissue remodeling (mid-pupae), and neural homeostasis (adult). Differential expression analysis revealed that the most significant expression changes (>1,000 genes, F.C. >0.5, P-value <0.05) were observed during the neurogenic and homeostasis periods, suggesting that neural remodeling does not play a major role in shaping the asp MCPH phenotype. GO analysis revealed a number of enriched terms, including an upregulation of immune system (NF-κB, Toll) and stress-related MAPK pathways that were consistent between neurogenic and homeostasis stages. To functionally test whether these pathways contribute to the etiology of asp MCPH, we probed genetic interactions between asp and components of the NF-κB (rel) & Toll (dif) pathways. Brain size was partially restored in asp/rel and asp/dif double mutant animals, despite no rescue of the neuropil disorganization phenotype. These results suggest that the inflammatory response may partially contribute to the asp small brain phenotype through a growth control mechanism that is currently under investigation. We will also present ongoing studies functionally testing the importance of MAPK-stress related pathways in the etiology of asp MCPH.

845A Innate immune signaling sculpts neuron-glia interactions across lifespan Heather Broihier Case Western Reserve University

Developmental neurobiologists have been consumed with elucidating how circuits are built and remodeled since the 1940s when Roger Sperry formulated the chemoefficiency hypothesis. Their work has led to key insights into molecular mechanisms responsible for driving axon guidance, target recognition, and synaptogenesis. What is much less clear is how these generative processes are balanced with degenerative processes. Neurons that are damaged, those with aberrant electrical activity, or lacking trophic support often undergo apoptosis during development and in the adult. And at a smaller scale, pruning of neurites and synapses is key to circuit remodeling and plasticity. Toll-like receptor (TLR) pathways were first identified for their roles in embryonic patterning in flies and have since been defined as a conserved centerpiece of innate immunity. My lab elucidated functions of TLR signaling in neural development, specifically at the NMJ. We then made the unexpected discovery that one of the most pronounced phenotypes associated with loss of Toll-6 function, a dramatic increase in the number of apoptotic neurons during development, is caused by selective loss of Toll-6 function not in neurons, but rather in glia. We demonstrated that release of the Toll-6 ligand, Spz5, from dying neurons activates a novel Toll-6 pathway in glia to prime them for phagocytosis. I will discuss recent progress my laboratory has made defining the functions of TLR signaling in regulating neuron-glia interactions in synapse, neurite, and neuron elimination across lifespan.

846B Response to and regulation of codon bias in Drosophila neural lineages. Rebecca Stewart, Scott Allen, Don Fox Duke University

Codon bias occurs when certain codons appear more frequently in the coding genome than their synonymous counterparts. This phenomenon occurs throughout all forms of life and is important in determining protein expression at the steps of both transcription and translation. Although the impact of codon bias is only beginning to be studied during animal development and in different tissues, an understanding of how individual cell types respond to codon
biased transcripts is beginning to emerge. We have undertaken one of the first studies to reveal cell and tissue-specific differences in codon bias in the model organism Drosophila and have uncovered some of the first evidence that tissues in the same animal respond differently to codon usage in vivo. Specifically, in cell types within the developing Drosophila brain, we find robust protein expression from reporters enriched in rarely used codons, whereas most other tissues in the fly do not express such reporters. Within the brain itself, neural stem cells are unable to express proteins encoded by rare-codon enriched genes, while differentiated neurons can. Cell identity appears tied to the ability to express transcripts enriched in rare codons, as experimentally blocking neuronal differentiation drastically reduces brain expression of a rare-codon enriched reporter. To uncover molecular regulators that enable neurons to specifically express rare-codon enriched reporters, we have initiated a reverse genetic screen. This screen used RNAi and the UAS-GAL4 system to knock down potential regulators enriched in neuroblasts or neurons. We assessed the role of each gene in enabling translation of rare-codon enriched transcripts by measuring expression of our rare-codon enriched reporter in neurons. Results of the screen will be presented at the poster. Using our Drosophila model, we may ultimately identify a new regulatory mechanism during neuronal development and function: cellular response to codon bias.

847C  Charting the development of leg sensory organs at the single-cell level Ben Hopkins, Olga Barmina, Tiezheng Fan, John Larue, Artyom Kopp University of California – Davis, Davis, CA

To respond to the world around them, adult flies rely on the input of a network of diverse sensory organs that are distributed across the body. The developmental genetic processes that drive divergence between different classes of sensory organs remain poorly resolved. Yet resolving these processes is critical to understanding how sensory systems function and evolve. Progress in this area has been limited by the difficulty of characterizing transcriptomic differences between sensory organs using bulk and single-cell RNA-seq (scRNA-seq): many sensory organs are rare, patchily distributed, difficult to dissociate into constituent cell-types, and embedded in undigestible cuticle. To overcome these difficulties, we developed a fine-scale dissection technique and dissociation protocol to target the pupal first tarsal segment of D. melanogaster males for scRNA-seq at multiple time points. This region is uncommonly enriched for sensory organs, containing large numbers of mechanosensory and chemosensory bristles, campaniform sensilla, the distal tibial chordotonal organ, and the sex comb, a male-specific evolutionary innovation present in a subset of Drosophilid species. Our datasets have allowed us to resolve distinct expression profiles for constituent bristle cells (sockets, shafts, sheaths, and neurons) separately for mechanosensory and chemosensory bristles and sex comb teeth. Furthermore, by subclustering neuronal populations, our data point to a combinatorial transcription factor code that specifies each of the four different chemosensory neuron classes (including male- and female-pheromone sensing neurons) and mechanosensory neurons. We verify this code using a combination of transgenic reporters and antibody staining. Finally, we recover a small population of neurons, enriched for the Pax family transcription factors eyg and toe, that mark campaniform sensilla. Using overexpression and RNAi experiments, we show that the expression of eyg and toe in campaniform sensilla suppresses bristle growth, giving rise to the characteristic dome morphology of these strain-detecting organs. Collectively, our work describes an array of developmental genetic differences that define the diverse sensory capabilities of the adult leg.

848A  Uncovering the mechanism of slit function in PNS development Maria Alejandra Pizarro Salazar, Afshan Ismat University of St. Thomas

Slit is a secreted ligand expressed in the ectoderm and plays an important role during neurogenesis. In the peripheral nervous system (PNS), removing slit results in defects in the morphology and the migration of the Ich5 chordotonal neurons. In order to gain a deeper understanding of how slit functions in this developmental process, we have over-expressed slit in the neurons, glial cells, or ectoderm of Drosophila embryos, and examined PNS neurons in each situation. Preliminary results show that over-expression of slit in these tissues results in a range of defects in comparison to wild type embryos. Over-expression of slit in glial cells results in the loss of certain dorsal neurons. The over-expression of slit in neurons shows that the cluster tails from the dorsal section are shorter than those found in wild type embryos. Additionally, over-expression of slit in the ectoderm results in defects including an axonal branch from one section that connects to another and shows a neuron that is completely out of place. The out-of-place neuron appears mostly close to the dorsal tail of sections A2-A3 with the direction of the axon going towards the ectoderm. These results suggest that Slit could be working as an attractant instead of a repellent for the neurons of the PNS. Currently, we are in the process of testing this hypothesis further. Clearly, this data demonstrates the importance of slit in PNS development, and we hope to provide insight into the mechanism of slit function in the PNS.

849B  Delta/Notch signaling inhibits expression of the early temporal factor Imp to promote termination of neurogenesis during development Chhavi Sood, Sarah Siegrist University of Virginia

In most metazoa, neural stem cells (NSCs) proliferate throughout development and then terminate their cell divisions prior to developmental completion. This ensures the formation of an adult neural circuit system that is functional and stable. To better understand the cellular and molecular mechanisms that regulate termination of NSC cell divisions and neurogenesis during development, we carried out a large scale GAL4/UAS-RNAi screen in Drosophila. From this screen,
we identified the Notch receptor and its ligand Delta. When Notch or Delta are knocked down, neuroblast divisions in the central brain region continue into adulthood, whereas when Notch is constitutively activated, neuroblast divisions terminate prematurely. We found that defects in Notch-mediated timing of termination are due to defects in neuroblast temporal patterning. When either Notch or Delta are knocked down, expression of the early temporal factor Imp (IgF-II mRNA binding protein) is maintained longer compared to controls, concomitant with delayed expression of the late acting temporal factor, Syp (Syncr). This leads to an increase in the number of Imp and Syp double positive neuroblasts, or neuroblasts with mixed temporal identity and reduced expression of the even later temporal factor, E93 (Eip93F). However, surprisingly, when we knocked down Imp and Notch signaling or overexpressed Syp and knocked down Notch signaling, the central brain neuroblasts still persisted into adulthood. This suggests that Delta/Notch cell signaling may regulate both temporal patterning and neuroblast cell cycle exit. To better understand whether Notch regulates neuroblast cell cycle exit independent of Imp/Syp temporal patterning, we assayed neuroblast quiescence during the embryonic to larval transition. When Notch is knocked down, some neuroblasts fail to enter quiescence and continue proliferating during the embryonic to larval transition. We conclude that the evolutionarily conserved Delta/Notch cell-cell signaling pathway regulates the Imp to Syp temporal transition and regulates neuroblast cell cycle exit. While Notch signaling is well known for its role in regulating binary cell fate decisions, it is becoming more apparent that Notch signaling also plays important roles in binary temporal decisions, in this case, early versus late.

850C  Deciphering the molecular clock controlling the neurogenesis diversity in drosophila’s medulla Khaled Ben el kadhi1, Claude Desplan1,2 1) New York University Abu Dhabi, Abu Dhabi, UAE; 2) New York University, NY, USA

The Drosophila compound eye is composed of 800 unit eyes; each contains 8 photoreceptors (PRs). The visual information collected by the PRs is transferred to the 4 visual processing centers of the optic lobe, the lamina, medulla, lobula and lobula plate. The medulla is the most complex structure of the optical lobe, it consists of 40,000 neurons. These neurons are the progeny of 800 medulla NeuroBlasts (NBs) that derive from a larval neuroepithelium, the Outer Proliferation Center (OPC). The OPC’s NBs divide asymmetrically to self-renew and to produce a Ganglion Mother Cell (GMC) that will produce two different medulla neurons. It was shown that the sequential expression of 6 temporal transcription factors (tTF) (Hth-Klu-Ey-Slp-D-Tll) in NBs generates neuronal diversity. Although the tTF cascade was identified, we do not have dynamic information about the timing mechanisms, the duration of each temporal identity and how the transition occurs between tTFs.

The general aim of my work is to develop a NBs primary culture to define the molecular clock of the tTF cascade using live-imaging (L-I).

We used the CRISPR-Cas9 system to endogenously tag the tTFs with different fluorescent proteins and/or transcriptional reporters MS2 or PP7. Using L-I we quantified the duration of multiple competence windows, the number of cell divisions as well as the duration of the transitions. To test if the molecular clock of the tTFs cascade is intrinsic to the NB, we set a quasi-isolated NB culture (qINBC). We found that the transition between tTFs is maintained in qINBC, suggesting that the transition signal is intrinsic to NB or comes from its progeny. We tested this hypothesis by selectively destructing the GMC or neurons (N) using Laser microablation techniques. Obtaining these dynamic data will allow us to decipher the molecular clock of tTFs and provide essential information about the mechanisms responsible for the neuronal diversity in the Drosophila optic lobe, which will likely also apply to temporal patterning observed in mammals. More importantly, this will also enable us to understand how to program a naive neural stem cell to produce a specific type of neuron that could be used for cell replacement therapy.

851A  Exploring the Role of Retrotransposable Elements in the Development of Microcephaly Bert Crawford1, Sabrina Torres1, Michelle Longworth1,2 1) Cleveland Clinic Lerner Research Institute; 2) Cleveland Clinic Lerner College of Medicine of Case Western Reserve University

Microcephaly is a rare disorder in which babies are born with head sizes that are approximately three standard deviations below the mean. Mutations in twenty-five proteins (termed microcephaly proteins), including three condensin subunits, have been shown to cause microcephaly in humans and in mouse models. However, the mechanisms by which microcephaly protein depletion and/or dysregulation leads to the development of microcephaly are not well understood. Condensins are conserved, multi-subunit complexes that are important for regulating chromosome organization. Our lab discovered that condensins also repress the expression and movement of retrotransposons in both Drosophila melanogaster and in human cells.

To determine whether increased retrotransposon expression and activity might contribute to the development of condensin-deficient microcephaly, we first used the UAS-GAL4 system to deplete condensin proteins in specific cell types of the developing Drosophila brain. Depletion of various condensin proteins in stem cells (neuroblasts), but not glial cells or post-mitotic cells resulted in significantly smaller adult brains and adult heads. We observed increased transcript levels of several families of retrotransposons and endogenous retroviruses (ERVs) in larval brains deficient for the condensin II subunit dCAP-D3. Experiments using gypsy-CLEVR reporter stocks demonstrated that replication of the gypsy retrotransposable element/ERV is prevalent in wild-type larval neuroblasts, and is significantly increased in...
dCAP-D3-deficient neuroblasts. Further, dCAP-D3-deficient larval neuroblasts exhibited increased lagging chromosomes, a phenotype observed in microcephaly patients harboring condensin mutations. dCAP-D3-deficient pupal brains also exhibit increased cell death. Excitingly, condensin-deficient microcephaly was almost completely rescued by allowing flies to develop on food containing Nucleoside Reverse Transcriptase Inhibitors (NRTIs), which inhibit retrotransposition and ERV replication. Together, these findings suggest that condensins may repress retrotransposon and ERV expression and activity in the developing Drosophila brain to prevent microcephaly. We will also present data which suggests that other microcephaly proteins may be involved in these processes.

852B  Long-range temporal patterning of progenitors in the developing Drosophila optic lobe Ishrat Maliha Islam1,2, Urfa Arain 1,2, Priscilla Valentino 1,2, Ted Erclik 1,2 1) Department of Cell and Systems Biology, University of Toronto, Toronto, ON; 2) Department of Biology, University of Toronto (Mississauga), Mississauga, ON

The Drosophila optic lobe serves as an excellent model system in which to study the mechanisms that regulate neurogenesis. The largest neuropil of the optic lobe, the medulla, is comprised of 40,000 neurons belonging to over 90 neuronal types. These neurons are generated from a neuroepithelial crescent called the outer proliferation center (OPC). At the beginning of the 3rd larval instar, a proneural wave converts each neuroepithelial cell into a neuroblast (NB), which subsequently divides asymmetrically to generate two neuronal or glial cells. It has been previously shown that OPC NBs generate unique sets of neurons based on their spatial origin and temporal state. In the spatial axis, the OPC is subdivided by the expression of Vsx1, Optix, Rx and Hh to generate eight distinct compartments. In the temporal axis, each NB sequentially expresses a cascade of five temporal transcription factors (tTFs) - Hth, Ey, Slp1/2, D and Tll - as it ages. Therefore, as the NBs in each OPC spatial domain divide, they rapidly cycle through the tTFs to generate a set of unique neurons at each spatio-temporal birth address. Here, we describe a third patterning axis that further diversifies medulla neuronal fates. We show that the OPC neuroepithelium is patterned by the expression of five long-range temporal factors; Imp, Syp, Chinmo, Mamo and E93. Over the course of the two days of OPC neurogenesis, we find that Imp and Chinmo are expressed in descending gradients in the OPC neuroepithelium, whereas Syp is reciprocally expressed in an ascending gradient. We show that these genes are required for the expression of the transcription factors Mamo and E93 in early and late temporal windows, respectively. To determine whether these long-range temporal patterning genes are required for neuronal fate specification, we focused on the neurons that are generated from the Vsx1-Hth spatio-temporal window. Surprisingly, we find that four (not one) neuronal types – Tm23, TmY15, Pm3 and TmY12 - are sequentially generated by Vsx1+Hth+ NBs during the two days of neurogenesis. We further show that the long-range temporal genes are required for this unexpected diversity; mamo promotes early neuronal fates (Tm23 and TmY15), whereas syp is required for late fates (Pm3 and TmY12). Taken together, our findings show that medulla NBs integrate three inputs – one spatial and two concurrent temporal (rapidly changing tTFs and long-range) – to generate the extensive neuronal diversity observed in the medulla.

853C  Intrinsic and Extrinsic Cues Regulate the Early-to-Late Transition of Transcription Factors in Drosophila Type II Neuroblast Gonzalo Morales, Mubarak Syed University of New Mexico

The central nervous system of D. melanogaster is composed of numerous morphologically different neurons and glia which originate from a limited number of type II neural stem cells (NSC). Such neural diversity poses the important question of how these few NSCs give rise to a variety of neurons and glia; furthermore, NSCs’ failure to diversify in the brain results in severe brain pathologies and behavioral defects. D. melanogaster is a widely used model to test neuronal differentiation mechanisms and transcription-mediated development of neural circuits. Type II (TII) NSCs divide asymmetrically beginning in embryogenesis and continue to divide into late larval stages. Upon division, TII NSCs self-renew and produce an intermediate neural progenitor (INP) which will also divide asymmetrically into multiple progenitors that eventually compose most of the adult’s central complex.

Recently, TII NSCs were found to express transcription factors in a temporal manner to produce neuronal diversity. Among these temporal factors, the ecysdine receptor (EcR) is essential because it initiates the early-to-late transcription factor transition, and its production takes place 56 hours after larvae hatch (ALH). Seven-up (svp) activity is required from 0-36 hours ALH for induction of EcR transcription, but svp is downregulated at 36 hours; therefore, the 20 hour time gap between svp disappearance and EcR appearance suggests an additional regulatory mechanism is responsible for EcR induction and NSC temporal cascade transition from the early to late stage in larvae. Previous studies have shown these molecular transitions are linked to intrinsic and extrinsic cues which play important roles in the larval NSC transcription factor cascade. Cell division is an internal process essential for neuroblast development, and may trigger an intrinsic process via the cell cycle to establish neural diversity and activate transitions in the temporal cascade. We hypothesize that if the cell cycle is blocked, then transcription factors inducing later-stage development of neuroblasts will be absent while early-stage transcription factors will continue to be expressed into L2/L3 stages. To test this, two kinds of cell cycle disruptors are used: first, a cytokinesis cycle delay signal (CDK-1RNAi) which inhibits the formation of a protein kinase subunit controlling important aspects of cell cycle progression; and second, pavarotti downregulation (PavRNAi) which inhibits transcription of microtubule motor proteins in spindle apparatus formation. Cell cycle disruption at both 0 and 48 hours ALH, which are critical time points for proper EcR induction, showed lack of EcR in the NSC at 56 hours. These
results support our hypothesis in that cell cycle disruption leads to persistent early-stage transcription factor expression into late larval developmental.

854A Unraveling the mechanisms of early neurogenesis with single cell resolution Ana Costa Veloso1,2, Robert Zinzen1,2 1) Max Delbrueck Centre for Molecular Medicine (MDC), Berlin, Germany; 2) Berlin Institute for Medical Systems Biology (BIMSB), Berlin, Germany

The Drosophila embryonic nervous system develops from neuroblast stem cells that emerge in a characteristic spatiotemporal pattern per segment. The lineage relationships from neuroblast to neurons and glia are highly stereotypic and have largely been described, but the underlying mechanisms driving diversification, specification and differentiation are still poorly understood.

To uncover how neuroblasts produce distinct lineage trajectories, we have produced a gene expression atlas resolving over 65 000 neurogenic stem cells across 6 consecutive timepoints encompassing all waves of neuroblast delamination. The resulting transcriptomic blueprint of neuroblasts and ganglion mother cells allows unique and novel insights into the molecular mechanisms responsible for early nervous system development.

We have been able to identify specific neurogenic populations and their gene expression dynamics along early neurogenesis. Our resource allows assigning spatial and temporal identities to individual cells and discovery of new spatio-temporal markers, including transcription factors and signaling molecules. Moreover, we show that the gene expression programs of individual neuroblast identities can be linked to specific upstream regulators and give new insights into the downstream effectors of cellular behavior. Several neuroblasts with glial potential were particularly interesting, as their glial character becomes manifest not only much earlier than expected, but temporal ordering indicates ‘developmental pockets’ of signaling receptivity. Finally, we demonstrate that newly identified drivers of neurogenesis are functionally important for crucial features of neurogenesis, such as proper motoneuron projection.

We present a resource that will aid in identifying neurogenic populations and individual neuroblasts, further our understanding of how individual neuroblasts diverge molecularly, allow discerning how their expression programs are regulated and how these differences drive the emergence of specific cell identities such as distinct motoneurons, interneurons, and glia.

855B Building an integrative model of how nutrition and natural genetic variation interact during neurogenesis in natural populations of Drosophila melanogaster Taylor L. Nystrom, Sarah Siegrist, Alan Bergland University of Virginia

During early life neurogenesis, neural stem cells give rise to thousands of morphologically and functionally diverse neurons that will build the adult brain. Ultimately, these neurons form functional circuits, and it is these circuits that allow proper functioning and survival of the adult animal. While neurogenesis is known to be influenced by both intrinsic and extrinsic signalling, we still have a limited understanding of how these signals interact. My work seeks to elucidate the integration of intrinsic and extrinsic signalling by studying how an extrinsic factor, nutrient signalling, and an intrinsic factor, natural genetic variation, interact during nutrient-dependent processes of neurogenesis in Drosophila melanogaster (fly). In order to capture genetic variation reflective of natural populations, I have used a collection of geographically diverse, wild-caught, inbred lines of flies for all experiments. I first characterized neural stem cell, or neuroblast (NB), reactivation, a temporally regulated, nutrient-dependent process of neurogenesis. My work revealed significant differences in the timing of NB reactivation from quiescence when genetically distinct flies are raised in a calorically rich environment. These differences in reactivation could affect the total amount of time that a NB is producing neurons, which could result in changes in the total number of neurons in the adult brain and potentially cause functional differences for the adult fly. Moreover, these results may suggest a genetically encoded basis for variation in neurogenesis.

After observing variation in neurogenesis in a calorically rich environment, I characterized NB reactivation in a calorically poor environment. Preliminary results show that flies with distinct genetic backgrounds respond differently to caloric restriction. Such results may suggest a gene-by-environment interaction during the transition from quiescence to reactivation in flies.

Next, I will continue to build a full timeline from reactivation to termination in larval NBs in calorie rich and calorie poor environments, measure temporally regulated transitions between transcription factors that play a role in neuronal specification, analyze the genomes of the experimental fly lines for genes involved in nutrient dependent steps of neurogenesis, and quantify the neuronal make-up of the adult fly brains. Through this work, I will build a more integrative understanding of how intrinsic and extrinsic cues interact during neurogenesis.

856C Establishing anterior-posterior diversity in how stem cells give rise to neural circuits for somatosensory processing Deeptha Vasudevan, Yi-Wen Wang, Hannah Carr, Sean Corcoran, Chris Wreden, Ellie Heckscher Department of Molecular Genetics and Cell Biology, The University of Chicago

In different anterior-posterior (A-P) locations of the body, there are different circuits that control specialized functions.
For example, humans have appendages (arms that are anterior, legs posterior to the trunk) that have unique movement patterns. Circuits at different A-P locations must process different sensory stimuli to generate specific movements. Spinal cord injury at different A-P levels gives different phenotypes. Injury in more anterior regions leads to paralysis of all 4 limbs whereas more posterior injuries result in paralysis of lower limbs, and regenerative treatments would need to generate region-specific circuits. The vertebrate spinal cord is segmented and develops from stem cells. In each segment, there are ~10 different progenitor domains that give rise to specific sets of spinal neurons. These 10 progenitor domains are repeated in every segment along the A-P axis. Stem cells have enormous therapeutic potential to regenerate circuits, but we must understand how stem cells give rise to the A-P diversity of circuits during development. What are circuit variations in the spinal/nerve cord along the A-P axis? And how are these circuit variations established in development?

I am developing the Drosophila larval nerve cord as a model to address this question. Similar to the progenitor domains in the vertebrate spinal cord, the Drosophila nerve cord has a repeating set of ~30 stem cells called neuroblasts, which differentiate into neurons that wire up to form circuits. Here, I am characterizing the A-P circuit diversity of one neural stem cell lineage called neuroblast 3-3 (NB3-3). These NB3-3 stem cells are found in every segment of the Drosophila nerve cord and give rise to sensory processing even-skipped lateral (EL) interneurons. We are testing which properties of EL circuits are position-dependent vs position-independent. First, we assayed how ELs vary across the A-P axis of the Drosophila nerve cord in number, birth timing, and morphology using molecular and temporal identity markers, and stochastic labeling techniques. We are testing diversity in sensory encoding of ELs by determining anatomical and functional input synaptic partnerships using transsynaptic circuit tracing and calcium imaging. This will tell us the diversity of somatosensory processing circuits in the nerve cord, as well as the diversity in how stem cells give rise to these circuits along the A-P axis. In the future, we can test genetic and cellular mechanisms that establish this A-P variation in circuit development.

857A The OTUD6 deubiquitinase associates with the 40S ribosome to regulate translation and the response to stressors in Drosophila

Ribosome ubiquitylation is a highly regulated process that is essential for maintaining proper protein translation. While much is known about the role of ubiquitin in these pathways, less is known about the function of deubiquitinases. OTUD6 is an ovarian tumor (OTU) family deubiquitinase that is conserved from yeast (OTU2) to humans (OTUD6A & B). To study OTUD6 in Drosophila, CRISPR/Cas9 was used to create epitope-tagged wild-type and catalytically dead (DUB-dead) endogenous OTUD6. DUB-dead and loss-of-function OTUD6 mutants are markedly sensitive to oxidizing (paraquat) and alkylating (mms) agents that impact protein translation and ribosome function. Co-immunoprecipitation coupled with mass spectrometry in OTUD6.DUB-dead brains, where OTUD6 is highly expressed, identified 40S ribosome and RNA exosome proteins. The most significantly enriched 40S proteins, RACK1 and RP53, play key roles in ribosome quality control and turnover under cellular stress. Mutation of RACK1 rescued the mms sensitivity of OTUD6 mutants, indicating that OTUD6 is likely a negative regulator of RACK1. RACK1 regulates ubiquitylation of several 40S ribosome subunits: a genetic screen of E3 ligases that ubiquitylate the 40S ribosome in a RACK1 dependent manner revealed that OTUD6.DUB-dead interacts epistatically with two E3 ligases. Both E3 ligases are implicated in Rp53 ubiquitylation and in ribosome quality control. In support of OTUD6 regulating ribosome quality control or turnover, OTUD6 mutants have decreased protein translation, delayed development, and dramatically extended lifespan. Thus, we hypothesize that OTUD6 regulates protein translation through a novel deubiquitylation event on the 40S ribosomal subunit in response to stress conditions.

858B Rasputin – A mediator of translational activation for essential proteins in neurodevelopment

Mutations in the fragile X mental retardation 1 (FMR1) gene result in fragile X syndrome (FXS) and fragile X primary ovarian insufficiency (FXPOI) - leading causes of intellectual disability (ID), autism spectrum disorder (ASD) and premature ovarian failure. FMR1 encodes the RNA-binding protein FMRP, which binds to dozens of mRNAs associated with various ASDs. Both neurons and oocytes translationally regulate stored mRNAs that are associated with FMRP. Mature oocytes are large, transcriptionally silent cells which depend entirely on the translation of stored mRNAs, making them a useful model to study FMRP-dependent gene regulation. Our lab previously demonstrated that Drosophila Fmr1 functions in oocytes primarily as a translational activator, and promotes the production of many large proteins essential for neurodevelopment.

We developed a single molecule fluorescent in situ hybridization (smFISH)-based assay to identify novel regulators of Fmr1 during Drosophila oogenesis. The mRNA of the Fmr1 target, Poe, is localized to RNP particles in an Fmr1 and translation-dependent manner. The loss of the RNA binding protein Rasputin (Rin), led to the dispersal of these ‘Poe-particles,’ which we hypothesize is due to reduced Poe translation. We performed co-immunoprecipitation of endogenously tagged Fmr1 from ovarian extracts followed by mass spectrometry. Our results showed an enrichment of Rin in Fmr1-tagged extracts relative to controls. We also identified several other RNA binding proteins as novel Fmr1 interactors, including CG5726, an ortholog of the mammalian non-canonical translation initiation factor CTIF. Previous...
studies have led to a proposed role for Rin in the stabilization and translational upregulation of its target mRNAs. We hypothesize that Rin interacts with Fmr1 and CG5726 to facilitate translational activation of Fmr1 targets in oocytes and neurons.

**859C  Steroid hormone signaling activates a sensory switch during Drosophila peripheral nervous system development Jacob Jaszczak,1,2 Laura DeVault1,3, Lily Jan1,2, Yuh-Nung Jan1,2 1) Department of Physiology, Department of Biochemistry and Biophysics, University of California at San Francisco, San Francisco, California, USA; 2) Howard Hughes Medical Institute; 3) Department of Developmental Biology, Washington University Medical School, Saint Louis, USA

Sensory neurons which detect multiple types of stimuli enable animals to sense environmental changes and avoid harm. How the neuronal properties which facilitate sensation are activated during development and the mechanisms which allow neurons to distinguish sensory modalities are poorly understood. The Class 4 dendritic arborization (C4da) neurons are multimodal sensors which tile the body wall of Drosophila larvae and detect temperature, light, and mechanical force. The development of the sensory responses mediated by C4da neurons demonstrates that the sensory function and modality are developmentally tuned processes. While larvae respond to mechanical force throughout development (Almeida-Carvalho et al. 2017), a robust activation of the thermal nociceptive escape behavior is only present during the second half of larval development (3rd instar) (Sulkowski et al. 2011). In contrast to the increase in nociceptive behavior in the 3rd instar, we find that ultraviolet light-induced Ca2+ activity in C4da neurons decreases during the same period of larval development. This demonstrates that different modalities detected by the same sensory neuron can have different developmental patterns of activity. The steroid hormone receptor Ecdysone Receptor A (EcR.A) is required for thermal nociception in 3rd instar C4da neurons (McParland et al. 2015); thus, we examined the role of ecdysone in the sensory switch. We find that EcR nuclear localization increases early in the 3rd instar, and demonstrate that both increasing ecdysone titer and overexpression of EcR.A is sufficient to promote precocious thermal nociceptive responses in 2nd instar larvae. Additionally, we find that both 2nd instar and 3rd instar nociception requires EcR.A ligand dependent activation to promote thermal nociception. We next measured the expression of known nociceptive channels. While most of the channels have changing expression from 2nd to 3rd instar, we find EcR.A suppresses expression of subdued (encoding a TMEM16 channel) with indiscernible effect on the other nociceptive genes. RNAi reduction of subdued expression in 2nd instar C4da neurons not only increases thermal nociception, phenocopying EcR.A overexpression, but also decreases the response to ultraviolet light. Thus, steroid hormone signaling suppression of subdued expression facilitates the sensory switch of C4da neurons, and suggests that ion channel balance is a key target for tuning the development of sensory modalities.

**860A  Genetic regulation and protein interactions necessary for proper formation of Drosophila rhabdomeres and the inter-rhabdomeral space Jonathan Rylee,1 Simpla Mahato1, Lalitha Sastry1, Lauren Feder1, Shaun Grega1, Brandon Sizemore1, Andrew Zelhof1,2 1) Department of Biology, Indiana University, Bloomington, IN; 2) Drosophila Genomics Resource Center, Indiana University, Bloomington, IN

Drosophila Eyes Shut (EYS) is responsible for creating and filling the inter-rhabdomeral space (IRS), a unique extracellular matrix that spatially separates the apical rhabdomeres of photoreceptors in each ommatidium, a unique adaptation in higher order Diptera. This organization represents an important step in terminal differentiation of photoreceptors as well as establishing a functional visual system in Drosophila. A major goal of our lab is to identify and characterize the genes and protein products necessary for this adaptation. We have taken two approaches towards this goal. First, we carried out an RNAi screen to identify transcription factors necessary for proper rhabdomere and IRS formation. Second, we have identified interactors with EYS through the use of proximity labeling and mass spectrometry. We will present results of both screens and how these newly identified genes contribute to the differentiation and organization of photoreceptors in insects.

**861B  Molecular instructions for the production of sparse inputs Vanessa Puñal1, Najia Elkahlah1, Jackson Rogow2, Jamal Jenkins3, E. Josie Clowney4, Mark Lewis1, Mona Saeed1, E. Josie Clowney1 1) University of Michigan; 2) The Rockefeller University

Detection of environmental stimuli is essential for survival. The total number of sensory stimuli encountered by an organism is unpredictable and nearly limitless, making genetic coding of detectors for all possible stimuli impossible. How then does development build neural circuits capable of processing unforeseen and expansive arrays of sensory input? One strategy is to harness the power of combinations. Consider a palette of 100 colors: when combined, this limited number of colors can paint a rich picture containing ~10^{198} hues. Similarly, evolution has selected for amplifying the number of discriminable stimuli from a limited number of genetically encoded sensors to the number of combinations among them. This is achieved by expanding sensory neuron inputs via sparse, combinatorial wiring to higher order neurons involved in perceptual processing. It is not understood in any species what developmental mechanisms give rise to the input sparseness necessary for sensory amplification. To address this knowledge gap, I use the fly olfactory system where 50 odor channels are dispersed, via Projection Neurons, among ~2,500 higher order neurons called Kenyon Cells in the mushroom body calyx. Previous work in the lab demonstrated that in the absence of Kenyon Cells during
development, adult Projection Neuron axons no longer provide input to the calyx. We have subsequently found that this is because developing Projection Neurons never initiate collateral formation when Kenyon Cells are not present. These results have led to the hypothesis that Kenyon Cells dictate their input density via retrograde feedback to Projection Neurons. To test this hypothesis, we generated a bulk RNA sequencing dataset for developing Kenyon Cells and used this as the basis for a candidate screen to search for Kenyon Cell-derived retrograde signals required for the production of odor channel inputs. Results suggest a role for a variety of molecules secreted by Kenyon Cells as part of a molecular pathway that drives Projection Neurons to meet Kenyon Cell demands to generate odor channel inputs.

**862C**  Analysis of sexually dimorphic gene expression in *Drosophila* legs Jude Icoy, Pratyajit Mohapatra, Karen Menuz University of Connecticut, Storrs, CT

Insects have gustatory (taste) sensilla on their wings, labella, and tarsi. These sensilla house gustatory receptor neurons which express gustatory receptors. The *Drosophila* genome contains two known families of gustatory receptors, the Grs and the IRs. Such receptors can be involved in tasting food or conspecific pheromones. Although the expression of these receptors is best-characterized in the labellum, their expression is not as well defined in the legs. There are different combinations of sensilla in the hindlegs, midlegs, and forelegs due to their different roles in behavior. In particular, the forelegs are used by males during courtship. Thus, gustatory receptor expression is expected to be sexually dimorphic and to vary between the different leg-types. Additionally, other genes may also be differentially expressed by leg-type and gender. We have utilized RNASeq to profile gene expression in male and female forelegs, midlegs, and hindlegs. Differentially expressed genes are identified and categorized by gender and by leg-type. This work can inform future studies on the role of sexually dimorphic genes in *Drosophila* gustatory behavior.

**863A**  Elucidating the interaction between the chromatin reader Kismet and histone deacetylases in the promotion of axon pruning Emily Sterner1, Daniel Marendà2, Jennifer Stanford1 1) Drexel University, Philadelphia, PA; 2) Division of Biological Infrastructure, National Science Foundation, Alexandria, VA

In *Drosophila*, the chromatin reader Kismet (Kis) is required for proper axon pruning in the developing mushroom body during pupation. Kis is a homolog of the human transcription factor and chromatin reader CHD7. Haploinsufficiency of CHD7 accounts for two-thirds of CHARGE syndrome cases in humans. This neurodevelopmental disorder causes intellectual disability and defects in facial structure and sensory organs. The axon pruning defect caused by loss of Kis in *Drosophila* can be rescued pharmacologically by HDAC inhibition. Previously, our lab has shown the general HDAC inhibitor SAHA is able to rescue axon pruning defects. To narrow down which HDAC(s) are relevant to the axon pruning process, we are taking a two-pronged approach of: 1) pharmacological inhibition using more specific HDAC inhibitors, and 2) RNAi knockdown of specific HDACs in a *Drosophila* model for CHARGE syndrome. The Class I HDAC inhibitors Beta-hydroxybutyric acid and sodium butyrate are both able to rescue defects caused by loss of Kis, suggesting that Class I HDACs are relevant to axon pruning and could be putative targets for potential therapeutic for CHARGE syndrome. Follow-up work to confirm the involvement of HDAC1 and/or HDAC3 in the axon pruning process, using RNAi, is ongoing.

**864B**  Bisphenol A exposure impacts neurodevelopmental gene expression, cognitive function, and synaptic morphology in *Drosophila melanogaster* Judith Anderson1, Chloe Welch1, Rana Ghobashy2, Salma Eliessy3, Eden Johnson4, Angelina Tupikova5, Johnathan Newman1, Adam Alfareh1, Erin Widman1, Alexandra Davis1, Wendy Lee2, Kimberly Mulligan1 1) California State University, Sacramento; 2) California State University, San Jose

Bisphenol A (BPA) is an environmentally prevalent endocrine disrupting chemical that may be a risk factor for neurodevelopmental disorders. BPA has been associated with behavioral impairment in children and causes a variety of neurodevelopmental phenotypes in model organisms. We used *Drosophila melanogaster* as model to explore the consequences of developmental BPA exposure on gene expression, cognitive function, and synapse development. Following RNA-sequencing, we performed Gene Set Enrichment Analysis (GSEA) and found neurodevelopmentally relevant genes were significantly impacted by BPA (1mM). Among the top misregulated genes were those associated with learning and synapse development. To measure learning, we used a behavioral paradigm called conditioned courtship suppression. In this paradigm, male flies are exposed to an unreceptive female (an “aversive stimulus”) for an hour. Flies with unimpaired learning exhibit a significant decrease in their courtship activity in the final ten minutes compared to the initial ten minutes of the assessment period. We found that BPA-treated flies did not reduce their courtship activity in the final ten-minute interval, indicating that BPA impaired learning. We also used immunofluorescence to examine synapse morphology within the larval neuromuscular junction and found BPA significantly increased the number of axonal branches. Our findings align with studies of BPA in mammalian model organisms, suggesting that BPA impairs functionally conserved neurodevelopmental pathways. Further, because *Drosophila* do not possess classic estrogen receptors, this research indicates that BPA can impact neurodevelopment by molecular mechanisms distinct from its role as an estrogen mimic.

**865C**  Enhancing Mask activity in dopaminergic Neurons extends lifespan in flies Xiaolin Tian LSU Health Science Center
Dopaminergic neurons (DANs) are essential modulators for brain functions involving memory formation, reward processing, and decision-making. Here I demonstrate a novel and important function of the DANs in regulating aging and longevity. Overexpressing the putative scaffolding protein Mask in two small groups of DANs in flies can significantly extend the lifespan in flies and sustain adult locomotor and fecundity at old ages. This Mask-induced beneficial effect requires dopaminergic transmission but cannot be recapitulated by elevating dopamine production alone in the DANs. Independent activation of Gq in the same two groups of DANs via the drug-inducible DREADD system also extends fly lifespan, further indicating the connection of specific DANs to aging control. The Mask-induced lifespan extension appears to depend on the function of Mask to regulate microtubule (MT) stability. A structure–function analysis demonstrated that the ankyrin repeats domain in the Mask protein is both necessary for regulating MT stability (when expressed in muscles and motor neurons) and sufficient to prolong longevity (when expressed in the two groups of DANs). Furthermore, DAN-specific overexpression of Unc-104 or knockdown of p150
d demonstrates a novel DANs-dependent mechanism that, upon the tuning of their MT dynamics, modulates systemic aging and longevity in flies.

866V  Glia-neuron signaling induced by distinct sources of two different BMPs regulate synaptic growth Mathieu BARTOLETTI, Tracy KNIGHT, Aaron HELD, Laura RAND, Kristi WHARTON Brown University

The nervous system is a complex network of cells whose interactions provide circuitry that enables an organism to perceive and move through its environment. Both neuron-neuron and neuron-non-neuronal cell communications are essential for proper nervous system function. Considerable research has focused on communication between motor neurons and muscles, albeit the molecular basis of glial cell-neuron communication is still not well understood and its elucidation will advance our understanding of complex neural responses and abnormalities of the nervous system that arise when such communication fails. In Drosophila melanogaster, retrograde BMP signaling, activated by muscle-expressed Gbb (BMP5/6/7), has been shown to influence synaptic growth, function and plasticity. Here, we report Gbb also acts from glial cells to initiate signaling in motor neurons, an additional source of BMP ligand required for synaptic growth at the neuromuscular junction (NMJ). Furthermore, we find Dpp (BMP2/4), expressed in a subset of motor neurons (RP2 and aCC), facilitates NMJ growth through autocrine neuronal signaling and its expression in RP2 versus aCC motor neurons appears tightly controlled via an autoregulatory loop. Dpp also initiates neuron to glial cell signaling, but not neuron-neuron signaling to neighboring motor neurons. Together, our findings demonstrate that bi-directional communication between glia and neurons requires two different BMP ligands. Gbb, broadly expressed in glia, as well as muscles, and Dpp, from a discrete set of neurons, activate Smad-dependent BMP signaling in motor neurons to control synapse size.

867V  Early lineage segregation of the retinal basal glia in the Drosophila eye disc Chia-Kang Tsao1,2, Yu Fen Huang1,2, Y. Henry Sun1,2 1) Institute of Molecular Biology, Academia Sinica, Taipei, Taiwan, ROC; 2) Department of Life Sciences and Institute of Genome Sciences, National Yang-Ming University, Taipei, Taiwan, ROC

The retinal basal glia (RBG) is a group of glia that migrates from the optic stalk into the third instar larval eye disc while the photoreceptor cells (PR) are differentiating. The RBGs are grouped into three major classes based on molecular and morphological characteristics: surface glia (SG), wrapping glia (WG) and carpet glia (CG). The SGs migrate and divide. The WGs are postmitotic and wraps PR axons. The CGs have giant nucleus and extensive membrane extension that each covers half of the eye disc. In this study, we used lineage tracing methods to determine the lineage relationships among these glia subtypes and the temporal profile of the lineage decisions for RBG development. We found that the CG lineage segregated from the other RBG very early in the embryonic stage. It has been proposed that the SGs migrate under the CG membrane, which prevents SGs from contacting with the PR axons lying above the CG membrane. Upon passing the front of the CG membrane, which is slightly behind the morphogenetic furrow that marks the front of PR differentiation, the migrating SG contact the nascent PR axon, which in turn release FGF to induce SGs’ differentiation into WG. Interestingly, we found that SGs are equally distributed apical and basal to the CG membrane, so that the apical SGs are not prevented from contacting PR axons by CG membrane. Clonal analysis reveals that the apical and basal RBG are derived from distinct lineages determined before they enter the eye disc. Moreover, the basal SG lack the competence to respond to FGFR signaling, preventing its differentiation into WG. Our findings suggest that this novel glia-to-glia differentiation is both dependent on early lineage decision and on a yet unidentified regulatory mechanism, which can provide spatiotemporal coordination of WG differentiation with the progressive differentiation of photoreceptor neurons.

868V  Organizing the Drosophila olfactory circuits by interacting Ig superfamily adhesion molecules Qichen Duan, Scott Barish, Allison Carson, Rachel Estrella, Pelin Volkan Department of Biology, Duke University, Durham, NC

The Drosophila olfactory system provides an excellent model to study how complex neuronal circuits are assembled. In Drosophila, each olfactory receptor neuron (ORN) class exclusively expresses a unique olfactory receptor (OR) gene and target each class-specific and uniquely positioned glomerulus in the antennal lobe. How ORN axon terminals are organized into these dedicated structural compartments is not well understood. Through transcriptome profiling of
the antennal tissues during development and RNAi screen, we identified two protein subfamilies belonging to the Immunoglobulin Superfamily, Beats and their heterophilic binding partners Sides, as novel regulators of ORN glomerular organization. Many Beats and Sides are expressed at low levels at early pupal stages but increase their expression levels later. Beats and Sides are also differentially expressed across ORN classes, making them good candidates for encoding the ORN class-specific cell surface codes to mediate neuron-neuron recognition. Perturbing the functions of many Beats and Sides in all or a subset of ORNs at later stages leads to diverse local defects of ORN glomerular organization, associated with the expanding, split, or flipped glomerular morphology. Binding Beat-Side pairs are co-expressed in the same class of ORNs and knockdown of either member of these interacting pairs could lead to similar ORN axonal disorganization. These defects are not likely to be resulted from synaptic mismatching, but rather the loss of axonal adhesion. Furthermore, OR function regulates the expression of Beats and Sides, as some of these genes show altered expression levels in OR mutants. Our data suggest the context-dependent control of ORN glomerular organization by Beat/Side combinatorial codes, and bring new insights into how diverse neuronal populations are coordinated into hardwired circuits.

**869V** Chordotonal neurons have dendritic spike initiation zones that are controlled by Para, the *Drosophila* sodium channel *Thomas A. Ravenscroft*1,2, Ashleigh Jacobs3, Mingxue Gu1,2, Daniel F. Eberl4, Hugo J. Bellen1,2,4 1) Department of Molecular and Human Genetics, Baylor College of Medicine, Houston, TX; 2) Jan and Dan Duncan Neurological Research Institute, Texas Children’s Hospital, Houston, TX; 3) Department of Biology, University of Iowa, Iowa City, IA; 4) Department of Neuroscience, Baylor College of Medicine, Houston, TX

The fruit fly *Drosophila melanogaster* has provided important insights into how sensory information is transduced by Transient Receptor Potential (TRP) channels in the Peripheral Nervous System (PNS). In bipolar chordotonal neurons (CN), TRP channels alone are not sufficient for the transduction of mechanical stimuli. We use an intronic *Minos*-mediated integration cassette to show that the sole voltage-gated sodium channel (Na+) in *Drosophila*, Para, is localized to the dendrites of these bipolar neurons. Para is localized to the distal tip of the dendrites in all CN and is colocalized with the mechanosensitive TRP channel NompC, distal to the mechanosensitive TRP channel Inactive. This localization is unique to bipolar CNs as we do not observe dendritic localization of Para in other peripheral neurons. Reducing para expression using RNAi in chordotonal neurons of the adult Johnston’s organ severely affects sound-evoked potentials. However, in addition to the dendritic localization, Para also localizes to a proximal spike initiation zone (SIZ) at an axonal initial segment-like region in axons of both bipolar and multidendritic PNS neurons. Therefore, to determine the dendritic specific role of *para* in CN neurons, we expressed a genetically encoded calcium indicator in CNs and recorded the fluorescence spike changes in dendrites upon mechanical stimuli. We show that the observed spikes in dendrites are nearly completely abolished by exposure to the Na+ inhibitor Tetradotoxin (TTX). We conclude that Na+ channels are essential for mechanosensation in CN dendrites and that Para localization marks a dendritic SIZ in the distal dendrite as well as a SIZ in axons near the cell body. Hence, the TRP channels present in the dendrite of chordotonal neurons must cooperate with Para to trigger a neuronal response.

**870V** The post-transcriptional regulation of TFs in immature motoneurons shapes the axon-muscle connectome *WenYue Guan*1, Stéphanie Bellemin1, Mathilde Bouchet1, Lalanti Venkatasubramanian2, Camille Guillermin1, Anne Laurençon1, Kabir Chérief1, Aurélien Darnas1, Christophe Godin1, Séverine Urdy1, Richard S. Mann1, Jonathan Enriquez2 1) Institut de génomique fonctionnelle de Lyon, ENS de Lyon; 2) Mortimer B. Zuckerman Mind Brain Behavior Institute, Columbia University; 3) Laboratoire Reproduction et Développement des Plantes, Univ Lyon, ENS de Lyon

Temporal factors expressed sequentially in neural stem cells, such as RNA binding proteins (RBPs) or transcription factors (TFs), are key elements in the generation of neuronal diversity. The molecular mechanism underlying how the temporal identity of stem cells is decoded into their progeny to generate neuronal diversity is largely unknown. Here, we used genetic manipulations and a new computational tool to study the unique fates of the progeny of a stem cell producing 29 morphologically distinct leg motoneurons (MN) in *Drosophila*. We identified 19 combinatorial codes of transcription factors (TFs) expressed in immature MNs (iMNs) just before their morphological differentiation. Importantly these TF codes are progressively established as a function of their birth date. The comparison of the RNA and protein expression patterns of 6 TFs in the 29 iMNs in different combinations revealed that post-transcriptional regulation plays an essential role in shaping these TF codes. We found that the two known RBPs, Imp and Syp, temporally expressed in the NB and maintained in iMNs according to their birth date, are key players in the construction of the axon-muscle connectome. Both RBPs post-transcriptionally regulate 5 of the 6 TFs examined, which in turn control the establishment of the axon muscle-connectome. By deciphering the function of Imp in the iMNs with respect to the stem cell, we propose a model whereby these two crucial RBPs act in iMNs to specify the unique morphological fate of individual MNs. Taken together, our study reveals that iMNs have the potential to acquire different morphological fates because of a broad expression of different TF mRNAs. However, these potentials are post-transcriptionally narrowed down to single iMN or subpopulations of iMNs by Imp and Syp, and potentially by other RBPs that remain to be discovered, to ultimately fine-tune the morphological fates of individual MNs.
Synaptic refinement is a neuroplasticity process leading to the withdrawal of ectopic synapses formed during the initial phases of neuronal development. Extensive research has shown evidence of synaptic refinement occurring in both the central nervous system (CNS) and peripheral nervous system (PNS). However, the molecular mechanisms underlying synaptic refinement remain incompletely understood. The process of synapse elimination is crucial during development in multiple organisms as it has also been linked to the onset of neurodevelopmental disorders like autism. Evidence from postmortem studies of autistic patients that have mutations in genes associated with autism, show increased synaptic density and longer dendritic spines, suggesting a possible defect in the elimination process. The goal of this study was to determine the anatomical effects of candidate autism genes in vivo using the Drosophila model. Starting from the hypothesis that candidate autism genes would lead to the presence of ectopic synapses that branch off stereotypic connectivity patterns, anatomical synaptic innervations of cIV nociceptive sensory neurons were assessed during larval development. The candidate autism genes nrx-1 or nlg-3, respectively coding for the scaffolding proteins neurexin-1 and neuroligin-3, were downregulated using RNAi constructs. Anatomical defects were assessed by counting the number of ectopic synapses and by measuring the fluorescent area covered by synapses. Preliminary data on the anatomical effects of the downregulation of the candidate autism gene nrx-1 shows increased number of ectopic synapses in the ladder structure formed in the CNS by the axonal projection of nociceptive neurons. These findings offer the basis for investigating the processes leading to the failure in the elimination of ectopic synapses providing insights into the molecular mechanisms regulating synaptic refinement.

Histone deacetylase 4 (HDAC4) is implicated in several neurodevelopmental and neurodegenerative diseases that involve deficits in memory and cognition. Increased expression of HDAC4 in the Drosophila brain impairs neuronal development and memory, thus Drosophila is an ideal model to investigate the molecular pathways through which HDAC4 acts. A recent genetic screen in Drosophila, for genes that interact in the same molecular pathway as HDAC4, identified the cytoskeletal adaptor Ankyrin2 (Ank2). Knockdown of Ank2 in the brain resulted in deficits in axon morphogenesis (Fisher’s, p < 0.01) with reduced elongation and guidance defects as well as significantly reduced dendritic branch lengths (Student’s t-test, p < 0.05), all of which are similar phenotypes to those resulting from increased expression of HDAC4. HDAC4 contains a putative ankyrin-binding motif, suggesting that it may interact physically with Ank2, however no interaction was detected via co-immunoprecipitation. Further investigation revealed that expression of HDAC4 with a mutated ankyrin-binding motif retained the ability to interact genetically with Ank2 to synergistically impair photoreceptor development (ANOVA, p < 0.01), furthermore, this genetic interaction was dependent on the presence of HDAC4 in the nucleus. Together these data show that Ank2 and nuclear HDAC4 indirectly interact to regulate neuronal morphogenesis and function.

Synapses are connections between a presynaptic neuron and a postsynaptic cell, proper formation of synapses are needed for neuronal signaling. The ability to make and prune synapses is referred to as structural synaptic plasticity. The perturbation of plasticity is associated with many disease states. We use the Drosophila neuromuscular junction (NMJ) to study synaptic plasticity. Previous work in our lab shows that the Drosophila homolog of Activity-regulated cytoskeleton associated protein (dArc1), a retrovirus-like protein, forms capsids, transfers its own RNA in a viral-like fashion across the NMJ modulates, and finally that this transfer is needed for structural synaptic plasticity. However, the mechanism of how dArc1 regulates plasticity, and how its transfer is involved in this regulation is unknown. Here, using dArc1 knock downs in motor neurons followed by deep sequencing, we have found a subset of mRNAs that are decreased in muscle.

These represent RNAs that might depend on dArc1 for transfer across the synapse. Using this approach, we found that the Drosophila homolog of muscleblind (mbl) in fly muscles is decreased significantly with dArc1 knocked down in motor neurons. Consistent with a transfer of mbl across the NMJ, over-expression of mbl in the neuron causes a significant increase of mbl RNA both pre- and post- synaptically. Further, co-immunoprecipitation experiments suggest dArc1 protein binds to mbl RNA. Strikingly, both mbl knock down and mbl-over-expression cause a reduction of structural synaptic plasticity. Taken together, mbl mRNA is transferred across the NMJ and may be a cargo of dArc1 capsids and genetically it appears mbl is needed for structural synaptic plasticity. We are now investigating if the known functions of Mbl, such as regulating alternative splicing or RNA localization is the mechanism by which changes in Mbl expression affects plasticity.
Peroxisome function in glial cells has been shown to impact neuronal functions. In the present study, we aim to elucidate the mechanism behind peroxisome-mediated glia-neuron communication using *Drosophila* as a model organism. I used the GAL4/UAS system to drive RNAi knockdown or overexpression in either glia tissue or motor neurons. The morphology of adult abdominal neuromuscular junction (NMJ) was visualized using confocal microscopy and the axonal volume was quantified via ImageJ. Glial-specific knockdown of peroxisome import protein, Pex5, results in swollen axon as compared to the control. Consistent with our previous work showing that defective peroxisomes upregulate inflammatory cytokine upd3 and JAK-STAT signaling, overexpression of upd3 in glia phenocopies glial-specific knockdown of Pex5 and increases axonal volume and area. We further show that neuronal-specific activation of the JAK-STAT pathway through hop overexpression results in an increase in axon size. Taken together, our findings suggest that impairment of peroxisomes in the glial impacts axonal morphology and potentially functions via inflammatory response, specifically the JAK-STAT pathway.

875V  A comprehensive temporal patterning gene network controls developmental timing in *Drosophila* medulla neuroblasts

During development, neural progenitors are temporally patterned to sequentially generate distinct neural types. Previous studies showed that sequential expression of five temporal transcription factors (TTFs), Homothorax (Hth), Eyeless (Ey), Sloppy paired (Slp), Dichaete (D) and Tailless(Tll), in medulla neuroblasts (NBs) of *Drosophila* larval brain is necessary for generating the full spectrum of neurons in a defined order. However, these TTFs identified through candidate antibody screening may not compose the complete TTF sequence, and several gaps remained unfilled concerning the initiation, termination and regulation of this temporal cascade. To address these questions, we applied single cell RNA sequencing (scRNA-seq) to our model system to discover all unknown TTFs and additional determinants, as well as to get a global view of the dynamic temporal patterning process of medulla neuroblasts. After analyzing the scRNA-seq data of medulla neuroblasts, two sets of genes showed high to low or low to high gradients of expression as neuroblasts age. With known TTFs as markers, we identified a list of novel TTF candidates, among which, SoxNeuro (SoxN), doublesex-Mab related 99B (Dmrt99B), Odd paired (Opa), Earmuff (Erm), Scarecrow (Scro), BarH1, BarH2 and Glial cells missing (Gcm) are necessary for medulla temporal patterning. We identified extensive cross-regulations among these novel TTFs and known TTFs, that generally follow the rule that a previous TTF is required to activate a later TTF, while a later TTF would repress a previous TTF. Our study revealed a comprehensive temporal patterning cascade: Hth + SoxN + dmt99B -> Opa -> Ey+Erm -> Ey+Opa -> Slp+Scro -> D -> B-H1&2->Tll, Gcm, which controls the sequential generation of different neural types by regulating the expression of specific neuronal TFs. With Dmrt99B, Opa and Gcm discovered as TTFs, the mechanisms for the initiation and termination of the temporal cascade were uncovered. Moreover, in pursuit of the mechanism behind the regulation of the temporal cascade, we found that the initiation, progression and termination of the TTF temporal cascade require genes differentially expressed along the differentiation axis (NBs -> neurons) including Lola and Nerfin-1, providing clues as to why the temporal progression only happens in neuroblasts but not their differentiated progeny. In summary, our study revealed a comprehensive temporal gene network that controls developmental timing in neuro progenitors.

876B  Natural genetic modifiers of sensitivity to dopamine-level perturbations in *Drosophila* melanogaster

Dopamine (DA) plays a major role in many animal behaviors, and yet its level is highly variable. It naturally changes over the lifetime in concert with physiological states and environmental stresses. In order to maintain stable expression of many important downstream behaviors, DA level homeostasis needs to be regulated. The way DA homeostasis is achieved on a cellular level has been under extensive investigation, due to its role in Parkinson’s disease. However, genetic variation underlying these traits remains relatively unexplored. The extensively characterized genetic diversity existing in the *Drosophila melanogaster* Genetic Reference Panel, as well as the utility of *Drosophila* neurogenetic toolkit, enables us to pursue this question in a systematic way. Here, we use pharmacological interventions to perturb DA level in diverse genetic backgrounds of the DGRP. This enabled us to simulate exogenously induced dopamine perturbations while avoiding confounding the effects of dopamine level with the systemic phenotypic response to a specific environment. We administered L-DOPA (dopamine precursor) and 3IY (dopamine-precursor agonist) to the DGRP lines and then measured changes in locomotion across genotypes. This enabled us to explore the interaction in 193 genotypes and three dopamine states (nominal, elevated and depleted). Using genome-wide association study we then identified new genetic modifiers of perturbed dopamine phenotypes. These variants point to novel as well as known pathways that affect DA signaling. Namely, we find that the natural genetic variation of enzymes along the cAMP signaling cascade, in the octopamine synthesis, as well as sulfotransferase pathways may play an important role in maintaining locomotion upon perturbation of DA levels. In this study we explore and discuss their functional relevance.
Neuropeptides and their cognate receptors are critical molecular effectors that modulate many physiological processes and behavioral outputs. We recently reported that Glucose-6-phosphatase, one of the key gluconeogenic enzymes, is exclusively expressed in a small subset of about 30 large neurosecretory cells in the CNS of the fruit fly. Moreover, we demonstrated that neural activities of G6P neurons, but not glucose production per se, is critical for glucose homeostasis. These neurons are essential to maintain systemic glucose homeostasis, yet their identities are largely unknown. Here we show that G6P-GAL4 neurons express at least five distinct neuropeptides such as NPF, Orcokinin, PDF, FMRFa and Nplp1. Neural silencing of each class of neuropeptide neurons showed that only the FMRFa-producing neurons in the thoracic ganglion are essential for glucose homeostasis. FMRFa mutants show a similar hypoglycemic phenotype as G6P mutants, suggesting that G6P maintains systemic glucose homeostasis via FMRFa signaling. To support this idea, we measured secretory capabilities of FMRFa neurons using a neuropeptide reporter, a mammalian atrial natriuretic peptide fused to GFP (ANF-GFP). We found that G6P mutants show significantly lower amounts of ANF-GFP in the hemolymph, compared to G6P wildtype flies. These results indicate that G6P cell-autonomously increases circulating neuropeptide amounts by facilitating neuropeptide processing, transport and/or release. Taken together, our data suggest that G6P maintains systemic glucose homeostasis by increasing FMRFa amounts in the hemolymph.

Effectively acting on the demands of internal states is vital for survival. For example, when an animal experiences hunger, signals from nutrient sensors must be transmitted to its brain to orient its behavior towards acquiring sustenance. The internal state of hunger and the sensory inputs relevant to finding food must be integrated and transformed into action. Neuropeptides signal on a variety of scales and change the dynamics of neural circuits (Bargmann 2012). The majority of neuropeptide functions are unknown, yet their purported mechanisms make them well suited to signal internal state.

Recent advances allow for specific genetic access to around 40 neuropeptide producer and receptor neuron populations (Deng 2019). By placing these flies in a 1x1 meter enclosed environment and tracking their movement, we can precisely track individuals during behavioral bouts. An automated fly dispenser allows for the testing of various genotypes and conditions. Screens completely surrounding the fly allow for both static and dynamic visual stimulus presentation. To test the effects of various neuropeptide neurons on behavior, we activate them via optogenetic stimulation during the trial. This system allows for the first unbiased behavioral screen of the effects of a large variety of neuropeptides in response to stimuli, as well as a comparison between those effects and that of behavioral and physiological states of individual flies.

Through measurements of the fly’s position and heading, we quantify various behavioral metrics of individuals in response to visual stimuli, states, and neuropeptide neuron stimulation. We have found that internal states such as hunger and mated status of flies lead to dramatic changes in the animal’s locomotor behavior and that state changes alter the relationship between behavior and visual stimuli. Optogenetically activating the neuropeptide neuron populations also alters locomotor behavior generally and its relationship to visual stimuli, sometimes with similar effects to changing internal states.

These experiments enable us to probe the effects of many neuropeptides and states on behavior. The goal is to characterize the effect of neuropeptides in interpretable ways as they reflect and manipulate internal states of the animal. The ultimate goal is to build a circuit-level understanding of neuropeptide and state computations that transform sensory inputs into behavior, helping animals reach their goals.
supporting that glia (and not unintended neuronal expression) contribute to the motor behaviors. To identify which glial subtype(s) produced the accordion phenotype and full body relaxation, we expressed GtACR1 and dTRPA1 in various glial populations. We found that GtACR1 expression in the perineurial and subperineurial glia produced the accordion phenotype, and dTRPA1 expression in the cortex, ensheathing, and wrapping glia produced the full body relaxation. Our results support that changes in glial ion gradients allow glia to directly influence the production of motor outputs.

**880C** A non-nuclear NF-κB modulates behavioral alcohol sensitivity but not immunity Thilini Wijesekera, Zheng Wu, Nicole Stephens, Rahul Godula, Linda Kao Lew, Nigel Atkinson The University of Texas at Austin

NF-κB proteins are well known as transcription factors important in immune system activation. In this highly conserved role, they contribute to changes in behavior in response to infection and in response to a variety of other insults and experiences. In some mammalian neurons, NF-κBs can be found at the synapse and translocate to the nucleus when activated by synaptic activity. In these areas, NF-κBs are believed to produce change by acting in the nucleus to alter gene expression. Here we demonstrate that in *Drosophila melanogaster*, NF-κB action is important both inside and outside the nucleus and that the Dif gene has segregated nuclear and non-nuclear NF-κB action into different protein isoforms. The DifA isoform is a canonical nuclear-acting NF-κB protein that enters the nucleus and is centrally important for combating infection. The DifB variant, but not the DifA variant, is expressed in central nervous system neurons. DifB does not enter the nucleus and co-localizes with a synaptic protein. Mutations specific to DifB alter alcohol behavioral sensitivity but have no obvious effect on combating infection, whereas DifA mutants do not affect alcohol sensitivity but compromise the capacity to combat infection. These data are evidence that the unusual, non-nuclear DifB variant, alters behavior by a synaptic mechanism that is local and nongenomic and that diverges from the canonical NF-κB transcriptional effects used in the peripheral immune system. The Drosophila DifB variant is an ideal system for studying non-nuclear NF-κB effects in isolation from NF-κB action in the nucleus.

**881A** The functionally conserved neuronal pseudokinase Aln retrogradely regulates homeostatic UPR and autophagy responses in photoreceptor neurons. Shashank Shekhar, Andrew T Moehlman, Minh-Nguyet Hoang, Charles Tracy, HelmutKrämer UT Southwestern Medical Center

Abstract:
Homeostatic responses are essential for neuronal circuits to adapt to changes in the environment. To investigate the molecular mechanisms controlling homeostatic responses in neurons we use a reversible in-vivo model [1]. Extended exposure to ambient light triggers homeostatic responses including reduced rhabdomere size and loss of synaptic active zones in *Drosophila* photoreceptor neurons. These responses are reversible and therefore offer an excellent opportunity to study signaling events controlling neuronal homeostatic responses. Here, we identify aln as a novel gene required for the regulation of these responses. *aln* encodes a secreted pseudokinase that is functionally conserved in humans. Flies mutant for *aln* have normal visual responses when maintained under a regular 12h:12h light:dark cycle (LD), but lose photoreceptor structural integrity and postsynaptic responses after prolonged exposure to ambient light. This response is transient, and photoreceptors recover when returned to LD. Phenotypes are rescued by expression of Aln or its human homolog, the divergent protein kinase FAM69C, independent of its kinase activity. Aln is expressed in many neurons but, surprisingly, not in photoreceptors. Instead, it is secreted by lamina neurons, most prominently L4, and retrogradely regulates photoreceptor responses including UPR and autophagy, two key pathways of homeostatic responses. Perturbation of photoreceptor output by expression of Shi or Tetanus Toxin light chain drastically decreased Aln expression in L4 lamina neurons, and other brain regions. This suggests that the Aln is part of feedback loop in the Drosophila visual system. Aln homologs are highly expressed in mammalian brains suggesting important yet to be discovered functions of these molecules in the mammalian brain.


**882B** The CHD protein, Kismet, regulates both clathrin-mediated and activity-dependent bulk endocytosis at the *Drosophila* neuromuscular junction Emily Hendricks, Taylor Delaney, Faith Liebl Southern Illinois University Edwardsville

Chromodomain helicase binding domain (CHD) proteins, including CHD7 and CHD8, remodel chromatin to alter transcriptional programs. Both proteins are important for proper neural development as heterozygous mutations in *Chd7* and *Chd8* are causative for CHARGE syndrome and correlated with autism spectrum disorders, respectively. Their roles in mature neurons are poorly understood despite influencing the expression of genes required for cell adhesion, neurotransmission, and synaptic plasticity. The *Drosophila* homolog of CHD7 and CHD8, Kismet (Kis), promotes neurotransmission, endocytosis, and larval movement, and restricts synaptic levels of cell adhesion molecules. Endocytosis is essential in neurons for replenishing synaptic vesicles, maintaining protein localization, and preserving the size and composition of the presynaptic membrane. Endocytosis can occur via clathrin-mediated endocytosis (CME), which is coupled with neural activity and is the most prevalent form of synaptic endocytosis, and activity-dependent bulk endocytosis, which occurs during periods of intense stimulation. We assessed transcripts important for endocytosis...
in kis mutant central nervous systems. AP2α, dap160/intersectin, and endophilin B (endoB), required for CME, show differential expression in kis mutants. Rab11, akt, PI3K92E, and sgg/GSK3β, required for ADBE, exhibit reduced transcripts levels. Finally, syndapin transcripts, required for both CME and ADBE, were reduced while several other transcripts were unaffected by mutations in kis. We were able to restore endoB, PI3K92E, rab11, and sgg transcripts in kis mutants by expressing kis in either neurons or postsynaptic muscle. These data indicate that Kis contributes to both CME and ADBE. We further tested this hypothesis by pharmacologically inhibiting either CME or ADBE in kis mutants and examined endocytosis both electrophysiologically and functionally. Collectively, our data indicate that Kis promotes both CME and ADBE by promoting transcription of several endocytic genes.

883C Investigating the Effects of Rab11 on Synaptic Proteins FasII and APPL in kismet Mutants Ireland Smith, Joshua Preston, Faith Liebl Southern Illinois University Edwardsville

Chromodomain Helicase DNA-Binding Domain (CHD) proteins are chromatin remodelers that are critical for early neuronal development and the expression of genes important for synaptic plasticity. Mutations in CHD7/8 can lead to a variety of disorders in humans such as CHARGE syndrome, a neurodevelopmental disorder, and autism spectrum disorders. Mutations in kismet (kis), the Drosophila ortholog of CHD7/8, lead to an accumulation of cell adhesion molecules and the transmembrane protein amyloid precursor protein-like (APPL) at the synapse as well as reductions in endocytosis, synaptic transmission, and larval locomotion. Kismet regulates rab11 expression. Rab11 is a GTPase that traffics vesicles from the recycling endosome to the plasma membrane. Because of its role in membrane trafficking, Rab11 might facilitate some of these phenotypic changes observed in kismet mutants.

To investigate this possibility, we expressed constitutively active or dominant negative isoforms of Rab11 and examined Fasciclin II (FasII), a cell adhesion molecule. Similar to kis mutants, constitutively active Rab11 increased synaptic FasII. Expression of dominant negative rab11, however, decreased FasII protein levels. When rab11 was knocked down, we found a decrease in APPL, which directly contrasts kis mutants that show an increase in APPL levels. In addition, when Rab11 dominant negative and constitutive active isoforms were expressed, we saw no change in endocytosis or endocytic proteins Endophilin A (EndoA) and Dynamin (Dyn) unlike kis mutants, which show reductions in each. Kis affects genes, possibly rab11, that localize synaptic proteins to phosphatidylinositol-4,5-\(\text{P}_2\) (PIP\(_2\)), a phospholipid that helps organize proteins into lipid rafts in the plasma membrane. When rab11 was knocked down, we did find a reduction in PIP\(_2\) levels at the synapse. These data may suggest that while Rab11 might only play a minor role in contributing to kis mutant phenotypes, Rab11 may be able to restore kis mutant phenotypes. Therefore, we will express wildtype Rab11 in kis mutants to determine whether Rab11 can restore the synaptic levels of FasII and APPL.

884A The Drosophila CD63-related tetraspanins, Tsp42Ee and Tsp42Eg, regulate synaptic structure, function, and vesicle pool dynamics Emily Hendricks, Faith Liebl Southern Illinois University Edwardsville

Tetraspanins are a well-conserved class of transmembrane proteins that play a regulatory role at the cell membrane by mediating the spatiotemporal distribution of their binding partners—often receptors, cell adhesion molecules, and intracellular signaling proteins. The tetraspanin, CD63, is a classical marker of exosomes and late endosomes and likely regulates vesicular traffic through the exocytic pathway. However, CD63’s role in mediating endocytic and exocytic dynamics at the synapse is not well described. Thus, we aim to use the Drosophila neuromuscular junction (NMJ) to characterize the role of CD63 orthologs, Tsp42Ee and Tsp42Eg, in regulating synaptic vesicle dynamics. We find that CD63-related tetraspanins negatively regulate endocytosis, likely through control of synaptic vesicle pool dynamics and localization of the synaptic proteins, Endophilin A and Dynamin. We also find that Tsp42Ee and Tsp42Eg regulate synaptic morphology and function as both tsp mutants show reductions in locomotor function and mEJC frequency. Furthermore, tsp42Eg loss of function mutants show reduction in quantal content and eEJC amplitude suggesting an overall reduction in synaptic vesicle release. These alterations in synaptic function may, at least in part, be attributed to disruptions in cytoskeletal structure as we observe an increase in Futsch loops at overall reduction in synaptic vesicle release. These alterations in synaptic function may, at least in part, be attributed to disruptions in cytoskeletal structure as we observe an increase in Futsch loops at overall reduction in synaptic vesicle release. Furthermore, tsp42Eg loss of function mutants show reduction in quantal content and eEJC amplitude suggesting an overall reduction in synaptic vesicle release. These alterations in synaptic function may, at least in part, be attributed to disruptions in cytoskeletal structure as we observe an increase in Futsch loops at overall reduction in synaptic vesicle release. We will further explore this work by attempting to rescue the tsp mutant phenotype through expression of human CD63 in the tsp mutant backgrounds. These experiments will aim to validate the functional redundancy of hCD63 in Drosophila and further clarify the role that tetraspanins play at the synapse.

885B Uncovering the Genetic Basis of Variation in Learning and Memory Phenotypes using the Drosophila Synthetic Population Resource Victoria Hamlin, Huda Ansf , Patricka Williams-Simon , Elizabeth King University of Missouri

Learning and memory are complex traits in which many genes and regulatory regions affect the phenotypic output of an individual. For animals, these traits are vital functions necessary for adapting to and surviving in an ever-changing environment. Within a given population, there is variation between individuals in their ability to perform these functions, however, the mechanisms underlying this variability are still largely unknown. In addition, some genotypes may show more variable performance than others. Utilizing variance quantitative trait loci (vQTL) mapping, we can identify regions of genetic variation associated with differences in the variability of learning and memory performance in the Drosophila
Synthetic Population Resource (DSPR). This multiparent population consists of about 1800 recombinant inbred lines (RILs) which provides us with ability to perform high resolution genome wide scans to identify quantitative trait loci with significant contribution to the residual variability in learning and memory performance. To test the performance of RILs, flies underwent operant conditioning for place and olfactory learning and memory. In the aversive place learning assay, flies were placed in a heat box that would increase to an intolerable temperature when flies crossed into one side of the chamber followed by testing to see if they remained on the cool associated side after training. In the appetitive olfactory assay, flies are starved for eighteen hours then provided a sugar reward paired with an odor in training followed by a Y-maze choice test to see if they selected the positive conditioned odor. Using a double generalized linear model to detect both mean and variance QTLs, we identified several QTL influencing these key phenotypes.

886C Investigating the role of tRNA methyltransferase ALKBH8 in learning and memory Shanzeh Sayied1, Kim Madhwani1, Caley Hogan2, Kate O’Connor-Giles1,3 1) Brown University; 2) University of Wisconsin-Madison; 3) Carney Institute for Brain Science

Regulation of neuronal gene expression is crucial for proper neuronal development and function and occurs at numerous steps, from DNA methylation to protein degradation. Transfer RNAs (tRNAs) are emerging as an important point of regulation. tRNAs are heavily post-transcriptionally modified to regulate tRNA structure and stability and strengthen anticodon-codon interactions. Recently, variants in tRNA-modifying enzymes have been associated with multiple neurological disorders. In particular, mutations in ALKBH8, which encodes a highly conserved tRNA methyltransferase, have recently been linked to intellectual disability in four families. While ALKBH8 has been studied in the context of oxidative stress, its role in the nervous system remains unknown. Due to its role in translation and link to intellectual disability, we hypothesized that ALKBH8 may promote the translation-dependent processes of learning and memory. We generated ALKBH8 null mutants and analyzed sensory memory using a taste aversion assay. The taste aversion assay measures a reduction in the sucrose-induced proboscis extension reflex (PER) after starved female flies learn to associate sucrose with the aversive taste quinine. Our findings reveal a deficit in the formation of aversive taste memories in ALKBH8 null mutants. These results indicate learning deficits consistent with intellectual disability in humans and establish Drosophila as a model for understanding the underlying mechanisms of ALKBH8-associated intellectual disability.

887A Utilizing Y-mazes to Investigate Olfactory Learning Phenotypic Variations in Drosophila Huda Ansaf, Victoria Hamlin, Elizabeth King University of Missouri, Columbia, MO

Fruit flies, and other animals, can navigate in a complex chemosensory environment. Certain odors can act as signals for food, danger, or as pheromones released by conspecifics to elicit innate behavioral responses. Since not all odor stimuli are informative, the brain must make sense of the complexity of the odor signals and interrupt their actual relevance. Fruit flies perceive odors through olfactory sensory neurons that are located in the sensilla on the third antennal segments and the maxillary palps, which exhibit diverse morphological types. The olfactory system of Drosophila resembles systems found in vertebrates in its overall anatomical organization but is significantly reduced in terms of cell number, which makes it an ideal model system to investigate odor processing in the brain. Associative learning represents one process by which new or altered relevance is assigned to a stimulus through experience. For example, an odor that is repetitively paired with a food reward becomes attractive while an odor that often occurs simultaneously with a punishment becomes a predictor for a negative situation and will be avoided in the future. Drosophila melanogaster can easily perform such learning and memory tasks and represents an excellent organism to investigate the neuronal mechanisms that control olfactory learning processes. In this work, we identify phenotypic variations based on learning and memory abilities among different recombinant inbred lines (RILs) in a mapping population. Operant conditioning is induced by presenting an odor with or without the availability of a sucrose reward to condition odors as positive (sugar associated) or negative stimulus. During the testing phase, the Drosophila are simultaneously presented with the previously conditioned CS+ and CS- odors. After sufficient time has passed, fly distribution is recorded to allow for associative appetitive conditioning to be reliably measured without bias due to innate preferences. Various control experiments are also done to test whether all genotypes respond normally to odor and light alone. Learning and memory tasks are tested using a custom-built Y-maze apparatus. After repeating Y-maze testing three times, we identified high and low learning phenotype groups and high and low memory phenotype groups.

888B A survey of cis-regulatory fragments from the dissatisfication gene identifies a subpopulation of abdominal interneurons that regulate the opening of the vaginal plates during courtship Julia Diamandi1, Julia Duckhorn1, Jessica Cande1, Troy Shirangi1 1) Villanova University, Villanova, PA; 2) Cooley LLP, Boston, MA

In Drosophila, virgin females carrying mutations in the dissatisfication gene (dsf) are delayed in mating with males and exhibit deficits in opening their vaginal plates during courtship. Dsf contributes to female behavior in part by functioning in a sexually dimorphic population of interneurons in the abdominal ganglion called the dda neurons. The dda neurons in females are composed of a variety of neuroanatomical subtypes, but which subtypes matter for female behavior is unknown. Here, we screened seven cis-regulatory fragments from the dsf gene for reporter expression in dsf-
expressing neurons in the larval and adult central nervous system. We find that most of dsf’s expression in the central nervous system is driven by cis-regulatory sequences within dsf’s third intron. Using these fragments, we identified a subpopulation of local female-specific ddag interneurons that is sufficient for the opening of the vaginal plates in virgin females during courtship. The regulatory fragments targeting this subpopulation of neurons contain putative binding sites to proteins encoded by the sex determination gene, doublesex, suggesting that doublesex may directly regulate dsf expression in a subset of ddag neurons. Our results provide new insights into the neural circuits that mediate the opening of the vaginal plates and the mechanisms that regulate dsf expression in the central nervous system.

889C Pleiotropy and the rapid coevolution in reproductive traits in Drosophila Mehrnaz Afkhami, John Masly University of Oklahoma

Reproductive traits are some of the most rapidly evolving traits within species, and some of the most rapidly diverging traits between nascent species. Sex-specific enhancer pleiotropy can provide a base for male and female coevolution. Here, we study two genes, Pox neuro (Poxn) and CG14567, that specify the morphology of a male-specific reproductive structure. The epandrial posterior lobes (ePLs) are novel reproductive structures on the male genitalia that are structurally important for mating success. In addition, morphological variation in shape and size of the ePL affects the male's mating success and mating position. When males with a smaller or misshapen posterior lobe mate with a normal female, the female reduces her egg-laying amounts. Previous studies show that Poxn directs aspects of male courtship behavior and genital development, and recent studies from our lab show that variation in expression level of CG14567 in the developing ePL primordia directs variation in genital morphology among members of the D. melanogaster species complex.

Interestingly, one Poxn enhancer region that directs expression in the developing male genitalia is also expressed in three neurons in the adult female abdominal ganglion. These three neurons enervate the female reproductive tract as well as regions of the gastric tract. This Poxn enhancer is also expressed in collections of cells throughout the female VNC and Brian, but compared to males the domain of expression is much reduced. Our preliminary in-situ experiments show that CG14567 is also expressed in the brain and VNC of adult females and appears to localize to these Poxn-expressing neurons. We are currently working on behavioral assays to pinpoint the exact function of three neurons in the abdominal ganglion and we are using the GFP Reconstitution Across Synaptic Partners (GRASP) technique to identify the presynaptic cell(s) to these neurons in the females. Our initial functional genetic experiments show that both Poxn and CG14567 expression in these neurons affect female reproduction. In particular, overexpression of CG14567 in Poxn expressing cells increases the eggs laid by the female. Our preliminary genetic and behavioral data suggests that enhancer pleiotropy at Poxn and CG14567 could contribute to coevolution of reproductive traits.

890A Effects of L-DOPA on D. simulans and D. sechellia Mating Behavior Alyssa Cortés, Sofia Pogliano, Joseph Coolon, Ivy Lam, Sadie Gregory, Charlotte Freeland, Charity Russell Wesleyan University

The absence of mating between groups of organisms is one criterion for the separation of species. There could be multiple reasons for this absence, including geographical, behavioral, and genetic causes. The fruit flies Drosophila sechellia and Drosophila simulans inhabit the same geographical space, the Seychelles archipelago, yet remain separate species due to considerably low levels of hybridization. Previous studies have shown that their lack of interspecies mating is due to male mate discrimination, and that this mate choice is made by discrimination against cuticular hydrocarbon (CHC) profiles, which are unique to the two species. Additionally, it has been shown that chemicals uniquely present in D. sechellia’s food source, Morinda citrifolia (eg L-DOPA), affects the expression levels of genes related to CHC production. Given the likely effects of these species’ differing diets, and more specifically their consumption of L-DOPA on their CHC profiles, and the importance of these profiles in male mate choice, we investigated the consequence of dietary L-DOPA on intraspecific and interspecific mating behaviors. Here we show that L-DOPA does affect mating behavior, and this is mostly seen in D. sechellia flies, those that regularly consume L-DOPA in the wild. These results reveal that mating seen among these flies in the wild is influenced by the presence of L-DOPA. Given this, L-DOPA should be included in laboratory studies regarding mating behavior to mimic the flies’ natural conditions. Including L-DOPA would create a more natural environment and lead to mating behaviors more like those produced in the wild, generating more accurate results. Our results could be indicative of complex interactions between the environment and the genetic basis of mating behaviors, and thus support motivations for more in-depth studies of the effect of L-DOPA on various genes and the overall CHC profiles of D. simulans and D. sechellia. This would further clarify the mechanisms of their speciation.

891B Impact of histamine deficiency on accessory gland secondary cell differentiation, persistence, and post-mating responses in Drosophila melanogaster Cazmir Sarnacki, Justin Lilley, Ryan Blackmer, Jonathan Wassink, Erika Spafford, Jordan Yokubonusi, Eric Gonzales, Lauren Gerritsen, Anna Prince, Martin Burg

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The study of histamine and its function in Drosophila melanogaster has primarily focused on its role as a neurotransmitter used by photoreceptors and its effects on a number of other behaviors, such as grooming, thermal
preference, and sleep. Recently, we have identified histamine-like immunoreactivity in a vacuole-like compartment (VLC) of secondary cells in the male accessory gland. This immunoreactivity is eliminated by null mutations in the \textit{Hdc} gene that encodes the enzyme histidine decarboxylase. Previous studies have shown that secondary cells are responsible for enabling a number of post-mating responses including increased egg laying, enhanced sperm viability, and reduced female receptivity to copulation. For this analysis, Oregon-R female flies were conditioned with either wild-type, \textit{Hdc}^{K910}, or \textit{Hdc}^{P211} males, followed by wild-type males at 1, 4, and 10 days after the conditioning copulation. In the current study, we demonstrate that elimination of histamine via the \textit{Hdc}^{P211} or \textit{Hdc}^{K910} mutations appears to alter the female receptivity to copulation 1 day after copulation, increasing receptivity when compared to wild-type. To determine whether the loss of histamine was affecting this phenotype by disrupting the differentiation of secondary cells (or altering their numbers), we examined wild-type, \textit{Hdc}^{K910}, or \textit{Hdc}^{P211} male accessory glands in which secondary cells were marked by GFP labeling. The wild-type and mutant flies were examined at several different ages for the presence of histamine, \textit{dlg} protein as well as GFP to determine whether large deviations were observed in the number of secondary cells or in the patterning of the accessory gland. Results for both mutant \textit{Hdc} alleles indicate that secondary cells are still able to differentiate when histamine is absent in the accessory gland, but the number of cells present appears to be consistently reduced when compared to wild-type flies.

**References:**
2. Oh et al., 2013, PLoS ONE 8(7):e68269
3. Burg et al., 2019, 60th Drosophila Research Conf., Poster 676

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**892C**  **Mechanisms of D2R signaling in the blood-brain barrier that regulates courtship in \textit{Drosophila melanogaster}** **Sumit Gautam,** Cameron R. Love, Brigitte Dauwalder University of Houston, Houston, TX

The blood-brain barrier (BBB) is a highly selective cell layer that separates the nervous system and the circulating hemolymph. The BBB protects the brain from the contents of the circulating fluids that could impede neuronal function. Two layers of glial cells, Perineural Glia (PG) and Subperineural Glia (SPG) make up the BBB in \textit{Drosophila melanogaster} which ensheathes the brain. We have previously shown that adult feminization of BBB cells in male \textit{Drosophila} leads to reduced courtship levels. In a microarray screen of isolated BBB cells we have identified several male-enriched transcripts. One of them encodes the \textit{Dopamine-2 like receptor (D2R)}. We discovered that conditional knockdown of D2R in adult male \textit{Drosophila} BBB decreases courtship levels. The D2R receptor is highly conserved and intriguingly, this receptor has been found to act via biased signaling (via G protein or arrestin) in mammals. We are interested in understanding the downstream signaling of D2R in the BBB of \textit{Drosophila} that mediates the effect on courtship. We are examining biased signaling by the D2R receptor by mutagenizing D2R receptor residues that favor G protein or arrestin signaling, respectively. These constructs will be tested for their ability to rescue the D2R courtship defects when they are expressed in adult male \textit{Drosophila} BBB. This will help us better understand how D2R contributes to the regulation of courtship in \textit{Drosophila}.

**References:**

**893A**  **Regulation of sexually dimorphic abdominal courtship behaviors in \textit{Drosophila} by the Tlx/tailless-like nuclear receptor, Dissatisfaction** **Julia Duckhorn,** Jessica Cande, Mary Metkus, Hyeop Song, Sofia Altamirano, David Stern, Troy Shirangi 1) Villanova University, Villanova PA; 2) Janelia Research Campus, Ashburn VA

Sexually dimorphic courtship behaviors in \textit{Drosophila melanogaster} develop from the activity of the sexual differentiation genes, \textit{doublesex (dsx)} and \textit{fruitless (fru)}, functioning with other regulatory factors that have received little attention. The \textit{dissatisfaction} gene (\textit{dsf}) encodes an orphan nuclear receptor homologous to vertebrate \textit{Tlx} and \textit{Drosophila tailless} that is critical for the development of several aspects of female- and male-specific sexual behaviors. Here, we report the pattern of \textit{dsf} expression in the central nervous system and show that the activity of sexually dimorphic abdominal interneurons that co-express \textit{dsf} and \textit{dsx} is necessary and sufficient for vaginal plate opening in virgin females and abdominal curling in males during courtship. We find that \textit{dsf} activity results in different
neuroanatomical outcomes in females and males, promoting and suppressing, respectively, female development and function of these neurons depending upon the sexual state of dsx expression. We posit that dsf and dsx interact to specify sex differences in the neural circuitry for dimorphic abdominal behaviors.

894B  A Drosophila model for understanding the perception and central processing of chronic social isolation  Wanhe Li Texas A&M University

Social isolation and loneliness have potent effects on public health. Social psychologists have suggested that compromised sleep quality is a key factor that links persistent loneliness to adverse health conditions. We report that chronic, but not acute, social isolation reduces sleep in Drosophila. We use quantitative behavioral analysis and transcriptome profiling to differentiate brain states under acute vs. chronic social isolation. Despite animals’ uninterrupted access to food, chronic social isolation altered the expression of metabolic genes and induced a brain state that signals starvation. Chronically isolated animals exhibit sleep loss accompanied by overconsumption of food, which resonates with anecdotal findings of loneliness-associated hyperphagia in humans. Chronic social isolation reduces sleep and promotes feeding through neural activities in the fan-shaped body columnar P2 neurons of the fly. Activation of these neurons causes misperception of acute social isolation as chronic social isolation and thus results in sleep loss. These and other new results reveal mechanistic links between chronic social isolation, metabolism, and sleep, presenting a new genetic model to understand the perception and central processing of chronic social isolation.

895C  Toll Family Receptor Function in Neuronal Recognition of Immune State  Tim Lebestky, Nicole Alvarez, Minjun Kim, Raphael Rakosi-Schmidt Williams College

The Toll family of receptors possess a number of critical functions in Drosophila, including roles in development and immunity. Recently, our group has begun looking for behavioral roles for the Toll family of receptors in the nervous system, given a breadth of expression in the adult nervous system. We hypothesize that Toll receptors function as an immune sensors in neurons to communicate immune challenge and infection and triggers behavioral responses for sickness behaviors in Drosophila. We will present our findings primarily on Toll1 and Toll7 with regards to sleep and feeding behaviors and their potential relationship to immune state when challenged by Metarhizium anisopliae, entomopathogenic fungi known to trigger robust responses from the Toll1 pathway. We will also share behavioral characterization of sleep and feeding behaviors for wild type animals exposed to M. anisopliae. We believe that immune sensors in the nervous system are essential for coordinating immune-behavioral responses in both sensing presence of pathogens as well as circuit-specific responses during chronic illness.

896A  The Drosophila serotonin transporter (dSERT) is required for proper sleep amount and sleep architecture  Elizabeth Knapp1, Henrike Scholz2, Jeffrey Donlea3, David Krantz1  1) University of California, Los Angeles, Psychiatry, Los Angeles, CA; 2) University of Cologne, Animal Physiology, Köln, Germany; 3) University of California, Los Angeles, Neurobiology, Los Angeles, CA

Sleep is a critical process essential for life and is evolutionarily conserved from insects to mammals. Although sleep disruption has been linked to a variety of neurological and psychiatric disorders, the cellular mechanisms and neural circuitry involved in sleep regulation are not well understood. The biogenic amine serotonin (5-hydroxytryptamine, 5-HT) functions as a key neuromodulator of sleep behavior in both Drosophila and mammals. The relationship between serotonergic signaling and sleep has been studied for several decades, however its complex role in sleep regulation remains uncertain, with many studies showing it to play a role in both wakefulness and conversely sleep propensity. These complexities are further compounded by the fact that many antidepressants and antianxiety medications that work through selective inhibition of serotonin reuptake (SSRIs) have been shown to produce contradicting and various effects on sleep, ranging from insomnia to daytime somnolence.

In both Drosophila and mammals reuptake of serotonin from the synaptic cleft is mediated via the serotonin transporter (SERT). The relationships between sleep, neurological disorders, and SSRI medications strongly suggest that variations in extracellular serotonin levels as a result of increased or decreased SERT activity could play a key role in modulating sleep behaviors. The molecular mechanisms underlying this process, however, are not well understood in either humans or model systems.

In this study, we use Drosophila as a model system to study the mechanisms by which altering Drosophila SERT (dSERT) activity impacts sleep behavior. Here we use novel dSERT mutants to demonstrate that dSERT is required for regulating both proper sleep amount and architecture. In addition, we discovered the increased sleep drive exhibited in these dSERT mutants is differentially impacted by certain circadian and environmental factors. Lastly, our data suggests that distinct serotonergic circuits modulate daytime and nighttime sleep behaviors independently. Overall, our work provides new insight into the molecular mechanisms of neuromodulation in the context of sleep and enhances our understanding of how serotonergic signaling is involved in sleep behavior.

897B  rhodopsin 3 regulates circadian periodicity  Menglin Li, Geoff Meyerhof, Kara Simones, Anvitha Aluri, Craig Montell University of California, Santa Barbara
Circadian rhythms are 24 hour cycles that align behavior, physiology, and gene expression with predictable daily changes in the environment. The primary entrainment cue for endogenous circadian clocks is light. Herein, we sought to test the impact of individual rhodopsins, the light-sensing G-protein-coupled receptors primarily expressed in the eye, on circadian rhythms. We found that flies harboring mutations in rhodopsin 3 (rh3), which is expressed in a subset of R7 photoreceptor cells, have an elongated circadian periodicity. Under constant darkness, wild-type flies maintain a locomotor rhythm with a 24 hour periodicity, whereas the rhythm of rh3 flies is dramatically extended to ~28 hours. This defect in rh3 flies suppresses the morning and evening anticipatory locomotion that usually precedes changes from lights on to lights off. Furthermore, the timing of sleep and feeding is also disrupted in rh3 flies. These defects in circadian behavior are associated with a decreased amplitude of mRNA expression of core-clock components in the periphery, as well as a delay in the phase of clock mRNA expression. When we housed rh3 flies under a 28 hour light:dark cycle (14h light:14h dark), the timing of their sleep and feeding was restored, and their lifespan was extended. Our future work aims to map the neuronal connections linking rh3-expressing photoreceptor cells and the core clock neurons in the brain, which set 24 hour locomotor rhythms. In conclusion, we have shown that rh3 is required for 24 hour circadian periodicity, which affects the daily timing of sleep and feeding. These results suggest that distinct populations of photoreceptor cells are required for maintaining circadian rhythms in Drosophila.

898C  Neuronal E93 Regulates Metabolic Homeostasis Cecilia Yip1, Steven Wyler1, Syann Lee1, Adrian Rothenfluh2, Young-Jai You1, Joel Elmquist1 1) University of Texas Southwestern Medical Center, Dallas, TX ; 2) University of Utah, Salt Lake City, UT

The central nervous system integrates environmental and internal cues to drive physiological and behavioral responses to control energy homeostasis. E93 plays an essential role during metamorphosis, yet its neuronal function in adults has not been fully understood.

To determine the central role of E93, we knocked down E93 expression using nSyb-Gal4 (nSyb>E93), a pan-neuronal driver. Unlike whole-body knockout flies, which cannot progress to eclosion, the nSyb>E93 flies became adults. The nSyb>E93 adults are obese and hyperphagic, with increased energy stores, revealing a novel role of E93 in metabolism. Assessments of whole-body glucose, glycogen, and triglyceride levels indicated a sexual dimorphism in their fuel stores, with males having increased triglyceride stores while virgin females show increased glycogen stores. In order to identify the sites of E93 action, we targeted E93 RNAi in subsets of neurons using 17 Gal4 lines. We found that E93 knockdown specifically in myoinhibitory peptide (MIP) producing neurons and GABA-ergic neurons mimics the phenotypes of nSyb>E93. Repression of the Gal4 driver by Gal80, specifically in MIP neurons, partially yet significantly rescues the E93 phenotypes, confirming the role of E93 in MIP neurons. Knockdown of the ecdysone receptor (ECR), the known upstream signaling molecule of E93 specifically in MIP neurons phenocopies the E93 phenotypes, suggesting that steroid hormone signaling to E93 in MIP neurons regulates metabolism. Metabolic dysregulation is often associated with abnormal circadian rhythm. Indeed, nSyb>E93 exhibits a disrupted circadian rhythm; both males and females show reduced activity during the day and decreased day time period.

Together, our study reveals an important neuronal function for E93 in controlling metabolism and provides an insight into how a transcriptional network downstream of a steroid hormone regulates metabolism and circadian rhythm.

899A  Adaptive variation in taste detection of carboxylic acids Manali Dey1, Anupama Dahanukar1,2 1) Interdepartmental Neuroscience Program, UCR, Riverside; 2) Department of Molecular, Cell and Systems Biology, UCR, Riverside

Comparative studies between generalist and specialist species enable us to learn how adaptations for a particular environment may lead to speciation. Drosophila sechellia, endemic to the Seychelles, is an obligate specialist that almost exclusively uses the fruit of Morinda citrifolia (noni) for feeding and oviposition. By contrast, its generalist relatives D. melanogaster and D. simulans take advantage of overripe fruits but avoid noni fruits, which are rich in short to medium chain fatty acids (FA) that are toxic for them. D. sechellia, however, are attracted to noni and are able to metabolize the fatty acids. To understand the contribution of the taste system in behavioral adaptation of D. sechellia, we undertook a comparative study of gustatory function in the three related species. In two independent behavior assays D. sechellia showed a feeding preference for noni acids, whereas the generalist species showed aversion. Genetic and surgical ablation experiments uncovered a contribution of olfaction in stimulating feeding preference for lower concentrations of noni acids, but largely olfaction-independent behavioral responses at the higher end of the concentration range. A systematic comparison of cellular responses with extracellular tip recordings identified two differences in D. sechellia that may facilitate their interaction with noni fruit – first, the responses of their deterrent (bitter-sensing) neurons to noni acids are weaker, and second, their appetitive (sugar-sensing) neurons are not strongly inhibited by noni acids. Further, D. simulans appears to exhibit phenotypic parameters that are like those of D. melanogaster in some cases, and those intermediate between D. melanogaster and D. sechellia in other cases, suggesting that multiple genetic changes may account for gustatory differences between D. melanogaster and D.
sechellia. Ongoing studies are directed towards understanding the molecular basis of variation in *Morinda* acid taste between the three *Drosophila* species. Preliminary results suggest the involvement of at least three molecularly distinct pathways involving ionotropic receptors, gustatory receptors, and odorant-binding proteins. We are testing evolutionary roles of candidate genes by manipulating function in both *D. melanogaster* and *D. sechellia*.

900B Functional Genetic Screen to Identify Interneurons Governing Behaviorally Distinct Aspects of *Drosophila* Flight Motor Programs

Chung Lee1, Matthew Clark1, Michael Dickinson2 1) Bucknell University, Lewisburg, PA; 2) Caltech, Pasadena, CA; 3) Boston Childrens Hospital, Harvard Medical School, MA

Muscles provide the force necessary to generate movements that drive locomotor behaviors. Muscles innervated by single motor neurons comprise synergistic motor units that can be recruited to perform a given task. *Drosophila* use only a dozen pairs of flight steering motor units to regulate wing motion during both quick maneuvers and slow compensatory reflexes. Steering muscles can be functionally grouped according to their size and patterns of activity. Small tonicly active muscles allow the fly to steer straight via subtle adjustments in the phase of firing relative to wing strokes. Large phasically active muscles allow the fly to quickly alter heading via rapid changes in muscle activity. Although features of these motor neurons and muscles are relatively well characterized, the neural network of interneurons controlling them is poorly understood. To study the relationship between neural circuitry and flight, we performed an optogenetic screen of Split Gal4 drivers specific to ventral nerve cord interneurons (VNC INs) and recorded changes in stroke amplitude and wingbeat frequency. Results showed that activation of VNC INs had a variety of effects compared to controls, suggesting that dedicated groups govern specific wing motions. Using the MultiColor FlpOut method we have begun matching morphological features of individual behaviorally relevant neurons to those reconstructed via comprehensive connectomic mapping. Future functional studies will determine how these various interneurons alter the activity of specific steering motor units and thus alter wing motion.

901C The Effect of Cannabidiol on Central Nervous System Development and Function using *Drosophila* as a Model System

Cameron Lowery, Annika Fischer, Parag Bhatt, Sandra Leal Harris-Stowe State University

Cannabidiol, or (CBD), is a non-psychoactive cannabinoid found in *Cannabis sativa*. While cannabidiol is used by individuals of all ages, little is known about its short and long-term neurological impact on the central nervous system (CNS) or function and development. For this reason, we sought to determine whether CBD oil affects one or more basic behavioral outcomes of the CNS using *Drosophila* as a model system. Several major neurotransmitter systems of *Drosophila* share high conservation with mammalian systems. We assayed three distinct behaviors of Oregon-R wild-type, third-instar larvae under acute (24-hour) and chronic (48-hour) CBD exposure. These behaviors included body wall contractions (BWC), mouth hook contractions (MHC), and the righting reflex (RR). We found that CBD oil differentially affected motor circuits regulating larval MHCs under specific experimental conditions. Acute pretreatment of larvae with 0.5 mg/mL CBD oil significantly reduced larval locomotor behavior but had no significant effect on larval feeding behavior. Conversely, 1.0 mg/mL CBD decreased third-instar larval feeding rates. In comparison to all control group animals, the majority of which righted within 5 seconds or less, the latency of animals righting within the passing score parameters increased for several populations of acute and chronic CBD-treated animals. Since we observed the effects of CBD oil treatment on motor behaviors, we examined neurotransmitter expression levels of tyrosine-hydroxylase (TH) in larval brains exposed to 3.0 mg/mL of CBD. There was no change in the expression level of TH. However, preliminary data suggest an increase in glial cell populations in the brainstem. In conclusion, these studies confirm that CBD is affecting CNS function and development, using a simple model system.

902A *Drosophila* larval burrowing: a parasitoid avoidance behavior? Meagan Ash, Todd Schlenke University of Arizona, Department of Entomology, Tucson, AZ

*Drosophila melanogaster* is a widely used model organism to uncover molecular mechanisms regulating natural behaviors. In the wild, fly larvae experience high rates of attack from endoparasitoid wasps. These wasps inject their eggs into the body of the insect wherein the wasp egg hatches, and the resulting wasp larva eventually consumes the host tissue. *Drosophila* larvae mount a cellular immune response against infection whereby the wasp eggs are encapsulated, melanized, and killed, which wasps attempt to suppress by injecting venoms along with their eggs. *Drosophila* larvae also perform defensive behaviors in the presence of wasps to avoid parasitism, such as moving away from wasp odors, rolling to interrupt wasp oviposition, and self-medication with ethanol to increase their toxicity to the wasp larvae inside them. Here, we describe a new larval defensive behavior in which more larvae burrow into the food substrate in the presence of wasps. We investigate the causes and adaptive value of this behavior, as well as the neurobiological mechanisms underlying it.

903B Characterization of *Drosophila* sugar receptors LINNI JIN1, Seungyun Yu2, Jae Young Kwon2, Seok Jun Moon1 1) Department of Oral Biology, BK 21 FOUR Project, Yonsei University College of Dentistry, Seoul, Korea; 2) Department of Biological Sciences, Sungkyunkwan University, South Korea
Taste allows animals to discriminate nutritious food from toxic substances. *Drosophila* gustatory receptors (GRs) for aversive chemicals are relatively well characterized; complexes consist of three or four GRs for detecting bitter chemicals. Sweet gustatory receptor neurons (GRNs) express eight closely related GRs: Gr5a, Gr61a, Gr64a, Gr64b, Gr64c, Gr64d, Gr64e, and Gr64f. However, little is known regarding the molecular basics of how sweet GRs detect sugars. Loss of function of Gr64 cluster GRs causes distinct phenotypes, presumably due to the polycistronic expression of mRNA. In addition, findings on ectopic expression of sugar receptors are contradictory to previous studies. To elucidate the function of sugar GRs, we employed the GAL4/UAS overexpression system to overexpress one or a combination of Gr64 genes in the Gr64 null deletion mutant to examine the function of the Gr64 genes. However, this approach also resulted in results inconsistent with the loss of individual Gr64 genes. To overcome the potential artifacts of overexpression of sweet GRs, we generated transgenes to express individual Gr64 cluster genes under the control of the 5′ regulatory element of the Gr5a gene, to mimic the endogenous expression of each Gr64 gene. We will present our progress on exploring the necessity and sufficiency of sweet GRs.

**904C** Meeting a threat of the Anthropocene: Robust taste avoidance of metal ions *Shuke Xiao*, Lisa Baik, John Carlson Department of MCDB, Yale University, New Haven, CT

The Anthropocene era poses a critical challenge for all organisms: they must cope with new threats at a faster rate than ever before. These threats include toxic chemical compounds released into the environment by human activities. Here, we examine high concentrations of heavy metal ions as an example of anthropogenic stressors. We find that different subsets of taste receptors contribute to avoidance behaviors towards eight metal ions when present at high concentrations that flies experienced rarely if ever until the Anthropocene. We analyze feeding and oviposition avoidance of metals at concentrations that flies experienced rarely if ever until the Anthropocene. We identify taste organs, neurons, and receptors that contribute to the avoidance of metals. We find that metals activate some taste neurons and inhibit others. Receptor mutations have different effects on different avoidance behaviors, and we identify taste organs, neurons, and receptors that contribute to the avoidance of metals. We find that metals activate some taste neurons and inhibit others. Receptor mutations have different effects on different avoidance behaviors. Some responses to metals are conserved across diverse dipteran species. Our results suggest mechanisms that may be essential to insects as they face challenges from environmental changes in the Anthropocene.

**905A** How acetic acid alters interactions of parasitoids with their *Drosophila melanogaster* hosts *Kayla Reddy*, Corinne Stouthamer², Todd Schlenke¹ 1) University of Arizona; 2) University of Georgia

*Drosophila melanogaster*, the common fruit fly, is frequently infected by parasitic wasps in nature. Flies have a cellular immune response to encapsulate and kill wasp eggs, but this can be suppressed by injected wasp venom. Flies also display defensive behaviors to prevent infection or cure themselves once infected. For example, it was previously shown that parasitized fly larvae, which live in rotting fruits, medicate themselves by consuming more alcohol. Acetic acid is another potentially toxic product of fermentation sought out by *D. melanogaster*. Here, we test whether acetic acid is beneficial to flies by affecting wasp parasitism. We performed infections on *D. melanogaster* using two parasitic wasp species, a generalist and a specialist, and measured parasitism rates in acetic acid, parasitism rates after long term exposure to acetic acid, fly and wasp tolerance of acetic acid, and fly substrate choice with various levels of acetic acid. Our data indicate that acetic acid can protect fly larvae from infection in some scenarios.

**906B** Dissecting the subcellular mechanisms of signal processing in the *Drosophila* visual system *Michelle Pang*, Thomas Clandinin Stanford University

Nervous systems process sensory information to extract relevant features from complex environments. While sensory processing algorithms are well-studied, their underlying circuit and molecular implementations remain poorly understood. How do neurons shape signal processing as they transform information from neurotransmitter input, to membrane voltage, to calcium influx, and finally to neurotransmitter release? How do these processes differ among cell types, and what molecular mechanisms drive those differences?

The Drosophila visual system is a highly tractable model for dissecting neural circuits at a subcellular level. Interneurons in the early visual system are genetically accessible, and their synaptic wiring diagram and gene expression profiles are well-characterized. Additionally, many studies have characterized their visual responses and their impact on visually-evoked behavior.

We have assembled a panel of genetically-encoded indicators of voltage, calcium, and neurotransmitter release for in vivo two-photon imaging in order to trace how visual information is transformed as it passes through neurons and across synapses. We display light and dark flash stimuli at multiple contrasts to probe the linearity and kinetics of these subcellular neural responses. Our lab previously showed that voltage and calcium sensors can be used to study visual processing with subcellular resolution, and we have now added an ultrafast calcium sensor and neurotransmitter sensors to our imaging toolkit. We are using these indicators in conjunction with pharmacological and cell type-specific genetic manipulations of ion channels and of neurotransmission to investigate how visual interneurons relay and transform information, as well as what classes of genes regulate signal processing in different neural cell types. By diversifying the types of observations we can make within neurons and synapses, we will gain novel insight into not only how Drosophila visual interneurons participate in signal processing, but also how central nervous system interneurons of different cell types function more broadly.
Alcohol research has largely been focused on understanding the neural mechanisms underlying excessive or compulsive alcohol intake. However, humans display a wide-variety of drinking behaviors with some escalating alcohol consumption over time while others moderately drink or even abstain. Investigating the circuits and neural dynamics underlying this individual variation is critical for developing more effective treatments for alcohol use and abuse disorders. In *Drosophila melanogaster*, the neural circuits required for encoding valence include identifiable connections, genetic and/or biochemical profiles and characterized temporal changes underlying learning, making flies an ideal model for investigating escalation of ethanol self-administration. We developed a 3-day operant paradigm to evaluate the spectrum of behaviors associated with self-administration of a pharmacologically relevant dose of volatilized ethanol and compared these responses to an ethologically relevant ethanol odor. Similar to mammals, individual variation in self-administration behavior for a pharmacologically relevant dose of ethanol occurs with consecutive training sessions. Approximately 35% of flies escalate self-administration whereas 61% of flies remain stable and 4% of flies decrease self-administration. This contrasts significantly with self-administration of ethologically relevant ethanol odor where 8% flies escalate self-administration. Thermogenetically inactivating a simple two neuron cholinergic and dopaminergic mushroom body circuit altered population ethanol preference to decrease and increase ethanol self-administration, respectively. Our data provides the behavioral and neuroanatomical groundwork to subsequently investigate variability in physiology of identified circuits contributing to alcohol use disorder.

All animals must process and respond accordingly to both internal and external stimuli. Modifying behavior is an essential strategy animals employ to respond to a changing environment. Octopamine (OA)—the insect equivalent of norepinephrine—functions as a neurotransmitter, neurohormone, and neuromodulator involved in modulating various behaviors based on sensory input and internal states. The relationship between OA and aggression has been well studied in insects and other arthropods. Because of its expansive genetic toolkit, Drosophila is one of the most useful model organisms for exploring the mechanisms employed by OA neurons to influence aggression. Past studies using Drosophila have demonstrated OA’s role in modulating aggression based on various sensory inputs, such as sleep, gut microbiome health, and social experience. Importantly, OA neurons in Drosophila are known to consist of up to 27 anatomically distinct subtypes. Aside from a few examples, it remains in question whether each subtype modulates a specific behavioral process (including context-dependent modulation of aggressive behavior), and if so, which of these OA neurons is responsible for each process. Genetic reagents (such as GAL4 lines) that label specific subtypes of OA neurons will help advance our understanding of the circuits driving octopaminergic modulation of behavior. Currently, only a few OA subtypes are genetically accessible. To investigate the functions of the remaining OA neuronal subtypes, we are isolating split GAL4 lines that label specific subtypes of OA neurons in the central brain. After using immunohistochemistry to visualize the labeling patterns of 367 combinations of split GAL4 Drosophila lines, we have so far found 24 candidates that partially or completely isolate several octopaminergic cell types. Our new genetic reagents will enable further investigation of not only OA’s role in aggression circuitry but also cell-type specific control of other behaviors modulated by OA. With more refined access to the circuits underpinning behavior modulation, we can achieve a better understanding of how animals process and respond to the barrage of stimuli they constantly encounter.

Amineergic neurons modulate cellular mechanisms involved in almost every human behavior. Norepinephrine in particular has been shown to modulate physiological processes ranging from heart rate to ovulation. Work from our lab and others has shown that the *Drosophila* equivalent to norepinephrine, octopamine (OA), also has a role in fly ovulation. Processes such as follicle cell rupture in the ovaries, muscle contraction in the oviduct, and sperm storage in the spermathecae have all been shown to be regulated by OA signaling. The specific neurons, receptors, and cellular mechanisms underlying such signaling, however, remain mostly unknown. OA signals to the *Drosophila* reproductive system from a central cluster of OA neurons in the abdominal ganglion, and both alpha and beta types of OA receptors, which are analogous to the alpha and beta types of mammalian adrenergic receptors, are expressed throughout the reproductive organs. Our lab has developed a set of novel preparations and assays for the optogenetic interrogation of this egg laying circuit. We show that stimulation of the OA cluster innervating the reproductive system produces rhythmic contractions of the lateral oviduct muscle. Our work also suggests that the circuit pathway facilitating this action is indirect, including OA signaling to other intermediate neurons rather than directly to the oviduct muscle, which lacks OA receptor expression. This indirect pathway includes a small group of neurons that express insulin-like peptide 7 (ILP7) and a variety of different neurotransmitters/peptides. OA-driven contractions of the lateral oviducts may therefore incorporate contributions from multiple different types of neurotransmission. By mapping the coexpression of other neurotransmitters and receptors in this circuit, such as glutamate and proctolin, and comparing optogenetically-
driven behaviors to those of single-neurotransmitter/peptide perfusions, we aim to define the specific presynaptic and postsynaptic components of this indirect pathway in the OA egg laying circuit. We hypothesize that OA signaling in the egg laying circuit has multiple distinct roles governed by both direct and indirect pathways as well as differences in which OA receptors are expressed in each pathway. This work will help establish a model that can be used to uncover how aminergic neurons coordinate within a single central cluster to regulate multiple different behaviors.

910C  Parallel processing of polarized skylight from the optic lobes towards the central brain Juliane Uhlhorn1, Emil Kind2, Gizem Sancer2, Thomas Mathejczyk3, Mathias Wernet1 1) Freie Universitaet Berlin, Berlin; 2) Yale University, New Haven, CT

Many animals rely on polarized UV skylight for orientation and navigation. The underlying neuronal circuitry for mediating this behavior in many insects, including Drosophila melanogaster, is formed by the Anterior Visual Pathway (AVP). Specialized photoreceptors in the dorsal periphery of the eye, the so-called dorsal rim area (DRA), detect polarized skylight. Additional cell types then process the information from within the optic lobes, before conveying it to a visual glomerulus – the anterior optic tubercle (AOTU) - towards the central complex. The retinotopic information encoded by the angle of polarization (AoP) is conserved throughout this pathway.

Of particular importance are visual projection neurons connecting the medulla neuropil to the central brain. To understand these parallel pathways, we characterize the morphological similarities and/or differences between subtypes using light microscopic techniques such as MultiColor FlpOut (MCFO). Additionally, using GRASP and connectomic EM reconstruction, we investigate potential differences in connectivity. Finally, functional approaches like calcium imaging and quantitative behavior assays are used. Although the overall retinotopic representation is conserved in most subtypes, we find distinct functional differences between them. Hence, simple redundant roles for these parallel channels seem unlikely. We propose a role for these cell types in processing specific navigational cues.

911A  Characterization of the mode of transmission of ethanol resistance to progeny of repeatedly intoxicated parental flies Michelle Bonilla, Jocelyn Coreas, Merna Massoud, Antony Ibrahim, Mariano Loza-Coll  California State University, Northridge, Los Angeles, CA

Characterization of the mode of transmission of ethanol resistance to progeny of repeatedly intoxicated parental fliesThere has been a renewed interest in the transmission of acquired traits, particularly those related to tolerance to drugs and environmental toxins. Since tolerance is an accepted pre-requisite and strong predictor of addiction, a firmer understanding of the mechanisms underlying the transmission of tolerance could facilitate novel research avenues of importance in public health. Despite the significant advances made in characterizing the epigenetic and molecular mechanisms that underlie the transmission of acquired traits across diverse species, it is still unclear what genetic pathways may connect acquired traits that operate primarily in metabolic, physiologic or nervous system levels with the necessary epigenetic modifications in the germline for transmission to take place. While classic model organisms, such D. melanogaster and C. elegans are prime candidates to dissect conserved aspects of the genetic and epigenetic underpinnings of acquired trait inheritance, little is known about their capacity to transmit acquired drug tolerance to their progeny.

Numerous previous studies over the last 20 years have demonstrated that fruit flies can develop tolerance to ethanol when repeatedly exposed to the drug. We have recently reported that parental flies that are intoxicated multiple times (once a day, for 10 minutes, over a 2 weeks period) give rise to progeny that is significantly more resistant to the sedative effects of ethanol. We observed that parental flies need to be exposed multiple times before transmission of resistance to progeny can be observed, that there is residual transmission of tolerance to the F2 and that the ability of females to transmit tolerance to their F1 progeny lasts several days after their last intoxicating exposure to ethanol. We have since more carefully explored whether the transmission of resistance to progeny of repeatedly intoxicated flies is matrilineal, patrilineal or both, which will delineate future research efforts aimed at elucidating the genetic and molecular bases of the transmission. Overall, our findings present D. melanogaster as a suitable experimental model for forward and reverse genetics approaches to investigate the pathways connecting repeated exposures of an adult animal to ethanol and the necessary germline modifications that underlie the transmission of increased resistance in their progeny to the drug.

912V  Pre-copulatory reproductive behaviours are preserved in Drosophila melanogaster infected with bacteria. Saloni Rose1, Esteban Beckwith1,3, Charlotte Burmester1, Robin May1, Marc Dionne2, Carolina Rezaval1 1) University of Birmingham; 2) MRC Centre for Molecular Bacteriology and Infection and Department of Life Sciences, Imperial College London, London SW7 2AZ, United Kingdom; 3) Instituto de Fisiología, Biología Molecular y Neurociencias (IFIBYNE), UBA-CONICET, Buenos Aires, Argentina; 4) Institute of Microbiology & Infection, University of Birmingham, Birmingham B15 2TT, United Kingdom

The activation of the immune system upon infection exerts a huge energetic demand on an individual, decreasing available resources for other vital processes, such as reproduction. However, the factors that determine the trade-off between defensive and reproductive traits remain poorly understood. Here, we exploit the experimental malleability of
the fruit fly Drosophila melanogaster to systematically assess the impact of immune system activation on pre-copulatory reproductive behaviour. Contrary to our expectations, we found that male flies undergoing an immune activation continue to display high levels of courtship and mating success. Similarly, immune-challenged female flies remain highly sexually receptive. By combining behavioural paradigms, a diverse panel of pathogens and genetic strategies to induce both arms of the fly immune system, we show that pre-copulatory reproductive behaviours are preserved in infected flies, despite the huge metabolic cost of infection.

**913V** Identification of individual essential amino acid sensors in *Drosophila* Jinhyeong Lee, Jong-Hoon Won, Boram Kim, Greg Suh KAIST

A balanced intake of macronutrients such as proteins, carbohydrates and fats is essential for the organism, and insufficient protein intake leads to several diseases, including kwashiorkor. Protein-deficient animals have a dietary behavior which is selecting food sources that contain essential amino acids (EAAs) rather than non-essential amino acids. Recently, a neuropeptide called CNMamide (CNMa) was found in *Drosophila melanogaster* to be highly expressed in enterocytes of the midgut during protein depletion to modulate EAA-specific selective dietary behavior. CNMa expression in these enterocytes is up-regulated in response to protein depletion through the GCN2-ATF4 and TOR-MITF pathways, each of which trigger a compensatory appetite for EAA (Kim, et al. 2021, *Nature*). Remarkably, flies fed a holidic diet lacking a single L-EAA had enhanced CNMa expression, suggesting the presence of a molecular sensor that recognizes each L-EAA. We have carried out RNAi screening to identify candidate molecules that sense or detect each EAA and are currently characterizing these putative sensors using molecular, genetic and biochemical approaches.


**914V** Neuroligin3 and dopamine are required for a response to social isolation, but recovery is complex and sex-specific. Ryley T Yost, Branden Walshe-Roussel, Anne F Simon University of Western Ontario

Social isolation causes profound changes in social behaviour in a variety of species including humans, monkeys, mice, bees, and vinegar flies. However, the genetic and molecular mechanisms modulating behavioural responses to both social isolation and social recovery remain to be elucidated. Here, we quantified the behavioural response of vinegar flies to social isolation using the flies’ social space preference. Flies with a loss of function of *neuroligin3* (ortholog of autism-related *neuroligin* genes) with known increased social space in socially enriched environment, responded similarly to genetically matched control to both social isolation and recovery. We next used a UAS-TH-RNAi driver in all neurons to investigate the role of dopamine, another known modulator of a response to social isolation. We show that dopamine is important for a response to social isolation and recovery in males but not in females. Furthermore, only in males, dopamine levels are reduced after isolation and are not recovered after group housing. Finally, in socially enriched flies with a loss of function of *neuroligin3*, dopamine levels are reduced in males, but not in females. We will share a model to explain how dopamine and *neuroligin3* are involved in the behavioural response to social isolation and its recovery in a dynamic and sex-specific manner.

**915V** Behavioral Characterization of *tecu* Mutants Laura Alejandra Lujano Perez¹, Juan Rafael Riesgo Escovar² 1) Maestría en Ciencias (Neurobiología), Universidad Nacional Autónoma de México; 2) Instituto de Neurobiología, Universidad Nacional Autónoma de México

We isolated mutations in a gene that we named *teczitécatl* (*tecu*), that codes for a phospholipase A₂ enzyme narrowly expressed in a group of Kenyon cells, which are the mushroom bodies intrinsic neurons in the brain of *Drosophila melanogaster*. Here we show that mutations in this gene lead to faulty behavioral responses in visual paradigms. We directly compared responses of control (yw) and two mutant *tecu* strains (*tecu¹* and *tecu²*), because all of them share the same genetic background. We first performed a larval phototaxis assay. Results show that they present significantly different responses (migrate more towards the light). We also performed an adult countercurrent assay, where they have a deficient response to light (have significantly less positive phototaxis). Taken together, these results show that *tecu* has an important role processing visual responses in *Drosophila melanogaster*.

**916V** Local 5-HT signals bi-directionally modulate the coincidence time window of associative learning Xuelin Li, Jianzhi Zeng, Zimo Zhangren, Yulong Li Peking University, School of Life Sciences

Temporal coincidence between a conditioned stimulus (CS) and an unconditioned stimulus (US) is an essential component in Pavlovian associative learning. Despite its ubiquitous presence, it remains unknown whether the coincidence time-window can be regulated. Here we assign this function in Drosophila to the dorsal paired medial (DPM) neuron, which innervates the mushroom body (MB) and releases 5-HT compartmentally in response to both an odorant (the CS) and electric shock (the US). The DPM neuron is activated by local acetylcholine (ACh) release from MB

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Kenyon cells through nicotinic ACh receptors, meanwhile releases 5-HT to reduce ACh release via the 5-HT1a receptor. Suppressing or promoting the 5-HT release from DPMs bi-directionally regulates the coincidence time-window for odor-shock pairing, which gates olfactory learning and synaptic plasticity changes in the MB. This mechanism provides a model for studying how an organism times the temporal continuity of environmental events and learns the underlying causal relationship.

917V Spying on the dynamics of octopamine by genetically-encoded GRAB _OA_ sensor in _Drosophila_ Mingyue _Lv_1,2, Ruyi _Cai_1,2, Huan _Wang_1,2, Yulong _Li_1,2,3,4 1) State Key Laboratory of Membrane Biology, Peking University School of Life Sciences, Beijing, China; 2) PKU-IDG/McGovern Institute for Brain Research, Beijing, China; 3) Peking-Tsinghua Center for Life Sciences, Academy for Advanced Interdisciplinary Studies, Peking University, Beijing, China; 4) Chinese Institute for Brain Research, Beijing, China

Small molecular neurotransmitters are essential for neuronal communication in the nervous system. Octopamine (OA), the analog of norepinephrine in invertebrates, is an important monoamine neurotransmitter involved in many physiological events of insects, such as the aggression, flight, ovulation, learning and memory. Although much is known regarding the properties of OA, its spatial and temporal dynamics are poorly understood at the in vivo level due to the limited detecting methods. Here, to directly visualize the release of OA, we developed a genetically encoded G-protein-coupled receptor (GPCR) – activation – based OA (GRAB _OA_ ) sensor. In response to extracellular OA, GRAB _OA_ shows large fluorescent increase (~500% DF/F0), high selectivity and fast kinetics to micromolar OA. In living flies, we observed GRAB _OA_ expressed in the Kenyon cells (KCs) responses to electrical stimulation in a dose dependent pattern. For optogenetic activation of octopaminergic neurons, we detected homogenous fluorescent increase across mushroom body, which is the center for _Drosophila_ olfactory associative learning. Moreover, GRAB _OA_ was able to reports physiological stimulation including odorant and abdomen shock induced OA release. Thus, GRAB _OA_ enables spatiotemporally precise measurements of OA dynamics in vivo in _Drosophila_, which provides the possibility to explore the role of OA in different interesting physiological process in invertebrates.

918V Two Individually Identified Paired Dopamine Neurons Signal Taste Punishment in Larval _Drosophila_ Denise _Weber_1, Katrin _Vogt_1, Andreas _Thum_1 1) Institute of Biology; Department of Genetics, University of Leipzig, Germany; 2) Department of Neurobiology, University of Konstanz, Germany

Dopaminergic neurons (DANs) perform multiple tasks in the brain. Among other things, DANs mediate teaching signals such as information about rewards and punishments throughout the animal kingdom. They serve to evaluate sensory input, store the resulting associations as memory, and continuously update them according to relevance and reliability. Therefore, to better understand the functioning of the dopaminergic system, it is crucial to know what specific roles are mediated by each DAN. To this end, we are studying _Drosophila_ larvae, whose brains consist of only about 12,000 neurons, of which only about 1% are DANs.

Only eight larval DANs presynaptically project to the mushroom body, a brain region in insects that is of central importance to associative olfactory learning. These eight DANs in turn anatomically subdivide into four cells of the primary protocerebral anterior medial cluster (pPAM) and four cells of the dorsolateral 1 cluster (DL1). As we have shown in previous studies, the activity of the pPAM DANs encodes the internal gustatory sugar reward signal. In this study, we investigate the four DANs of the DL1 cluster in terms of their cellular function with respect to gustatory teaching signals. We discover that two DANs (DAN-f1 and DAN-g1) innervating two specific compartments of the vertical lobe of the mushroom body are in combination acutely necessary for learning of odor-high salt punishment, but dispensable for appetitive learning and innate behavior toward the applied odors and salt. Optogenetical activation of DAN-f1 and DAN-g1 neurons in the larval brain is sufficient to encode punishment. The functional specificity of DAN-f1 and DAN-g1 within the DL1 cluster and towards the PAM cluster is further confirmed by synaptic reconstruction of sensory and DAN input neurons. Thus, the DL1 neurons convey a punishment teaching signal.

In summary, this indicates that a cellular division of labor of larval DANs exists with respect to the transmission of dopaminergic reward (pPAM cluster) and punishment (DL1 cluster) signals. Since the organizing principle of larval DANs is the same as that of its adult counterpart and that of the mammalian basal ganglion, it is possible that there are only a limited number of efficient neural circuit solutions to solve complex cognitive and behavioral problems in nature.

919V Single cell transcriptomic analysis of homologous courtship song neurons between species Justin _Walsh_1, Johaer _Jilani_1, Lihua _Wang_2, Yun _Ding_1 1) University of Pennsylvania; 2) Janelia Research Campus of the Howard Hughes Medical Institute

Understanding the evolution of complex traits, including behavior, is a main goal of evolutionary biology. Behavioral diversity across species is expected to be the result of evolutionary changes in neural morphology and function. In turn, a neuron’s morphology and function is the product of its molecular makeup. Therefore, comparing gene expression differences in homologous neurons between species is a powerful approach to identify genes that contribute to the
evolution of neural differences underlying behavioral diversity. *Drosophila* courtship song, wing vibrations generated by a male while courting a female, displays a wide range of diversity across species. For example, *D. melanogaster* males sing two types of song, pulse and sine, by extending and vibrating one wing. In contrast, *D. yakuba* males do not sing sine song but instead sing a song called clack by vibrating both wings without obvious extension. A male-specific single bilateral pair of neurons named pIP10 is necessary for pulse song in *D. melanogaster* but clack song in *D. yakuba*. In both species, we genetically labeled pIP10 neurons with a GFP marker and manually isolated them for transcriptomic profiling. We found that many genes are expressed in a species-specific or -biased manner including some with known neural or behavioral functions. Next, in *D. melanogaster* we used pIP10-specific RNAi to knock down genes upregulated in *D. melanogaster* pIP10 neurons relative to *D. yakuba* pIP10 neurons and found that some of our knockdowns resulted in mutant *D. melanogaster* song phenotypes. Overall, our study is one of the first to compare gene expression in homologous neurons across species and offers a novel perspective on how evolution has shaped gene expression patterns to result in functional adaptation across species.

920V Investigating the Role of SIFamide in the Effects of Food Deprivation on Female Reproductive Drive *Attilio Ceretti*, Jill Schneider Lehigh University

Adaptation requires differential reproductive success, but reproductive processes are energetically expensive and can compromise the chances of survival under energetic challenges. Thus, many species appear to have evolved mechanisms that inhibit reproduction when energy is scarce. For example, in a genetically heterogenous population of *Drosophila melanogaster* derived from an orchard in New Jersey, we found that 36-48 hours of food deprivation in females significantly decreased the incidence of copulation with fed males, whereas food deprivation in males had no effect on courtship or copulation attempts with fed females. One signal that could be mediating these results is the evolutionarily conserved neuropeptide, SIFamide, the insect ortholog of the mammalian peptidergic/hormonal signal RFamide-related peptide-3, which mediates the effects of energy on reproduction in hamsters. To test whether this peptide mediates the effects of energy on reproduction, RNAi towards SIFamide was expressed in SIFamide neurons. Wildtype control females decrease their mating rate and increase their latency to copulate following only 24 hours of food deprivation, while females that express RNAi toward SIFamide do not. Knockdown of SIFamide blocks the effects of food deprivation on female reproduction, consistent with the idea that SIFamide expression is important for the inhibition of reproduction when energy is scarce. Our results provide a potential neurological basis for an ancestral system that mediates tradeoffs in food intake and reproduction.

921V Chronic caffeine treatment disrupts circadian rhythm in *Drosophila* *Aishwarya Segu*, Nisha Kannan Indian Institute of Science Education and Research, Thiruvananthapuram

The circadian clock governs the timing of sleep-wake cycles as well as other behavioural, physiological and metabolic processes. While the endogenous circadian clock mediates the timing of sleep, homeostatic mechanisms modulate the amount and depth of sleep. Evidence from previous studies showed that caffeine intake promotes wakefulness, whereas adult-stage specific caffeine treatment not only suppresses sleep but also delays the phase of circadian rhythm in Drosophila. In humans, caffeine is consumed on a daily basis and hence it is important to understand the effect of prolonged intake of caffeine on circadian and homeostatic regulation of sleep. In the present study we examined the differential effect of acute and chronic caffeine treatment on sleep ontogeny as well as on circadian and homeostatic regulation of sleep in *Drosophila*. The results of our study showed that acute caffeine treatment reduces both day and night sleep in mature flies whereas it reduced only the day sleep in young flies. Chronic caffeine treatment did not exert any significant effect on sleep in young flies. On the other hand, it delayed the timing of sleep in mature flies and in addition these flies reduced the morning and evening anticipatory activity at higher caffeine concentration under 12 hour: 12 hour light: dark cycles. Apart from these it also changed the overt rhythm of these flies. These flies exhibited either a longer free running period or arrhythmicity under constant darkness. The results from our study shows that acute treatment of caffeine reduces sleep in flies and does not affect the homeostatic sleep whereas prolonged caffeine treatment affects the overt rhythm of the flies including the eclosion rhythm.

922V Aggression in *Hieroglyphus banian* (Rice grasshopper) vs. in *Drosophila melanogaster*: A Comparison *Abhilash Kondai*, Sudipta Saraswati University of Hyderabad, Hyderabad, India

While aggression has been reported to be a widely observed behavior across animal species, it has not been systematically studied in many organisms. For example, aggressive behavior in *Drosophila melanogaster* has been well-documented but that is not the case with grasshoppers. We undertook an extensive study of aggressive behavior in a species of rice grasshopper, *Hieroglyphus banian*. Here, we present our findings of aggression in male *Hieroglyphus banian* and compare our findings with published literature of aggression in male *Drosophila melanogaster*. We found that several components of aggressive behavior, such as (i) approach, (ii) holding or grabbing, (iii) low-level fencing or step on, (iv) tussling, and (v) walking away or running away or flying away, are conserved between grasshopper and *Drosophila*. However, we also observed that biting is an aggressive behavior that is exhibited by only grasshopper and is not reported in *Drosophila*. Interestingly, in grasshopper we did not observe some behavioral components of
agression, such as (i) wing threatening, (ii) lunging, and (iii) boxing, that are reported to be exhibited by Drosophila. Our findings and comparative analysis indicate that several components of aggressive behavior are conserved between two insect species of two different kinds—hemimetabolous insect grasshopper and holometabolous insect Drosophila. Results from our study will help us further investigate if the aforementioned conserved aspects of aggressive behavior are conserved across a variety of insect species. Moreover, our study also forms the basis for investigations into the functional significance of the conserved vs. non-conserved aspects of aggressive behavior across different insect species.

923V  Intestinal CNMα induction by protein deficit affects two distinct pathways in the brain to regulate the preference for protein-rich food  Boraem Kim1, Seongju Lee1, Yangkyun Oh2, Makoto Kanai2, Won-Jae Lee3, Greg S. B. Suh1 1) Department of Biological Sciences, Korea Advanced Institute of Science and Technology, Daejeon, Republic of Korea; 2) Skirball Institute of Biomolecular Medicine, Department of Cell Biology, New York University Grossman School of Medicine, New York, NY, USA; 3) National Creative Research Initiative Center for Hologenomics and School of Biological Sciences, Seoul National University, Seoul, Republic of Korea

The ability to maintain protein homeostasis is crucial for survival and fitness. While it is known that the taste sensory system recognizes the availability of dietary proteins in the environment, it remains unclear how animal detects and responds to the deficit of amino acid levels in the internal milieu. Essential amino acids (EAAs) cannot be synthesized and must be consumed through diet. CNMamide (CNMa) is released from a specific population of enterocytes in the Drosophila gut during EAAs deprivation and causes increased appetite for EAAs (Kim et al., Nature 2021). However, the mechanism by which CNMα influences the brain for the promotion of EAAs consumption remains poorly understood. Here, we show that among many regions of the brain that express CNMα receptors, two distinct regions of the brain that affect protein and carbohydrate appetite are activated or inhibited by CNMα. These findings reveal the interplay between the gut and the brain for the regulation of nutrient-specific appetite.


924V  Molecular and cellular basis of acid taste sensation in Drosophila Ting-Wei Mi1, John Mack1, Christopher Lee2, Yali Zhang1,1 1) Monell Chemical Senses Center, Philadelphia, PA; 2) Department of Biology, University of Pennsylvania, Philadelphia, PA; 3) Department of Physiology, The Diabetes Research Center, University of Pennsylvania, Perelman School of Medicine, Philadelphia, PA

Sour taste is one of the five commonly accepted basic taste modalities, alongside four other fundamental taste: bitter, sweet, salty, and umami. This taste sensation is evoked by acidic foods such as vinegar or certain fruits, and allows organisms to avoid consuming unripe or spoiled, or fermented foods. Species from insects to mammals are attracted by slightly acidic foods, but repulsed to highly acidic foods. However, the molecular and cellular mechanism of how animals distinguish low from high acid foods is still largely unknown. Here, we use the fruit fly (Drosophila melanogaster) as a model organism to address this question. Firstly, we show that fruit flies employ two competing taste sensory pathways to discriminate low from high acidicity, which trigger attractive or aversive behavior respectively. Secondly, we identify Otopetrin-like α (OtopLa), a member of an evolutionarily conserved gene family, as a proton channel in fruit flies expressed in bipolar type gustatory receptor neurons (GRNs) in the proboscis with central projection patterns in the subesophageal zone (SEZ), which receives taste input from the peripheral taste organs. Loss of otopla impairs flies’ acid sensation, causing an imbalance in the competing taste pathways leading to an abnormal aversive response to low acid. In summary, our behavioral, anatomical, and electrophysiological data strongly suggest that flies display a competing acid-sensing mechanism to distinguish between high and low acid, and the proton channel OtopLa is required for low acid sensation. This sensing mechanism that may be evolutionarily conserved between insects and mammals, given the similarity in their taste preferences for acid and homologous functions of Otop genes.

925V  Gastric mechanosensation and the peptidergic sugar sensing regulate the Drosophila nutrient sensor Yangkyun Oh1, Jason Lai1, Soohong Min2, Huai-Wei Huang1, Stephen Liberles2, Hyung Don Ryoo1, Greg Suh1,3 1) Department of Cell Biology, NYU Grossman School of Medicine, New York, NY, 10016; 2) Harvard Medical School, Howard Hugh Medical Institute, Department of Cell Biology, Boston, MA 02115; 3) Department of Biological Sciences, Korea Advanced Institute of Science and Technology, Daejeon 34141, Republic of Korea

Nutrient sensors allow animals to identify foods rich in specific nutrients. The Drosophila nutrient sensor, DH44 neurons, helps the fly to detect nutritive sugar. This sensor becomes operational during starvation; however, the mechanisms by which DH44 neurons or other nutrient sensors are regulated remain unclear. Here, we identified two satiety signals that inhibit DH44 neurons: 1) Piezo-mediated stomach/crop stretch after food ingestion; 2) Neuromedin/Hugin neuropeptide in the ventral nerve cord (VNC) activated by an increase in the internal glucose levels. A subset of Piezo+ neurons that express DH44 neuropeptide project to the crop. We found that DH44 neuronal activity and food intake were stimulated following a knockdown of piezo in DH44 neurons or silencing of the VNC Hugin neurons, even in fed flies. Together, we propose that these two qualitatively distinct peripheral signals work in concert to regulate the
The maintenance of synapses is dependent on several coordinated mechanisms, which involve cell-to-cell communication, molecules recycling and degradation. However, the mechanism that controls the synapse maintenance during aging is poorly understood. Here, we investigated the synapse maintenance at the neuromuscular junction (NMJ) of adult Drosophila melanogaster lines. Using a flight behavior tester, we evaluated the flight ability of 205 inbred fly lines from the Drosophila Genetic Reference Panel (DGRP), which harbor known SNP and non-SNP variants in or near genes, during aging. To flight test, we collected flies, which were reared on an incubator at 25°C for 1- or 2-days post pupae eclosion, separated by sex and matured them for 3 days and 21 days on an incubator at 29°C. We observed a widely variable flight ability in the DGRP lines from day 3 to day 21 during aging. However, 10 out of 205 DGRP lines had the most expressive decline of the flight ability during aging. To validate these findings, we tested mutant flies for these 10 genes and observed that these genes are prone to cause progressive loss of flight ability. In addition, to identify how these genes cause disruption on tripartite synapse of the neuromuscular junction (NMJ), we used a UAS-Gal4 system to direct the knockdown of those genes on glia (Repo-Gal4), muscle (MHC-Gal4) and motor neuron (BG380-Gal4). Knocking down the 10 genes on muscle and/or motor neurons led to flight ability decline, but one out of the 10 genes caused a progressive defect of the flight behavior directed by the glial driver. To further understand the cellular alterations on synapse at the NMJ, we are characterizing the morphology of the pre- and post-synaptic components. Our data is highlighting the mechanism regulating the synapse maintenance during aging and the progressive degeneration of the synapse cellular components.

Cilia are involved in a plethora of motility- and sensory-related functions. Ciliary defects cause several ciliopathies, some of which with late onset, suggesting cilia are actively maintained. While much is known about cilia assembly, little is understood about the mechanisms of their maintenance. Given that intraflagellar transport (IFT) is essential for cillum assembly, we investigated the role of one of its main players, IFT88, in ciliary maintenance. We show that DmIFT88/NOMPB, the Drosophila melanogaster orthologue of IFT88, continues to move along fully formed sensory cilia. We further identify the Drosophila Guanylyl Cyclase 2d (DmGucy2d/CG34357) and the TRPV channel subunit -inactive- as DmIFT88 cargoes. In particular, DmIFT88 binds and is important for the localisation of the intracellular part of DmGucy2d, which is evolutionarily conserved and mutated in several degenerative retina diseases. We find that acute knockdown of DmIFT88 and DmGucy2d in ciliated neurons of adult flies leads to defects in the maintenance of cilium function, impairing hearing, and gravitaxis behaviour.

Our results show that cilia function requires an active maintenance program and that DmIFT88, and two cargos, are important for this process. In addition, our work suggests that this maintenance program may be deregulated in degenerative human diseases where Guanylyl Cyclases are mutated.

Bark beetles (Coleoptera: Curculionidae: Scolytinae) are a highly diverse subfamily of weevils generally considered as economically highly relevant pests of forests. As a natural part of forest ecosystems, they rely on damaged trees, but can switch to healthy hosts during the ‘aggressive’ outbreak phase (Biedermann et al., 2019). Bark beetles employ species-specific aggregation pheromones to attract conspecifics to their host trees (predominantly conifers) for the purposes of mating and resource exploitation. Within the Scolytinae, a clear phylogenetic pattern has been observed in pheromone composition; some components are phylogenetically more conserved than others (Symonds and Elgar, 2004; Symonds and Gitau-Clarke, 2016). The differences in pheromone composition are greater between closely related species than with distantly related species. This indicates a rapid ‘switching on and switching off’ of certain compounds in aggregation pheromone composition (Symonds and Elgar, 2008, 2004). From the evolutionary perspective, we hypothesize a similar level of rapid and heritable switches in olfactory reception at molecular level, i.e. in the structure and expression of pheromone-detecting receptors. We therefore investigate the evolution of chemosensory gene families in closely related Ips species. Species were selected based on niche level differentiation (syntropy), feeding modes and availability in the Czech Republic (Lindgren and Raffa, 2013). An antennal transcriptome and, more recently, the genome of I. typographus have become available (Andersson et al., 2013; Powell et al., 2021) and our research extends this knowledge to other species using both transcriptomes and genomes. For functional analysis we
use a *Drosophila* transgenic expression system (so called “empty neuron”) in conjunction with electrophysiological measurements to characterize receptors by responses (Gonzalez et al., 2016). Overall, this research will provide better understanding of olfactory perception in bark beetles and thereby support development of olfaction-based pest-control strategies.

**929V Genetic dissection of physiological properties of local interneurons in the *Drosophila* larval visual circuit**

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Animals continually modify their behaviors in response to the changing environment. Understanding the molecular and circuit basis of how the nervous system modifies its function to adjust behavioral output is an active topic in the neurobiology field. The *Drosophila* larval visual circuit serves as a suitable system to study this fundamental question. Composed of only twelve photoreceptors (PRs) to detect visual stimuli, one pair of visual local interneurons (VLNs) to modulate the signals, and nine visual projection neurons (VPNs) to transfer the filtered information to higher brain regions, larvae employ a simple visual circuit for a variety of behaviors. Specifically, recent connectome and functional imaging analyses revealed these VLNIs, which have been identified as one cholinergic (cha-IOLP) and one glutamatergic (glu-IOLP), synaptically interact with each other as well as receive neuromodulatory input from serotoninergic and octopaminergic neurons, suggesting their potential role as plastic components in the circuit for internal or external signals to modulate visual information and impact behavior. Our previous study suggests that excitatory and inhibitory inputs into the VLNIs are mediated by glutamatergic and cholinergic signaling. To better understand how visual signals are processed and modulated at the level of VLNIs, we aim to evaluate the interactions between these two antagonizing pathways within larval VLNIs. First, to identify the neurotransmitter receptors expressed in each VLN, we performed a comprehensive survey on acetylcholine receptors (AChRs) and glutamate receptors (GluRs) using endogenously-tagged GAL4 and LexA lines. Our preliminary results offered an overview of the combinatorial expression of nicotinic AChR (nAChR) subunits and muscarinic AChRs (mAChRs) as well as that of ionotropic and metabotropic GluRs in larval VLNIs. Second, to validate the screen results, we are performing immunostaining and genetic manipulations, in combination with physiological and behavioral tests. Our results will provide molecular insights into how sensory information is computed and transmitted within a sensor sensory circuit and help us identify specific cellular and molecular components targeted by developmental and activity-dependent regulatory mechanisms.

**930V Manipulation of neuron transmission in the mushroom bodies and protocerebral bridge affects social behaviour**

Abigail Bechard, J. Wesley Robinson, Ryley Yost, Anne Simon Western University, London ON, Canada

Social interactions between animals depend upon where they settle in relation to each other; a behaviour called social spacing. This behaviour is defined as the distance between individuals in a stable group and has been shown to be affected by life experience, genetics, and the environment. Simple interactions like social spacing that mediate and precede more complex behaviours serve as a practical means to elucidate the basic neurogenetics involved in a variety of social behaviours.

Social behaviours are determined by perceiving attractive and repulsive cues from other individuals, followed by the neural integration of these cues. A key aspect in the determination of an organism’s behaviour (including social spacing) is how these cues are integrated in the brain. For example, in *D. melanogaster* the Mushroom Bodies (MB) have been demonstrated to be involved in specific behaviours like temperature preference, social approach, and food-seeking, as well as learning, and memory.

Here, we are interested in elucidating the neural circuitry which modulates social spacing at the level of brain structures. We are currently investigating whether two brain structures, the Mushroom Bodies and the Protocerebral Bridge (PB), play a role in social spacing neural circuitry. These structures are of interest because they show enrichment of Neuroligin-3 (Nlg-3), a postsynaptic cell adhesion protein that we previously found to have a strong association with social spacing. This behaviour is defined as the distance between individuals in a stable group and has been shown to be affected by life experience, genetics, or differential signalling of neural circuitry. So far, we and others have determined that the mushroom bodies (MB) of the *Drosophila* brain are important for social space. As well, synaptic proteins including Rugose a scaffolding protein at the postsynaptic density, Narrow-
abdomen a channel at the presynaptic terminal, and proteins involved with dopamine synthesis and vesicle loading all play a role in social space - highlighting the importance of synaptic functioning on this behaviour. The manipulation of global dopaminergic signalling also alters social space in a sexually dimorphic manner; however, we do not know how dopamine signalling, synaptic proteins, and the MB are all connected to modify fly social spacing. Here, we show that a deficiency of neuroligin 3 (nlg3), a gene encoding a post-synaptic protein that regulates transmission at the synapse, modifies social space. Neuroligin 3 is also found to be enriched in the MB, protocerebral bridge, and optic lobes regions of the brain. Hyperactivation and silencing of fruitless neurons recapitulates the sexually dimorphic effect we previously observed in fly social spacing, so lastly we targeted a dopaminergic knockdown within the P1 subset of fruitless neurons. Our results contribute to understanding the role of specific neural circuitry in modulating the social spacing behaviour of Drosophila.

932V  Don’t want to be all by myself BUT Don’t stand so close to me Anne F Simon, Ryley T Yost, J Wes Robinson, Abigail T Bechard University of Western Ontario

Before engaging in complex social interactions, flies must determine their preferred social space. Indeed, all motile organisms, from bacteria to humans, including D. melanogaster, display a preferred inter-individual (or social) distance that can be affected by genetics, experience and/or the environment.

One of the focus of my lab, is to better understand the neurogenetic underpinnings of social space determination. We and others have shown that social spacing in D. melanogaster can be influenced by a variety of intrinsic and extrinsic factors, such as mating status, social enrichment, genes, and environmental conditions, and an interplay between those.

A sex specific neural circuit is emerging as a modulator of social spacing: it involves dopaminergic signalling, and two major brain structures: the mushroom bodies, and protocerebral bridge. In addition, the neural bases of social spacing are starting to be elucidated. And several of the players – from neurotransmitters to post-synaptic proteins – are conserved throughout evolution. At the synaptic level, we have shown that Neurobeachin (an anchor protein) and Neuroligin (a cell adhesion protein) are implicated in social space, as well. Both of those postsynaptic proteins have human homologues that are candidate genes for Autism Spectrum Disorders (ASDs).

In another type of response to social cues, flies strongly avoid the volatile substance dSO emitted by stressed flies. CO2 has been identified as one of the compounds in dSO, although other unidentified compound(s) are required to elicit the full avoidance of dSO. Because dSO is emitted by stress flies and causes a response from conspecifics, it is considered a social cue. Compared to social space, dSO avoidance requires different sensory modalities such as olfaction.

Surprisingly, we found that only a very light stress is enough to elicit a strong dSO avoidance response by the flies, potentially influencing flies in all sorts of behavioural assays performed in typical Drosophila research laboratories. Furthermore, elucidating what chemical the flies are avoiding, and whether other species would avoid this chemical could open the door to pest management applications.

I will present the recent progress my research team has made in elucidating the basis underlying the decision-making process to come to around 2-body length away from another fly, and to avoid another one when an alarm signal is perceived.

933V  Some Innexin Family Members Are Required for Cold Nociception Responses Mediated by Class III Dendritic Arborization Neurons Nicolas Netteneyer1, Rachel Barborek1, Maddie Ward1, Susan Helsell1 1) James Madison University, Harrisonburg, VA; 2) University of Washington, Seattle, WA; 3) Virginia Commonwealth University School of Medicine, Richmond, VA

The class III dendritic arborization (da) neurons mediate the behavioral response of third instar larvae to noxious cold (Turner et al. 2016. Curr Biol. 26:3116). Larvae respond to noxious cold by decreasing their overall length in a manner referred to as cringing. Innexin proteins form gap junctions, and may function in electrical synapsis, development of the nervous center, or tissue morphogenesis. GAL4/UAS RNAi expression coupled with a cold behavioral response assay assessed the possible role of each of the eight Drosophila Innexin family members. Expression of innexin gene-specific RNAi’s was driven specifically in class III da neurons. When possible, more than one RNAi line was studied for each gene. Alteration of the wild-type cold behavioral response was quantified as a loss of the cringe response (non-cringing). Of the eight innexin genes, RNAi expression against innexins 1 (ogre), 2, and 5 significantly inhibited the cold nociception response with both tested RNAi lines. The requirement for Innexin 1 function was confirmed by examining the response in ogre complete loss-of-function mutants. innexins 3, 4 (zero population growth), 6, and 8 (ShakingB) showed inconsistent results between their two tested RNAi lines. In order to resolve the ambiguity, behavioral assays are underway to test double-stranded RNAi constructs for each of these genes. Finally, RT-PCR is being conducted to determine which of these genes are expressed in class III da neurons.

934C  A drosophila model depicting braak-like propagation of tau pathology Aarya Vaikakkara Chithran1, Ben Batchelor2, Benjamin Boehme1, Lovesha Sivanantharajah3, Tianshun Lian1, Brad Richardson2, Eva Ruiz2, Efthimios Skoulakis4, Amrit Mudher2, Douglas Allan1 1) Department of Cellular and Physiological Sciences, University of British Columbia, Vancouver, BC, Canada; 2) Department of Biological Sciences, University of Southampton, Southampton, United Kingdom; 3) Department of Biological Sciences, University of Bangor, Bangor, United Kingdom; 4) Biomedical
Prion-like propagation through circuits is believed to be the mechanism by which tau pathology spreads throughout the brain in tauopathies like Alzheimer’s disease (AD). This is reflected in the neuropathological Braak-staging of disease and manifests in the progressive cognitive decline evident clinically. Though various synaptic proteins are implicated, the precise players and mechanism(s) mediating the trans-cellular spread of pathological tau species remain unclear. Furthermore, though the trans-cellular spread of pathological tau species has been demonstrated in many experimental models, the neurobiological consequences in recipient neurons are largely unknown. Moreover, in all such studies, the tau species that propagates is invariably mutated or isolated from pathological fractions of brains of tauopathy patients. This is puzzling given that it is wild-type tau that becomes pathological and spreads in AD, and this process is accompanied by neurodegeneration. We report a novel Drosophila model in which wildtype human tau expressed in select neuronal subsets becomes pathological and undergoes trans-cellular spread through adult brain circuits, causing neurodegeneration reminiscent of late stages of disease in AD brain. The superior genetic tractability of this model makes it ideally suited for dissection of the key players that mediate this pathogenic process through genetic and pharmacological modifier screens. Furthermore, the availability of functional and behavioural assays for many adult brain circuits will enable future studies to reveal the neurobiological consequences of spreading tau pathology more directly.

935A  DDX17 modulates FUS toxicity in an RGG-domain dependent manner Tyler Fortuna¹, Sukhleen Kour¹, Eric Anderson¹, Caroline Ward¹, Dhivya Rajasundaram², Christopher Donnelly³, Udai Pandey² ¹) Department of Pediatrics, Children's Hospital of Pittsburgh of UPMC; 2) Division of Health Informatics, Department of Pediatrics, Children’s Hospital of Pittsburgh, University of Pittsburgh School of Medicine, Pittsburgh, PA, USA; 3) Department of Neurobiology, University of Pittsburgh School of Medicine, Pittsburgh, PA, USA

Mutations in Fused in Sarcoma (FUS) have been linked with juvenile and aggressive form of amyotrophic lateral sclerosis (ALS). Cytoplasmic aggregation and defects in DNA repair pathway have been linked with ALS pathogenesis. However, the molecular mechanisms and genetic modifiers of DNA damage repair pathway are yet not known. While doing RNA-sequencing analysis of the Drosophila brains expressing FUS, we identified significantly altered genes and pathways involved in FUS-mediated neurodegeneration in vivo. We found that the DEAD-Box Helicase 17 (DDX17) expression level was significantly downregulated in response to mutant FUS in Drosophila and human cell lines. Interestingly, mutant FUS recruits nuclear DDX17 into cytoplasmic stress granules and biochemically interacts with DDX17 through the RGG1 domain of FUS. Importantly, genetic upregulation of DDX17 reduces cytoplasmic mislocalization and sequestration of mutant FUS into cytoplasmic stress granules. We identified DDX17 as a novel regulator of the DNA damage response pathway whose upregulation repairs defective DNA damage repair machinery caused by mutant neuronal FUS ALS. In addition, we show DDX17 is a novel modifier of FUS-mediated neurodegeneration in vivo. Our findings indicate DDX17 is downregulated in response to mutant FUS, and restoration of DDX17 levels suppresses FUS-mediated neuropathogenesis and toxicity in vivo.

936B  Metabolic Dysregulation in Frontotemporal Dementia Jackson Diltz, Emily Peterson, Em Teixeira, Angela Brillantes, Taylor Barber, Kristen Burke, Simone Alvares, Madeline O’Connor, Arianna Libby, Maximus Morton, Marla Tipping Providence College

Metabolic reprogramming is a common hallmark of many diseases. In recent years the focus on metabolic change in cancerous tissues has increased. However, fewer studies have investigated the metabolic shifts in neurodegenerative diseases. Metabolic reprogramming in neurodegenerative disease has been well documented and glucose uptake is even used as a key diagnostic indicator for some of these diseases. We are utilizing an established Drosophila model of the neurodegenerative disease, Frontotemporal Dementia (FTD), to investigate metabolic changes using a whole brain energy utilization assay. We are also studying the underlying molecular mechanism by using quantitative PCR to measure expression of metabolic enzymes in the brain and whole body. The goal of this project is to determine if metabolism could be a potential target for treatment of FTD.

937C  Uncovering the Mechanisms Behind the Neuroprotective Effect of Glycolysis in a Drosophila Model of ALS Nicholas Mortimore¹, Suvithanandhini Loganathan¹, Hannah Ball¹, Maria Macias¹, MyDuyen Tran¹, Gabrielle Peterson¹, Gabriel Birchak¹,⁴, Ernesto Manzo¹,⁴, Daniela Zarnescu¹,⁴ ¹) Department of Molecular and Cellular Biology, University of Arizona, Tucson, AZ; 2) Department of Neuroscience, University of Arizona, Tucson, AZ; 3) University of Pennsylvania, Philadelphia, PA; 4) Vollum Institute, Oregon Health and Science University, Portland, OR

Amyotrophic Lateral Sclerosis (ALS) is a progressive neurodegenerative disease that affects upper and lower motor neurons primarily in individuals aged forty years or greater. ALS causes loss of motor control, paralysis, and eventually respiratory failure and death within 2-5 years of diagnosis; there is no known cure for this disease. TAR DNA-binding protein (TDP-43) is a pathological marker of ALS and aggregation of TDP-43 is observed in 97% of ALS cases. Our lab has generated a Drosophila model of ALS based on overexpression of human TDP-43 in a motor-neuron specific manner which recapitulates key disease phenotypes including cytoplasmic accumulation of TDP-43, neuromuscular...
juncture (NMJ) abnormalities, decreased lifespan, and locomotor defects. Glycolysis upregulation by an increase in Phosphofructokinase-1 (PFK1; PFK in Drosophila) has been shown to have a neuroprotective effect in Drosophila models of ALS based on TDP-43 proteinopathy. Here we aim to decipher the mechanisms underlying the restorative effect of PFK overexpression (OE) in ALS. One possible mechanism is that PFK assembles in complexes that support energy demands locally, at synapses. Supporting this scenario are our observations that PFK forms puncta within Drosophila NMJs in the context of mutant TDP-43 OE. This is consistent with recent evidence from other groups showing that PFK can form clusters at the synapses of C. elegans where it associates with other glycolytic enzymes to form Glycolytic bodies (G-bodies) and supports synaptic vesicle cycling under stress. Our preliminary data support this hypothesis as evidenced by FM1-43 dye uptake experiments showing that PFK OE rescues TDP-43 induced synaptic vesicle cycling deficits.

Another possible mechanism is that pyruvate, the end product of glycolysis, which was found to be increased in multiple ALS models may act as a scavenger of reactive oxygen species as shown in cancer cells. We are currently interrogating the composition of G-bodies with mass spectrometry analyses of PFK complexes to further test the hypothesis that G-body composition is altered in the context of TDP-43 proteinopathy. These findings may inform novel therapeutic strategies based on improved cellular energetics for ALS and related neurodegenerative disorders.

938A A CRISPR-Cas9 Mediated Knockout of RNaseZ in Drosophila Neurons Max Luf, Ekaterina Migunova, Jake Nelson, Edward Dubrovsky Department of Biological Sciences, Fordham University

The RNaseZ gene is a vital and highly conserved gene with homologs in all eukaryotes. It plays an essential role in the maturation of tRNA, being the only enzyme to process the 3'–ends of pre-tRNA in both the nucleus and mitochondria. Point mutations in the human homolog, ELAC2, have been linked to a heterogeneous group of neurological conditions, such as microcephaly and untreatable epilepsy. Because these conditions result in such a poor quality of life for patients, we seek to understand what unique role RNaseZ may play in the nervous system.

Here we report the importance of RNaseZ to the Drosophila nervous system and demonstrate its role using neuron-specific knockouts of RNaseZ. To determine if point mutations of RNaseZ homologous to those in humans could elicit neurological dysfunction in Drosophila we first generated two transgenic lines (gZ1541 and gZ5208) carrying mutant alleles of RNaseZ under its natural promoter. When the endogenous RNaseZ is removed these transgenes led to increased neurodegeneration in adult flies, decreased motor ability, and shortened life span. To then determine if this process was cell-autonomous, we generated a Cas9-mediated pan-neuronal knockout (KO) of RNaseZ in all Drosophila neurons. This resulted in developmental delays, debilitating motor deficits, and death shortly after eclosion. Because of the dual role of RNaseZ in the mitochondria and the nucleus, we created a Cas9-resistant rescue construct and used it to restore wt RNaseZ exclusively to the nucleus, creating a mitochondrial KO of RNaseZ in Drosophila neurons. The mitochondrial KO of RNaseZ produced a phenotype in longevity and locomotive ability almost identical to that of flies with the pan-neuronal KO. Overall, our findings improve our understanding of the unique role that RNaseZ plays in the nervous system and demonstrate how tissue-specific Cas9 engineering and Cas9-resistant rescue constructs can be applied to the modeling of neural degeneration in Drosophila.

939B Mechanism of adult neurodegeneration in drop-dead mutants Unmila Jhuti, Edward Blumenthal Marquette University, Milwaukee, WI

Adult neurodegenerative diseases such as Alzheimer’s and Parkinson’s are one of the main causes of death for elderly population worldwide. Using Drosophila models of neurodegeneration (ND) to study such diseases has been proven useful because they show striking similarities to the human diseases. This project aims to understand ND in flies mutant for drop-dead (drd). Absence of drd expression is associated with adult ND and short lifespan, but the molecular pathways responsible for ND in this model have not been identified. Hyperactivation of innate immunity (HII) has been linked to ND both in human diseases and in Drosophila. It is yet to be identified how HII pathways are activated in adult neurodegenerative diseases. Preliminary data suggest a causal connection of HII with ND in drd mutants. Expression of the antimicrobial peptide genes DptB, CecA1 and AttC was increased by 2-4 fold in drd mutant brains compared to heterozygous sibling controls. Furthermore, drd mutant flies that are heterozygous for mutations in the immune genes rel or imd showed an increased median lifespan (8-10 days) compared to drd mutant controls (4-7 days). Although ND has been identified in mutant drd brains, the spatiotemporal pattern and pathway(s) of cell death have not been characterized. Immunostaining with the apoptotic marker Dcp-1 suggested activation of caspase-dependent apoptosis in the brains of drd mutants as early as the day of eclosion (P0). Intense and spatially dispersed staining was observed in older (P2-3) flies. Staining with the non-specific cell death marker acridine orange revealed widespread cell death beginning at P0.

Previous work has shown that expression of drd in the respiratory tracheae is necessary and sufficient to prevent ND. Thus, brain cell death could potentially be connected to disruption of gas exchange and hypoxia. However, staining mutant brains with the ROS marker dihydroethidium showed the presence ROS only in older flies (P2-3), but not in younger flies (P0), suggesting that hypoxia is not a primary cause of ND in drd mutants.

Studying the role of drd will provide significant insight into the link between HII and adult ND. Identification of the pathways leading to cell death can help to understand cellular mechanisms potentially responsible for adult
neurodegeneration.

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Keywords: innate immunity, apoptosis, cell death, hypoxia, neurodegeneration.

940C The ketone body beta-hydroxybutyrate ameliorates molecular and behavioral pathological markers in a Drosophila model of glial tauopathy. Celya D. Dahmani1, Julia O’Connell2, Timothy O’Toole1, Kenneth J. Colodner2, Geoffrey R. Tanner1,3 1) University of Connecticut, Storrs, CT; 2) Mount Holyoke College, South Hadley, MA; 3) Institute for the Brain and Cognitive Sciences, Storrs, CT

Tauopathies are a class of neurodegenerative diseases characterized by abnormal intracellular aggregations of the microtubule-stabilizing protein tau. Symptomology of tauopathies features impaired cognition—including difficulties with learning and memory—caused by progressive neuronal and synaptic degeneration. Tau pathology is often characterized by an accumulation of hyperphosphorylated tau protein resulting in the formation of neurotoxic tau aggregates, also known as neurofibrillary (NFT) and gliofibrillary (GFT) tangles. These aggregates may hinder neural glycolytic metabolism and result in degradation of the myelin sheath and other glial components. Ketone bodies (KBs), metabolites of the putatively-neuroprotective high-fat and low-carbohydrate ketogenic diet (KD), bypass cytoplasmic glycolysis and enter directly into mitochondrial metabolism. The KD and KBs serve to elevate cellular ATP/ADP ratio and thus may rescue neural energetics. We have obtained evidence strongly suggesting that KBs may exhibit neuroprotective effects in a Drosophila melanogaster model of traumatic brain injury (TBI); repeated TBI may lead to the progressive neurodegenerative condition termed Chronic Traumatic Encephalopathy (CTE), which is characterized in part by intracellular aggregation of tau protein in neural cells. We employed a Drosophila model of tauopathy that allows for spatial restriction of exogenous tau protein overexpression to glial cells and for temporal restriction of tau overexpression to adulthood, so as to avoid developmental effects of neural tau protein overexpression. We directly applied a racemic mixture of the KD metabolite beta-hydroxybutyrate (β-HB)—the major circulating ketone body (KB)—to a standard high-carbohydrate Drosophila diet to determine whether dietary KB supplementation can ameliorate tauopathy-associated learning deficits. We found that β-HB markedly improves tauopathy-induced learning deficits in mixed-sex groups of adult Drosophila and reduces levels of phosphorylated tau in the brains of male flies subjected to induced glial tauopathy.

941A Phagocytic glia mediate prion-like spreading of mutant huntingtin aggregates in Drosophila brains Kirby Donnelly, Aprem Zaya, Graham Davis, Olivia DeLorenzo, Margaret Panning Pearce University of the Sciences

A key pathological and diagnostic feature of most neurodegenerative diseases is appearance of insoluble aggregates due to protein misfolding in the central nervous system (CNS). Accumulation of toxic protein aggregates in the brain is partially offset by a number of clearance mechanisms, including the ubiquitin-proteasome pathway, autophagy, and engulfment by phagocytic glia. A growing body of evidence supports the hypothesis that pathogenic aggregates associated with many neurodegenerative diseases behave similarly to infectious prions—they spread from cell-to-cell and nucleate the aggregation of natively-folded versions of the same protein, events which are thought to contribute to propagation of aggregates in the brain. We have recently reported that mutant huntingtin (mHTT) aggregates associated with the inherited neurodegenerative disorder Huntington’s disease spread in a “prion-like” manner between synaptically-connected neurons and glia in the Drosophila CNS. mHTT aggregates formed in presynaptic olfactory receptor neuron (ORN) axons directly nucleate the aggregation of soluble wild-type HTT (wtHTT) proteins expressed in post-synaptic partner projection neurons (PNs) or nearby glia. ORN-to-PN and ORN-to-glia transfer of mHTT aggregates is enhanced when ORN activity is silenced, slowed when caspase-dependent apoptosis in ORNs is blocked, and remarkably, require expression of the conserved glial scavenger receptor Draper/MEGF10. Further, mHTT aggregate transmission between synaptically-connected ORNs and PNs requires a transient visit to the glial cytoplasm, indicating that phagocytic glia directly mediate aggregate transfer between neurons in vivo. Preliminary data from our lab indicate that entry of phagocytosed neuronal mHTT aggregates into the glial cytoplasm involves certain Rab GTPases with proposed roles in intracellular vesicle membrane fusion related to phagosome maturation, suggesting that inefficient or incomplete progression of engulfed material in the phagolysosomal system may generate seeding-competent mHTT aggregate species in the glial cytoplasm. In support of this, expression of mHTT in ORNs accelerates age-related decline in glial phagocytic clearance of axonal debris. Together, our findings add to a growing understanding of phagocytic glia as double-edged players in neurodegeneration—these cells clear neurotoxic aggregates, but age- and/or disease-related defects in this process can paradoxically promote aggregate propagation in the brain.

942B Dynamic transcriptional changes in the adult Drosophila central nervous system highlights potential coordination of stress and repair responses following traumatic brain injury Eddie Cho1, Alec Candib1, Reina Hastings3, Gina Torabzadeh1, Jesse Rojas1, Sharon Zhang1, Marta Lipinski2, Eric Ratliff1, Kim Finley1 1) Shiley BioScience Center and Biology Dept, San Diego State University, San Diego, CA; 2) Department of Anesthesiology, University of Maryland School of Medicine, Baltimore, MD

Traumatic brain injury (TBI) from accidents, domestic violence, sports and combat are a major cause of worldwide...
mortality and disability. Several groups have shown that adult *Drosophila* can be an effective model to examine the physiological, behavioral, neuronal, and molecular responses following TBI exposure. In this study, we have continued our Drosophila TBI research using RNA-sequencing to examine dynamic acute and long-term changes to central nervous system (CNS) gene expression profiles in adult flies following severe (sTBI) and mild repetitive TBI (mTBI) exposure. Principal component analysis (PCA) highlight highly dynamic transcriptional changes in male and female flies, with over a third of the *Drosophila* transcriptome showing 1.5-fold (+/-) alterations 4-hrs post-TBI. By 24-hrs, both genders rapidly shifted toward baseline transcriptome levels. 4-days post TBI male fly cohorts maintained a more baseline transcriptome profiles, while female flies had a second round of gene expression changes. Transcriptional changes trend toward baseline levels at 7-days post. Genes with significant alterations were analyzed using DAVID to identify functional gene-pathway clusters. At 4-hrs both genders had significant changes to NFkB and IMD pathway components, indicating rapid activation of stress and inflammatory signaling. At this time dramatic changes to neuronal/ERAD stress responses, ribosomal, vesical trafficking, and membrane transport components also occurred. At 24-hrs, inflammation remains elevated, along with alterations to cytoskeletal, kinase-phosphatase and proteolytic pathway members. At 4-days, there is a significant increase in select metallopeptidases as well as changes to organelle, protein and basement membrane modifying processes. 7-days post-TBI, DAVID revealed changes to alternative splicing factors, ER, cytochrome P450, glutathione metabolism, and members of the scavenger receptor pathways. Cell cycle components demonstrated multiplex differences, with PCNA and several cyclin genes increased significantly (4-hrs). Our novel trauma injury paradigm illustrates the effectiveness of model systems to identify conserved genetic factors influencing the complex stress and repair mechanisms underscoring *in vivo* trauma exposure. The goal of these findings is to identify genetic factors that are linked to adverse acute or protracted TBI outcomes and in the discovery of novel therapies for those experiencing trauma.

**943C Assessing Novel Therapeutics with a Drosophila Model of Neural Aging and Stressors** Alec Candib, Brandon Molina, Jessica Mastroianni, Nicholas Lee, Natasha Sam, Eddie Cho, Kyle DeAlva, Andrea Gonzalez, Gina Torabzadeh, Josephine Vu, Joy Phillips, Kim Finley Department of Biology, San Diego State University, San Diego, CA

A wide range of potential therapeutics, including modified diets, cannabis, turmeric, and probiotics, are being examined as treatments for disorders associated with aging and neural stress. In this report, we detail a unified *in vivo* method to test the safety, efficacy, therapeutic dosage, and potential molecular mechanism of neuroprotective products using adult *Drosophila melanogaster* as a model system. This includes an integrated three-part platform that examines the impact of individual compounds and biologics on neural aging, TBI responses, and cytotoxic β-amyloid (Aβ). Adult flies were administered different dosages of cannabis-derived cannabinoids (CBD, CBN, THC), select drugs (1J47, metformin, donepezil), and active and inactive biologics (LGG, IAB). Aging studies show select dosages of several therapies preserved locomotor behaviors (negative geotaxis response, NGR) and promoted longevity, while other therapies exacerbated age-related NGR decline and reduced longevity. Pre-treatment with several therapies also helped preserve NGR profiles and promote longevity in traumatized fly cohorts. To examine the potential molecular mechanisms of these compounds, changes to the autophagy and inflammation (NF-kB) pathways were investigated in adult tissues. Several compounds impaired basal and fasting-induced autophagic flux, while others increased autophagic flux and reduced protein aggregate formation. Several therapies reduced the expression of downstream NF-kB signaling targets (antimicrobial peptides) after TBI. The impact of CBD, IAB, and Donepezil on longevity, NGR, NF-kB signaling, and autophagy were studied in flies expressing Aβ peptides. Together, these studies showed that these compounds have a significant impact on *Drosophila* physiology and molecular pathways. Generally, exposure to every compound but THC had similar neuroprotective and longevity-promoting effects, while THC had a limited positive impact only following mTBI. These findings demonstrate the versatility of *Drosophila* as a drug discovery model system and highlight the need for future studies to clarify the molecular interactions and downstream consequences of natural products on complex biological systems and tissues.

**944A Identify novel approaches suppressing stress granule assembly to mitigate TDP-43-mediated neurotoxicity** Quinlan Mewborne, Jessica Chalk, Junli Gao, Ke Zhang Mayo Clinic Jacksonville

Amyotrophic lateral sclerosis (ALS) is a degenerative disease of the motor neurons sometimes associated with frontotemporal dementia (FTD). These two neurodegenerative diseases are progressive, increasingly prevalent, and currently incurable. A major challenge of the therapeutic development of these diseases is our incomplete understanding of the pathogenic mechanism. Previous studies have suggested that stress granules (SGs), RNA/protein condensates assembled in cells under stress, play a critical role in ALS/FTD pathogenesis. Consistent with this notion, chemical inhibitors suppressing SG assembly suppress neurodegeneration in cellular and animal models of ALS/FTD. However, these inhibitors exhibit detrimental side effects in mammals in preclinical studies, preventing their translation into clinics. Thus, we propose to use a fly model to identify novel ways targeting SG assembly as potential therapeutic strategies for ALS/FTD.

Tar DNA-binding protein 43 (TDP-43) is an SG protein, whose cytoplasmic aggregation has been implicated in ~98% of...
ALS and ~45% of FTD. Indeed, SG assembly has been shown to trigger TDP-43 aggregation in multiple models of ALS/FTD. Consistent with these data, overexpressing a cytoplasmic form of TDP-43 causes eye degeneration in flies. Using this model, we screened through the RNAi of 250 fly genes whose human homolog have been shown to promote SG assembly in cultured cells. We identified ~20 genes whose RNAi strongly or moderately suppresses TDP-43-mediated eye degeneration in our flies and will further characterize their roles in ALS/FTD pathogenesis.

945B  Poly(ADP-ribose) Promotes the Condensation of C9ORF72 Arginine-rich Dipeptide Repeat Proteins Ke Zhang, Junli Gao, Ritika Punathil, Tania Gendron, Udit Sheth Mayo Clinic Florida

Arginine-rich dipeptide repeat proteins (R-DPRs), abnormal translational products of a GGGGCC hexanucleotide repeat expansion in C9ORF72, play a critical role in C9ORF72-related amyotrophic lateral sclerosis (ALS) and frontotemporal dementia (FTD), the most common familial form of ALS and FTD (c9ALS/FTD). R-DPRs undergo phase separation to form condensates in vitro and, as a c9ALS/FTD pathological hallmark, aggregate and sometimes co-aggregate with TDP-43 in patients. However, how these processes are regulated is unclear. Here, we show that poly(ADP-ribose) (PAR) interacts with R-DPRs, promotes R-DPR phase separation and toxicity in vitro, and correlates with insoluble R-DPR and TDP-43 in patient postmortem tissue. Loss of PAR suppresses R-DPR aggregation and neurodegeneration in fly or cellular models of c9ALS/FTD. Our findings identified PAR as a pathogenic contributor promoting R-DPR condensation both in vitro and in vivo, suggesting the potential of R-DPR condensation as a target for c9ALS/FTD therapeutic intervention.

946C  Behavioral changes and tau pathology in response to traumatic brain injury in Drosophila Roilea Maxson1, Christine Smoyer1,2, Kailea Wiese3, Kassandra Ori-Mckenney1 1) University of California, Davis; 2) University of Kansas

Traumatic brain injury (TBI) is a disruption of normal brain function that results from mild to severe impacts to the head, and can affect memory, as well as behaviors such as anxiety, aggression, and depression. In addition, TBI is the leading risk factor for late onset neurodegenerative diseases such as Alzheimer’s Disease. One major hallmark of TBI is the presence of hyperphosphorylated tau in the brain and cerebrospinal fluid. In a healthy brain, tau is a microtubule associated protein that controls microtubule based processes, but upon hyperphosphorylation, tau forms aggregates called neurofibrillary tangles (NFTs) that are thought to contribute to neurodegeneration. However, the series of events that cause tau hyperphosphorylation and NFT formation in response to injury, and how these contribute to neurodegeneration are not well established. We are utilizing a high impact method to subject flies expressing 2N4R human tau pan-neuronally to TBI in an effort to study the contribution of tau phosphorylation and oligomerization to neurodegeneration and behavioral decline. At 24 hours after inflicting TBI on 1 week old virgin males, we recorded flies to observe courtship and aggression. We found that pan-neuronal expression of tau caused an increase in inter-male aggression in flies subjected to TBI, both in terms of the amount of time spent in aggressive acts and in the total number of aggressive acts. We have screened a variety of specific drivers for neuronal types known to be involved in mating and aggressive behaviors, and have identified dopaminergic and serotonergic circuits as contributors to increased aggression. In addition to our behavioral studies, we are performing immunohistochemistry on fly brains from 24 hours, 1 week, or 3 weeks post-injury to determine if there are any differences in tau localization and/or neurodegeneration. The neurodegeneration associated with TBI and tau expression in Drosophila can be indicated via the presence of vacuoles. We aim to determine if coupling TBI and tau expression affects tau localization or exacerbates the occurrence of neurodegeneration seen with TBI or tau expression alone. In addition, we are analyzing the phosphorylation and aggregation of tau isolated from fly brains using biochemical techniques. Ultimately, we hope to elucidate how different molecular states of tau contribute to neurodegeneration and lead to alterations in brain function and behaviors.

947A  Comparing the Neurotoxic Effects of P3 (Aβ17-42) and Aβ1-42 using Drosophila as an Alzheimer’s Disease Model Marisa Fujimoto, Alfredo Rojas Moreno, Yuhao Pan, Emmanuel Akabogu, Gloria Qui, Joey Wong, Katrina Haas, Selma Alamari, Tatyannah Farsh, Annika Streeb, Jeremy Lee Department of Molecular, Cell, and Developmental Biology, University of California, Santa Cruz, CA

Alzheimer’s disease (AD) is a neurodegenerative disease associated with nearly 6 million deaths in the U.S. every year. AD is characterized by a loss of memory, decline in cognitive abilities, and changes in mood and personality. A hallmark of AD pathology is the presence of Aβ plaques, which are formed by the aggregation of Aβ peptides. The Aβ1-42 peptide is produced by the cleavage of amyloid precursor protein (APP), first by β-secretase and then γ-secretase. An alternative pathway uses α-secretase instead of β-secretase and generates a peptide called P3 (Aβ17-42), that is 16 amino acids shorter than Aβ1-42 and is composed of the hydrophobic portion of Aβ1-42. This P3 cleavage pathway is often characterized as non-amyloidogenic, however, recent evidence suggests P3 is amyloidogenic (Kuhn, et al., 2020). As such, there is reason to think that P3 could have similar deleterious effects as Aβ1-42. The purpose of our research is to characterize P3 peptide’s neurotoxicity and determine whether it interacts with Aβ. To investigate this, we generated P3 expressing flies, Aβ1-42 expressing flies, and Aβ1-42/P3 co-expressing flies, in which the peptides were expressed pan-neuronally, to examine the neurodegenerative effects of these two peptides independently, as well as their synergistic neurotoxic effects. RT-PCR experiments have confirmed expression of the transgenes. A longevity assay was conducted to measure lifespan in the transgenic flies and the Rapid Iterative Negative Geotaxis (RING) assay was performed to
monitor their locomotor performance in relation to age. These results, along with electron micrograph analysis of the peptides’ effects when expressed in fly eyes, indicate that P3 expression alone has similar, but less severe, effects as Aβ on lifespan, behavior, and neurodegeneration. Our results also suggest that P3 exacerbates the effects of Aβ1-42 when co-expressed. RNA sequencing is being used to further investigate the effects of these peptides by examining whether the expression of neurodegenerative-related genes and neuroplasticity-related genes, known to be affected by Aβ1-42, are similarly affected by P3. We will also conduct co-immunoprecipitation experiments using Aβ1-42/P3 co-expressing flies to determine whether the two peptides directly interact. Our research will help to determine whether P3 might be a direct contributing factor to AD pathology and/or have an effect via its interaction with Aβ.

948B  Observing the Effects of the Human Peptide, LL-37, on Aβ₄₂’s Neurotoxicity and Effects on Gene Expression Using a Drosophila Model of Alzheimer’s Disease Ruby Guevara, Kenneth Owyang, Jennifer Tan, Marissa Joe, Taylor Jones, Duyen Nguyen, Emmanuella Tetteh, Shannon Twardy, Celine Neudorf, Shelly Huynh, Jeremy Lee Department of Molecular, Cell, and Developmental Biology, University of California, Santa Cruz, CA

The hallmark of Alzheimer’s disease (AD) is the presence of amyloid plaques in affected areas of the brain, which are aggregates of the amyloid-beta (Aβ) peptide. Aβ, especially Aβ₄₂, is known to be neurotoxic and crucial in AD pathology. While the function of Aβ is not well understood, recent studies have shown it to have antimicrobial effects (Kumar et al., 2016). It has also been shown to interact with the human antimicrobial peptide LL-37 in binding assays in vitro (De Lorenzi, et al., 2017). If LL-37 and Aβ interact in vivo, it may affect the neurotoxicity of Aβ.

In order to determine whether LL-37 affects Aβ neurotoxicity, we generated four transgenic flies pan-neuronally expressing human Aβ₄₂ and/or LL-37: LL37-expressing, Aβ₄₂-expressing, Aβ₄₂/LL-37 co-expressing, and non-expressing controls. Longevity assays and RING climbing assays were conducted to determine whether LL-37 has an effect on Aβ₄₂’s neurotoxicity in vivo. The results showed that co-expression of LL-37 and Aβ₄₂ led to longer lifespans than expression of only Aβ₄₂, but shorter lifespans than non-expressing controls. In developmental assays, these co-expressing flies had higher survivorship to eclosion than Aβ₄₂-expressing flies. In addition, RING assays showed that dual-expressing flies had a significant improvement in locomotive functions compared to LL37-expressing and Aβ₄₂-expressing flies. These results indicate that LL-37 might attenuate the deleterious effects of Aβ₄₂ in Alzheimer’s disease.

We are assessing LL-37 and Aβ₄₂ expression levels in our transgenic flies by qRT-PCR and Western blot to determine whether co-expression of LL-37 has an effect on Aβ₄₂ transcript or protein levels. We also intend to assess whether the expression of LL-37 modulates Aβ’s known effects on the expression of a variety of genes involved in synaptic plasticity and neuronal function. To do this, we are performing RNA sequencing analysis to obtain a quantitative snapshot of RNA transcripts present within our four genotypes. Using a baseline comparison of Aβ₄₂-expressing flies versus wild type, we will focus on key genes of interest previously shown to be affected by Aβ₄₂. This will allow us to compare any modulating effect LL-37 has on Aβ₄₂’s effects on gene expression, by assessing co-expressing flies and noting any key differences in their RNA expression profiles. This will help provide insight into the mechanism by which LL-37 attenuates Aβ’s neurotoxic effects.

949C  An Analysis of the Microbiota of Various Drosophila melanogaster Parkinson’s Disease Models Evan Marshman¹, Samara Petersen¹, Zachary Pickup¹, Gerald Call¹, John Chaston¹ 1) Department of Plant and Wildlife Sciences, College of Life Sciences, Brigham Young University, Provo, UT; 2) Department of Pharmacology, College of Graduate Studies, Midwestern University, Glendale, AZ; 3) Biomedical Sciences Program, College of Graduate Studies, Midwestern University, Glendale, AZ

The gut-brain axis is defined as the bi-directional interactions that occur between the brain and the gut of an organism. Considering this interaction, the brain is also known to affect the gut microbiome of an organism. The microbiome has specifically been shown to be affected by many neurological diseases. Parkinson’s disease (PD) is a neurodegenerative disease that affects motor function. We are interested in studying the interactions between the gut microbiome and PD. Current research has shown that the microbiome of an individual with PD differs from the microbiome of a healthy individual. This has been exhibited in humans and mice. We predicted that we would detect this same variation in the microbiota of D. melanogaster (fruit fly) models of PD, relative to a genetically matched control. To test this hypothesis, we used five PD fruit fly models obtained from four different laboratories, each mimicking human PD by different mutations in their genome. We collected and analyzed 16S rRNA sequence, reporting the microbiota composition in our fly samples. We found various microbes that were differentially abundant between control and PD models at a genus level. We also show that the bacterial diversity of PD models is overall significantly lower than in control flies, consistent with the idea that individuals with declined health show lower diversity in their gut microbiome. The major conclusion of our work is that there is significant variation in microbiota composition between conventionally reared fly models of PD and their controls. This shows that the differences seen in humans and other models are also present in fruit flies.

950A  A photo-switchable assay system for dendrite degeneration in Drosophila melanogaster Han-Hsuan Liu¹, Chien-Hsiang Hsu¹, Lily Jan¹, Yuh-Nung Jan¹ 1) Howard Hughes Medical Institute, Departments of Physiology, Biochemistry and Biophysics, University of California, San Francisco, San Francisco, CA, United States.; 2) Department of Pharmaceutical Chemistry, University of California, San Francisco, San Francisco, CA, United States.
Neurodegeneration arising from aging, injury or disease has devastating health consequences. Whereas neuronal survival and axon degeneration have been studied extensively, much less is known about how neurodegeneration impacts dendrites. To develop an assay for dendrite degeneration in the Drosophila peripheral nervous system, we used photo-switchable caspase-3 (caspase-LOV) to induce neurodegeneration with tunable severity by adjusting illumination duration in class 4 dendrite arborization (c4da) neurons during development and adulthood. We found that c4da neurons were able to sustain mild caspase-3-induced neurodegeneration and continue to grow. To ask whether mechanisms underlying axon degeneration also govern dendrite degeneration, we tested the involvement of the Wallerian degeneration pathway by examining the effects of expressing the mouse Wallerian degeneration slow (Wld<sup>s</sup>) protein in larval and adult c4da neurons. Here we report Wld<sup>e</sup> expression protect c4da neurons from caspase-3 induced dendrite degeneration in larval and adult c4da neurons through reducing the dendrite elimination without affecting dendrite addition. At the functional lever, Wld<sup>e</sup> expression improved the recovery of thermal nocifensive behavior that was impaired by chronic low-level of caspase-LOV activity. By establishing ways to induce graded dendrite degeneration, we uncover a protective role of Wld<sup>e</sup> in caspase-3 induced dendrite degeneration.

**951B TDP-43 expression in dementia-relevant circuits causes axonal degeneration and behavioral deficits in Drosophila**

Reed Bjork<sup>1</sup>, Rebekah Keating Godfrey<sup>1</sup>, Christi Williams<sup>1</sup>, Hillary Cowell<sup>1</sup>, Allison Michael<sup>1</sup>, Grace Halaufia<sup>3</sup>, Erik Lehmkühl<sup>2</sup>, Eric Alsop<sup>2</sup>, Kendall van Keuren-Jensen<sup>2</sup>, Daniela Zarnescu<sup>1</sup> 1) The University of Arizona; 2) The Translational Genomics Research Institute

Trans active response (TAR) DNA-Binding protein 43 (TDP-43), an evolutionarily conserved RNA/DNA binding protein that regulates RNA processing, is known to form pathological, cytoplasmic inclusions in numerous neurodegenerative diseases. These cytoplasmic accumulations, collectively referred to as TDP-43 proteinopathy, are a hallmark of motor neuron degeneration in amyotrophic lateral sclerosis (ALS, 97% of cases) and are also detected in nearly half of frontotemporal dementia (FTD) and Alzheimer’s disease (AD) cases, establishing TDP-43 as an important candidate for modeling neurodegenerative disease. Drosophila models of TDP-43 proteinopathy in ALS and other neurological disorders have elucidated important molecular dynamics of neurodegeneration, yet such models have not yet been employed to understand TDP-43 dependent pathomechanisms in dementia. Given the striking overlap between ALS and FTD and our own success in modeling TDP-43 proteinopathy in a fly model of ALS, we have developed a novel fruit fly model of dementia by overexpressing human TDP-43 in the mushroom bodies (MBs) of the Drosophila brain. Insect MBs form an elaborate, associative network that overlaps in function and gene expression with neurons in the human hippocampus and frontal cortex, making it an appropriate brain structure to model human dementia. We demonstrate that TDP-43 mislocalizes from the nucleus to the cytoplasm and forms axonal inclusions accompanied by age-dependent behavioral deficits and axonal degeneration in MBs. Importantly, cognitive deficits in working memory and sleep appear prior to our ability to detect axonal degeneration, paralleling what is seen in human disease. Based on recent studies identifying a role for TDP-43 in translation dysregulation, we sought to identify the molecular targets of TDP-43 in the MBs using RNA immunoprecipitations and RNAseq. These experiments identified several candidate targets of TDP-43 in mushroom bodies, including Dally-like protein (Dlp/GPC6), a translational target that we recently reported in Drosophila ALS models and patients. These findings suggest that TDP-43 overexpression in MBs causes FTD-relevant phenotypes that may help uncover novel mechanisms of disease. Future experiments are aimed at identifying additional FTD-relevant targets of TDP-43, including novel therapeutic strategies.

**952C A small molecule ion channel screen to suppress gliopathic epilepsies**

Walt Krueger<sup>1,2</sup>, Bidisha Roy<sup>1</sup>, Jungsoo Han<sup>3</sup>, Benjamin Geier<sup>1,2</sup>, Lawrence Reiter<sup>1,4,5</sup> 1) Department of Neurology, UTHSC, Memphis, TN; 2) IBSP Program, UTHSC, Memphis, TN; 3) Department of Biochemistry, UT Southwestern, Dallas, TX; 4) Department of Pediatrics, UTHSC, Memphis, TN; 5) Department of Anatomy and Neurobiology, UTHSC, Memphis, TN

Duplication 15q syndrome (Dup15q) is caused by the presence of at least one extra copy of the 15q11.2-q13.1 region. Characteristics of Dup15q include hypotonia, intellectual disability, autism spectrum disorder and, in the majority of isodicentric 15 cases, pharmacoresistant epilepsy. We previously constructed a fly model that successfully recapitulates the seizure phenotype observed in Dup15q individuals by overexpressing Drosophila Ube3a (Dube3a) in glial cells. To identify new anti-epileptic drugs, our lab developed a medium throughput screening method to repurpose previously FDA or otherwise approved chemical libraries for their ability to suppress seizures. We recently used this model to screen 1,280 compounds from the Prestwick Chemical Library. Eight compounds were identified that reduce seizure recovery time by at least 50% in both male and female flies. Most of these compounds act through serotonin or dopamine receptors to increase Na<sup>+</sup>/K<sup>-</sup>-ATPase activity in glial cells (Roy et al. (2020) *Biol Psychiatry* 88(9):698-709). In the current study, we evaluated 70 compounds from the Screen-Well Ion Channel Ligand Library for their ability to suppress seizures in our glial cell specific epilepsy model. The primary screen was composed of 24 calcium channel modulators, 23 potassium channel modulators, 10 sodium channel modulators, 7 intracellular calcium modulators, and 6 other miscellaneous drugs. We identified 8 compounds that suppress seizures in repo>Dube3a flies by at least 50%. Seventy-five percent of these compounds are potassium modulators and 25% are calcium modulators. ATP-sensitive Inward Rectifying K+ channel (KATP) modulation is a shared commonality among 3 of the 8 compounds. To evaluate the
potentially critical role KATP modulation plays in seizure suppression, as well as to investigate the importance of glial-specific neuronal modulation, we are currently testing drug efficacy on flies that simultaneously express Dube3a and an RNAi against the Drosophila homologue for KATP (irk) in glial cells. We expect to find that KATP modulators fail to suppress seizures in the absence of sufficient irk channel expressions. These studies will lead directly to new candidate drugs, specific ion channel agonists and antagonists, that may eventually be used clinically to suppress seizures in Dup15q syndrome.

953A Characterization of the Fly Models for Glutaminase-related Neurological Disorders Zelha Nil1-2, Carlos Ferreiraa, Camilo Toro6, David Adams6, Oguz Kanca1-2, Shinya Yamamoto1-2, Hugo Bellen1-2, Undiagnosed Diseases Network 1) Department of Molecular and Human Genetics, Baylor College of Medicine, Houston, TX, USA; 2) Jan and Dan Duncan Neurological Research Institute, Texas Children’s Hospital, Houston, TX, USA; 3) Metabolic Medicine Branch, National Human Genome Research Institute, National Institutes of Health, Bethesda, MD, USA; 4) Adult NIH Undiagnosed Diseases Program, National Human Genome Research Institute, National Institutes of Health, Bethesda, MD, USA; 5) Human Biochemical Genetics Section, National Human Genome Research Institute, National Institutes of Health, Bethesda, MD, USA

Glutaminase (GLS) encodes a mitochondrial enzyme that converts L-glutamine into L-glutamate in mitochondria. In addition to serving as building blocks of proteins, these amino acids have many additional roles in vivo including maintaining the cellular redox status, fueling the Krebs cycle, and synaptic transmission. Recent studies have identified rare human diseases that are caused by loss- (LOF) or gain-of-function (GOF) variants in GLS. Due to the pleiotropic roles of glutamine, glutamate and other related metabolites in diverse cellular processes, the precise molecular mechanisms underlying the human diseases are unknown. Here, we generated and characterized different alleles of fly GLS and transgenic human GLS expressing lines to study these disorders. We found that complete loss of GLS (GLS<sup>ΔMito</sup>) or the loss of the isoforms with strong mitochondrial targeting signals (GLS<sup>Gal4ΔAll</sup>) do not cause any obvious defects. However, a dominant negative allele expressing the short N-terminal fragments of different GLS isoforms (GLS<sup>CON</sup>) shows decreased survival, shortened life-span, reduced body size, progressive loss of motor skills and glutamine accumulation similar to diseases caused by some GLS missense variants in humans. Moreover, we found that the ubiquitous overexpression of a potential GOF variant (p.Ser482Cys) in human GLS in wild-type flies is toxic, whereas overexpression of the reference or predicted LOF variants do not show any effect. These preliminary data argue that Drosophila is a promising model to explore molecular mechanisms of rare human GLS diseases.

954B Biallelic variants in OGDHL cause a neurodevelopmental spectrum disease featuring epilepsy, hearing loss, visual impairment, and ataxia Zheng Yie Yap1, Stephanie Ethfymiou2, Simone Seiffert3, Karen Vargas Parra1, Suyung Lee4, Alessia Nasca5, Reza Maroofian5, Isabelle Schrauwen6, Manuela Pendziwiat7, Sunhee Jung8, Pasquale Striano9, Kshitij Mankad10, Barbara Vona11, Sanmati Cuddapah12, Sriniyana Gannavarapu13, Costanza Lamperti14, Andrea Legati15, Vincenzo Salpietro16, Sarah von Spiczak17, Abigail Sandoval18, Massimo Zeviani19, Adi Reich15, Cholsoon Jang9, Ingo Helbig16,17, Tahsin Stefan Barakat18, Daniele Ghezzi15-19, Suzanne M Leal6, Yvonne Weber20, Henry Houlden2, Wan Hein Yoon1 1) Oklahoma Medical Research Foundation, OKC, OK, USA; 2) UCL Queen Square Institute of Neurology, London, UK; 3) University of Tübingen, Tübingen, Germany; 4) Baylor College of Medicine, Houston, TX, USA; 5) Fondazione IRCCS Istituto Neurologico “Carlo Besta”, Milan, Italy; 6) Columbia University Medical Center, New York, NY, USA; 7) Christian-Albrechts-University of Kiel, Kiel, Germany; 8) University of California Irvine, Irvine, CA, USA; 9) University of Genoa, Genoa, Italy; 10) Great Ormond Street Hospital for Children, London, UK; 11) Eberhard Karls University, Tübingen, Germany; 12) Children’s Hospital of Philadelphia, Philadelphia, PA, USA; 13) Western University, London, ON, Canada; 14) University of Padova, Padova, Italy; 15) GeneDx, Gaithersburg, MD, USA; 16) Children’s Hospital of Philadelphia; 17) University of Pennsylvania, Perelman School of Medicine, Philadelphia, PA, USA; 18) Erasmus University Medical Center, Rotterdam, Netherlands; 19) University of Milan, Milan, Italy; 20) University of Aachen, Aachen, Germany

The 2-oxoglutarate dehydrogenase-like protein (OGDHL) is a rate-limiting enzyme in the Krebs cycle that plays a pivotal role in mitochondrial metabolism. OGDHL expression is restricted mainly to the brain in humans. Here, we report nine individuals from eight unrelated families carrying biallelic variants in OGDHL with a range of neurological and neurodevelopmental phenotypes including epilepsy, hearing loss, visual impairment, gait ataxia, microcephaly and hypoplastic corpus callosum. The variants include three homozygous missense variants (p.Pro852Ala, p.Arg444Trp, p.Arg299Gly), three compound heterozygous single nucleotide variants (p.Arg673Gln/p.Val488Val, p.Phe734Ser/p.Ala327Val, p.Trp220Cys/p.Asp491Val), one homozygous frameshift variant (p.Cys553Leufs*16), and one homozygous stopgain variant (p.Arg440Ter). To support the pathogenicity of the variants, we developed a novel CRISPR/Cas9-mediated tissue-specific knockout with CDS rescue system for dOgdh, the Drosophila ortholog of human OGDHL. Pan-neuronal knockout of dOgdh led to developmental lethality, which was fully rescued by expression of wild-type dOgdh. Studies using the Drosophila system indicate that Arg673Gln, p.Phe734Ser, and p.Arg299Gly are severe loss-of-function alleles, leading to developmental lethality, whereas p.Pro852Ala, p.Ala327Val, p.Trp220Cys, p.Asp491Val, and p.Arg444Trp are hypomorphic alleles, causing behavioral defects. Transcript analysis from fibroblasts obtained from individual carrying the synonymous variant (c.1464T>C:[p.Val488Val]) in family 2 showed that the synonymous variant
affects splicing of exon 11 in OGDHL. Human neuronal cells with OGDHL knockout exhibited defects in mitochondrial respiration, indicating the essential role of OGDHL in mitochondrial metabolism in humans. Together, our data establish that the biallelic variants in OGDHL are pathogenic, leading to a Mendelian neurodevelopmental disease in humans.

955B Proteolytic activity via the ubiquitin-proteasome system (UPS) serves a crucial role in regulating proteins for degradation and redistribution in the cell. Several human disorders directly result from dysregulation of the UPS. Mutations that affect expression levels of the ubiquitin protein ligase E3A (UBE3A/E6AP) can result in DuplicatioN 15q or Angelman syndrome. While many researchers have paired cell culture methods with immunoprecipitation and blotting techniques to identify UBE3A interactors, to date, only a handful of validated UBE3A substrates have been identified. Through the implementation of a new biochemical method known as a ubiquitin activated interaction trap (UBAIT), we can now covalently link ubiquitin substrates to UBE3A in vivo. We generated two upstream activator sequence (UAS) lines that incorporate a 6x polyhistidine-tagged Dube3a UBAIT construct for expression experiments in all cells (Actin-GAL4), glial cells (Repo-GAL4), and neurons (elav-GAL4). The first UBAIT is a K48R wild-type ubiquitin line that actively binds ubiquitin substrates and covalently attaches them to His-tagged Dube3a for isolation. The second UBAIT employs a non-functional ubiquitin with a DGG mutation that cannot covalently attach substrates to His-tagged Dube3a, but can serve as a control for other proteins stuck to the expressed UBAITs. Using Ni-NTA column purification combined with unbiased whole proteome analysis we identified 27 proteins using actin-GAL4 and 59 proteins using repo-GAL4 that appear to be bonafide Dube3a substrates and were not present in the DGG control UBAIT group. These hits included the Na/K pump ATPalpha, which our group previously showed is regulated by Dube3a in glial cells. Other hits include trehalose transporter 1-1 (Tret1-1), wings up A (wupA), and rolled (rl). This work marks the first successful implementation of the UBAIT system in a model organism, opening room for a multitude of successive in vivo UBE3A proteomic experiments.

955C Proteomic characterization of Dube3a substrates in glia versus neurons using ubiquitin activated interaction trap (UBAIT) Benjamin Geier, David Kaknijashvili, Daniel Johnson, Jon Huibregtse, Lawrence Reiter University of Tennessee Health Science Center; 2) University of Texas at Austin

Proteolytic activity via the ubiquitin-proteasome system (UPS) serves a crucial role in regulating proteins for degradation and redistribution in the cell. Several human disorders directly result from dysregulation of the UPS. Mutations that affect expression levels of the ubiquitin protein ligase E3A (UBE3A/E6AP) can result in DuplicatioN 15q or Angelman syndrome. While many researchers have paired cell culture methods with immunoprecipitation and blotting techniques to identify UBE3A interactors, to date, only a handful of validated UBE3A substrates have been identified. Through the implementation of a new biochemical method known as a ubiquitin activated interaction trap (UBAIT), we can now covalently link ubiquitin substrates to UBE3A in vivo. We generated two upstream activator sequence (UAS) lines that incorporate a 6x polyhistidine-tagged Dube3a UBAIT construct for expression experiments in all cells (Actin-GAL4), glial cells (Repo-GAL4), and neurons (elav-GAL4). The first UBAIT is a K48R wild-type ubiquitin line that actively binds ubiquitin substrates and covalently attaches them to His-tagged Dube3a for isolation. The second UBAIT employs a non-functional ubiquitin with a DGG mutation that cannot covalently attach substrates to His-tagged Dube3a, but can serve as a control for other proteins stuck to the expressed UBAITs. Using Ni-NTA column purification combined with unbiased whole proteome analysis we identified 27 proteins using actin-GAL4 and 59 proteins using repo-GAL4 that appear to be bonafide Dube3a substrates and were not present in the DGG control UBAIT group. These hits included the Na/K pump ATPalpha, which our group previously showed is regulated by Dube3a in glial cells. Other hits include trehalose transporter 1-1 (Tret1-1), wings up A (wupA), and rolled (rl). This work marks the first successful implementation of the UBAIT system in a model organism, opening room for a multitude of successive in vivo UBE3A proteomic experiments.

956A Sex and reproductive differences in intestinal tumours Emily Strachan, Irene Miguel-Aliaga, Susumu Hirabayashi, 1) MRC London Institute of Medical Sciences, UK; 2) Institute of Clinical Sciences, Faculty of Medicine, Imperial College London, London, UK

Cancer risk, type and progression varies with sex, due to a range of genetic, hormonal and behavioural factors. Post pregnancy this is modulated further due to changes in hormones, growth factors and the immune system.

The Drosophila midgut, analogous to the human small intestine, is a highly sexually dimorphic organ. As well as changes in metabolism, size and shape of the midgut, the intestinal stem cells (ISCs) that replenish the intestinal epithelium of adult flies divide more in females than males. After mating the female ISCs proliferate further: the midgut undergoes extensive remodelling, changing its lipid metabolism and increasing midgut size to prepare for energy-intensive egg laying.

It has been previously reported that females produce more proliferative Notch and Ras12VAPC/ midgut tumours than males. We have confirmed this sexual dimorphism is also present in Hippo pathway tumours induced by activation of Yorkie (Yki12V) or downregulation of Warts (Wts800).

Both Notch and Ras12VAPC1 tumours show an enhancement in proliferation after mating in females, reflecting the mated females larger and more proliferative midguts. Unexpectedly, two genetically distinct Hippo pathway tumours (Yki80A and Wts800) instead have (i) unusually high proliferation in virgin females, and (ii) lack of enhanced proliferation in response to mating. This suggests that differences in Hippo signalling may normally contribute to the reproductive differences in intestinal proliferation.

This demonstrates that different midgut tumour models respond differently to sex and reproductive status. Next we will explore the underlying molecular mechanisms, focusing on the Hippo pathway and metabolism more broadly.

957B Imbalances in active and repressive chromatin states underlie phenotypes caused by the oncoproteins H3 K27M and EZHIP

Sam Krabbenhoft, Tyler Masuda, Truman Do, Siddhant Jain, Peter Lewis, Melissa Harrison University of Wisconsin-Madison

Central nervous system (CNS) tumours are the leading cause of solid tumor death in children. Among the deadliest and most common pediatric brain tumours are diffuse intrinsic pontine glioma (DIPG) and posterior fossa ependymoma type A (PFA). Two key molecular events drive nearly all cases of these two cancers: a lysine-to-methionine mutation at residue 27 on histone H3 (H3 K27M) in DIPG or elevated expression of the previously uncharacterized protein EZHIP in PFA. These tumors share striking similarities, including a near-complete loss of histone H3 trimethylation at lysine 27 (H3K27me3), a mark that contributes to transcriptionally silent chromatin. We have previously shown that H3 K27M and
Genetic mutations are the major driver of cancer development. Aberrations within the EGFR-Ras-MAPK signalling pathway are the most frequently observed in human cancer, making this pathway a good target for therapy. Another signalling pathway that is gaining increasing interest is the Hippo signalling pathway, due to its involvement in cancer. The Drosophila imaginal discs have been extensively used as a model to study patterns of cancer growth, as most cancers arise from the epithelial tissue. Using Drosophila imaginal discs, we have discovered that the output of the Ras signalling pathway in promoting proliferation versus differentiation is under the tight control of the Hippo pathway. This study identified a cluster of 26 genes that are only upregulated with simultaneous dysregulation of the EGFR and Hippo signalling pathways and are largely unaffected with a mutation in a single pathway. Here we used a Gal4/UAS system to drive Ras or Raf gain of function combined with Yki upregulation using an optogenetic driver to generate our cancer model. This leads to a dramatic overgrown phenotype of the wing imaginal disc compared to wild type discs and extends the larval developmental stage up to 9 days. We have used this model to perform an RNAi screening of the aforementioned 26 genes and have identified that downregulation of negative regulators of the EGFR signalling pathway led to a rescue of the observed overgrown phenotype. Between these screening hits, we are focusing our investigation on the gene sprouty. This is a negative regulator of the EGFR signalling pathway whose function as either a tumour suppressor or oncogene in cancer has shown conflicting findings. Upregulation of sprouty 2 (the human ortholog of Drosophila sprouty) is common in colorectal cancer where dysregulation of both the Hippo and EGFR pathways are reported. Several studies have shown that downregulation of sprouty 2 using siRNA led to decreased cancer proliferation and invasion. The similarity to our in vivo system has led us to investigate this gene further. Coupling an in vivo (Drosophila melanogaster) and in vitro model (colorectal cancer cell lines), we are investigating whether Sprotuy can be a better target for therapy in those cancers where dysregulation of this gene could lead to a worse outcome.

Salt-inducible kinases synergise with Homeodomain-interacting protein kinases to promote significant tumour growth Kewei Yu, Nivi Ramkumar, Esther Verheyen Simon Fraser University, Burnaby, British Columbia, Canada

Homeodomain-interacting protein kinases (HIPks) are conserved kinases that regulate cell proliferation, apoptosis and tissue development. HIPks have been shown to regulate the activity of many key conserved signaling pathways, including Wnt, Notch, Hippo and Jak/Stat. Overexpression of HIPk in Drosophila causes tumorigenic phenotypes. We primarily examine tissue overgrowth and distortion in larval imaginal discs. We found that co-expression of a constitutively active form of Salt-inducible kinase 2 (Sik2) or Sik3 with HIPk caused significant tissue overgrowth and tissue distortion, indicating that both Sik2 and Sik3 can synergize with HIPk to promote tumour growth. Larvae expressing these neoplastic growths also display an extended larval phase, characteristic of Drosophila tumor models. SIKs are serine/threonine protein kinases belonging to the adenosine monophosphate (AMP)-activated protein kinase (AMPK) family. While mammals have SIK 1-3, Drosophila only has Sik2 (mammalian ortholog of SIK1 and SIK2) and Sik3. Furthermore, depletion of Sik 2 or Sik3 can suppress overgrowth phenotypes due to HIPk overexpression. Examination of total protein levels from fly tissues showed that HIPk proteins were reduced when SIKs were depleted through RNAi or by using mutant alleles. These results suggest that SIKs may regulate HIPk protein stability and/or activity. Furthermore, SIKs are known to regulate several signal transduction cascades by phosphorylating key components of these pathways. Preliminary data from the lab has shown that Sik2 can be co-immunoprecipitated with HIPk. In summary, our research demonstrates a novel function of SIKs in synergizing with HIPk to promote tumour growth potentially. We are investigating the mechanism underlying these interactions and hypothesize that SIKs phosphorylate HIPks to modulate their stability and or activity. Gaining further insight into tumorigenic synergies has potential implications of finding drug targets in the future that could block tumour growth via inhibition of SIKs.
Myocardial infarction remains the leading cause of death and myocardial ischemia contributes to 2/3rds of all cases of heart failure, which is rapidly emerging among the leading causes of mortality and morbidity in the U.S. *Drosophila melanogaster* is a simple and powerful genetic model system to investigate the role of genes associated with human diseases, including cardiac diseases. As short-lived animals, flies represent an excellent opportunity to model age or disease dependent changes in cardiac function that can be traced throughout life. The fly’s heart tube is located on the dorsal side of its abdomen within 200 µm from the tissue surface allowing visible to near-infrared light to reach the heart tube. This enables non-invasive optical pacing of the fly heart using existing optogenetic tools. Morphological and rhythmic changes in a relatively simple fly heart can be readily analyzed with a non-invasive biomedical imaging technology, Optical Coherence Tomography (OCT). We developed transgenic system consisting of spatiotemporally regulated ChR2, ReaChR, and eNpHR2.0 opsins in *Drosophila*. We have demonstrated the tissue specific opsin expression and performed the heart pacing experiments. The next step will enable us to characterize changes of the fly heart function in response to different stress challenges providing insights into molecular mechanisms of ischemic preconditioning.

**961C Optogenetic control of *Drosophila* cardiac function with ChRmine and ReaChR opsins Fei Wang, Elena Gracheva, Abby Matt, Hongwu Liang, Matthew Fishman, Chao Zhou Washington University in St. Louis, Saint Louis, MO**

Cardiac optogenetics is a promising alternative to traditional electrical stimulations in controlling the activity of cardiac tissues non-invasively. In recent years, cardiac functions of animals such as rats, zebrafish, and fruit fly, have been controlled by opsin activation. In this study, we expressed red-shifted excitatory opsins (ReaChR and ChRmine) in the heart of *Drosophila melanogaster*. For ReaChR and ChRmine, we used red-light stimulation for deep penetration into the myocardial structures. Compared to existing opsins, the newly generated opsin ChRmine is able to induce spiking with much lower irradiance power. M-mode images acquired with custom optical coherence microscopy (OCM) system demonstrated controlled heart function in vivo and in real-time throughout the *Drosophila* life cycle (larva, early pupa, late pupa, and adult). Fast kinetics, low stimulation power, and broad heart-rate adjustable range were demonstrated for light pulses pacing. *Drosophila* heart rate can be enhanced to the frequency of the light excitation, faster than its resting heart rate, and reversible acceleration of heartbeats can be achieved at various developmental stages. This study demonstrated non-invasive cardiac control through promoting heart functions of an intact animal, which is promising for the nondestructive studies of cardiac diseases, such as congenital heart disease, posteriority bradycardia, tachycardia, and regional mechanical dys-synchrony.

**962A Mitochondria malfunction and RNaseZ-associated cardiomyopathy Ekaterina Migunova1, Stefania Bonanni1, Megan Kurz2, Alessia Vata1, Hongwu Liang2, Fei Wang2, Chao Zhou2, Edward Dubrovsky1 1) Fordham University, Bronx, NY; 2) Washington University, St Louis, MO**

Cardiomyopathy (CM) is a disease of heart muscle leading to changes in heart wall morphology, reduced heart contractility and obstruction of blood flow. It affects people of all ages; the adult form of CM is often secondary to systemic causes such as inflammation, malnutrition and other conditions, while infantile CM occurs due to inherited mutations. Mutations of ELAC2/RNaseZ gene, which encodes RNaseZ enzyme, lead to an especially aggressive form of HCM with early onset of symptoms and lethal outcome within one year after birth. RNaseZ is a highly conserved vital enzyme with homologs in all eukaryotes. It is essential for tRNA maturation and therefore for protein synthesis. It has a dual role within a cell as it is the only enzyme that processes 3’-ends of pre-tRNA in both nucleus and mitochondria. Previously we have generated a novel fly model of RNaseZ linked CM and demonstrated that it mimics the human disease in all major symptoms. Further investigation of the underlying mechanisms leading to RNaseZ linked abnormal heart morphology is challenged by the dual role of RNaseZ within a cell. Here we report the effect of CM-linked RNaseZ mutations on mitochondria function. We have found that RNaseZ mutations cause a decrease in mitochondrial complex I activity, which leads to a reduction in mitochondrial ATP synthesis rates. We also have separated the nuclear and mitochondrial roles of RNaseZ to investigate the contribution of each towards the CM phenotype. We have discovered that mutating only mitochondrial RNaseZ is sufficient to cause changes in heart morphology. Lastly, we investigated whether the heart damage originating from RNaseZ dysfunction in mitochondria is cardiac specific or if it is secondary to a systemic disorder, and found that the cardiac damage associated with mutant mitochondrial RNaseZ is heart specific. Overall, these findings improve our understanding of the cellular processes contributing to RNaseZ linked CM.

**963B A *Drosophila* model for human ARVC-5 caused by TMEM43<sup>S358L</sup> Nora Klinke1, Sandra Ratnavadivel2, Marcel Reinhardt1, Heiko Meyer1, Anders Malmendal1, Hendrik Milting1, Achim Paululat1 1) University of Osnabrueck, Department of Zoology and Developmental Genetics (Germany); 2) Erich and Hanna Klessmann Institute, Heart and Diabetes Center NRW, University Hospital of the Ruhr-University Bochum, Bad Oeynhausen (Germany); 3) Roskilde University, Department of Science and Environment (Denmark)**

Human *TMEM43* encodes a membrane protein with four transmembrane domains, which localizes to the ER- outer membrane compartment. A point mutation in TMEM43 results in arrhythmogenic right ventricular cardiomyopathy
(ARVC) type 5. Characteristics of ARVC-5 are ventricular tachycardia, heart attacks, and sudden cardiac death. Mostly, men between the ages of 20-40 years are affected. The identified point mutation is a missense mutation in the human TMEM43 gene that results in incorporation of the amino acid leucine rather than serine at position 358 of the protein, S338L. TMEM43 is a highly conserved protein throughout different species. Homologous proteins exist in chimpanzees, rhesus monkeys, dogs, cows, mice, frogs and fruit flies. The corresponding homolog in Drosophila melanogaster is CG8111. Interestingly, the serine, which is involved in the familial mutation in humans, as well as the surrounding region, are highly conserved in fruit flies (S333) and other species.

To investigate the effect of the point mutation in flies, we overexpressed CG8111S338L ubiquitously or in a tissue specific manner. Our studies showed that ubiquitous overexpression of the mutant, but not the wild type form of CG811, leads to lethality at pupal stages, less food uptake and increased lipid droplets in adipocytes of 3rd larvae. Furthermore, we applied mass spectrometry (MS)-based proteomics and are able to show that CG8111 S338L animals up-regulate proteins, which are involved in fatty acid metabolism pathways. Subsequent metabolomic analysis by nuclear magnetic resonance (NMR) spectroscopy revealed that short fatty acids accumulate in animals overexpressing CG8111S338L compared to controls.

Moreover, we examined the effect of heart specific overexpression of CG8111S338L. Therefore, we used the Semi-automatic Optical Heartbeat Analysis (SOHA) software to analyse heart beat parameters like heart rate, systolic and diastolic interval, fractional shortening and arrhythmicity index. We found that heart specific overexpression of CG8111S338L leads to arrhythmia in five week old male flies.

In the end, we compare our findings with models for TMEM43S358L in mouse and rat.

1 Baskin et al., Human Genetics, 2013
2 Christensen et al., Clinical Genetics, 2011
3 Milting et al., European Heart Journal, 2015
4 Fontaine et al., Circulation, 1998
5 Merner et al., The American Journal of Human Genetics, 2008

964C New genetic avenues in Congenital Heart Disease: Ribosomal protein genes as regulators of cardiac growth (via YAP/yorkie) and proliferation (via p53) along with cardiogenic transcription factors Tanja Nielsen4, Anais Kervadec2, Xin-Xin I. Zeng1, Analyne Schroeder1, Jeanne L. Theis1, Timothy M. Olson3, Karen Ocorr1, Paul Grossfeld4, Alexandre R. Colas2, Georg Vogler1, Rolf Bodmer1 1) Development, Aging and Regeneration Program, Sanford Burnham Prebys Medical Discovery Institute, La Jolla, CA, USA; 2) Human Genetics Program, Sanford Burnham Prebys Medical Discovery Institute, La Jolla, CA, USA; 3) Division of Pediatric Cardiology, Department of Pediatric and Adolescent Medicine, Mayo Clinic, Rochester, MN, USA.; 4) Department of Pediatrics, UCSD School of Medicine, La Jolla, CA, USA

Hypoplastic Left Heart Syndrome (HLHS) represents the most lethal Congenital Heart Disease (CHD) and is characterized by a severely underdeveloped left ventricle. The genetic mechanisms leading to HLHS are poorly understood, but likely of oligogenic origin.

Decreasing sequencing costs have led to the identification of thousands of putative human disease variants. However, establishing genotype-phenotype relationships remains challenging. In vivo functional analysis of patient-derived genes and gene clusters in a high-throughput manner using the fly heart model combined with human iPSC-cardiomyocytes (hiPSC-CMs) and zebrafish, provides a framework for prioritizing and interrogating the contribution of genetic variants in heart disease.

We performed whole genome sequencing in a familial HLHS case and a cohort of 25 HLHS proband-parent trios with poor clinical outcome. Candidate genes identified by Mendelian modeling in the 25 trios were subject to gene network enrichment analysis, revealing an over-representation of ribosomal protein (RP) genes. Segregation analysis in the familial case identified a rare promoter variant affecting RPS15A. Remarkably, patient-derived iPSC-CM proliferation was reduced compared to the parents.

Functional analysis in model systems showed that knockdown (KD) of RPs reduced proliferation in generic hiPSC-CMs and impaired cardiac differentiation in Drosophila resulting in a partial or ‘no’ heart phenotype in adult flies. Furthermore, Rps15Aa KD led to cardioblast misspecification during early cardiogenesis. Functional validation in zebrafish revealed that rps15Aa KD causes reduced cardiomyocyte numbers and contractility, and defective heart looping, without affecting overall embryonic development. Strikingly, RPS15A KD-induced defects were significantly reversed by p53 KD in hiPSC-CMs or zebrafish, and by myc KD or YAP/yorkie overexpression in flies, dependent on yorkie’s co-factor TEAD/scalloped.

When testing for cardiac-specific RP functions, we found synergistic interactions between RPS15A and cardiac transcription factor tinman/Nkx2.7 and dorsocross/Tbx5a, in both Drosophila and zebrafish suggesting conserved synergism between RPs and cardiogenic genes in both models.

In summary, we conclude that RP genes play a critical role as regulators of cardiac growth and cardiomyocyte proliferation along with cardiogenic transcription factors and we suggest that RP genes represent a novel class of genetic effectors in CHDs, such as HLHS.
**965A** Characterizing Robinow Syndrome-associated DVL1 mutations in Drosophila Katja MacCharles1, Sarah Gignac2, Katherine Fu2, Joy Richman2, Esther Verheyen1 1) Simon Fraser University; 2) University of British Columbia

Human development is regulated by intricate, and interconnected, signal transduction networks. Given the complexity, deciphering the effects of mutations that give rise to abnormal development can be challenging. Using Drosophila melanogaster can simplify the puzzle of studying human disorders as flies have little genetic redundancy and are significantly easier, cheaper, and faster to raise than other vertebrate models. I use Drosophila to characterize Dishevelled1 (DVL1) mutations obtained from patients with Robinow Syndrome (RS). RS is a rare genetic disorder associated craniofacial abnormalities and shortened stature. Most of the mutations associated with RS affect components of the non-canonical/Planar Cell Polarity (PCP) pathway of Wnt signaling. Wnt signaling is involved in embryonic development and homeostasis. The two main pathways, canonical and non-canonical/PCP Wnt signaling, require DVL but there is still much to learn about PCP signaling which mediates cytoskeletal rearrangement events and orient cell polarity within the epithelial plane. There are 3 DVL proteins found in vertebrates, and a single ortholog, Dsh, in Drosophila. Each of the DVL1 variants I study have unique frameshift mutations that replace the highly conserved C-terminus with the same novel peptide sequence of no known homology. I use the Gal4-UAS system to express wildtype human DVL1 and three DVL1 variants in Drosophila. My research has shown that these DVL1 patient variants disrupt the stability of Armadillo/β-catenin, ectopically induce PCP/JNK signaling and activate apoptosis. Furthermore, the variants induce several novel phenotypes in wing tissue such as anterior cross vein abnormalities, ectopic bristles, and vein thickening, suggesting novel functions in other conserved signaling pathways. By understanding how conserved signaling pathways are altered by these DVL1 variants, we gain insight into the underlying mechanisms of non-canonical Wnt signaling and more broadly, how development in individuals with RS is altered. This information may guide future therapeutics for RS patients.

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**966B** The Drosophila ortholog of POLR1D, an RNA Polymerase I & III assembly protein, is required for development Ryan Palumbo, Alana Belkevich, Bruce Knutson SUNY Upstate Medical University

rRNA expression and ribosome assembly are becoming appreciated as processes required for proper development. Defects in rRNA synthesis/processing, ribosomal protein expression, and ribosome assembly, all constitute a class of diseases referred to as ribosomopathies. Ribosomopathies are characterized by a tissue-specific requirement for sufficient translational output not being satisfied due to mutations in genes ultimately involved in translation. Several animal models of ribosomopathies have been established to study the molecular and physiological consequences of those mutations. Drosophila is emerging as an excellent model system to study ribosomopathies in various developmental contexts. Here, we establish Drosophila as a model to study the ribosomopathy Treacher Collins Syndrome (TCS). TCS is caused by mutations in several genes, one of which is POLR1D. POLR1D protein forms a heterodimer with POLR1C, which is a prerequisite to the assembly of RNA Pol I and Pol III, which synthesize 47S and 55S rRNAs, respectively. Loss of POLR1D function is thought to affect polymerase assembly, reduce the production of rRNAs, and impair ribosome assembly, which ultimately affects translation. We found that flies hemizygous for an allele of Drosophila POLR1D (dPOLR1D) are pre-adult lethal, and viable larvae exhibit reduced growth, and are arrested in the L1 and L2 instar stages. dPOLR1D mutant larvae have reduced levels of 18S and 28S rRNAs, and this is reflected in heterozygous dPOLR1D adults, which exhibit the bobbed phenotype characteristic of mutants lacking ample rRNA production. Thus, dPOLR1D is required for rRNA expression, which is necessary to complete larval development. We sequenced this allele of dPOLR1D, and identified a missense mutation causing a substitution in an evolutionarily conserved amino acid that has been found to be mutated to a different residue in a single case of non-familial TCS. We found that the “fly mutation” in both dPOLR1D and human POLR1D (hPOLR1D) does not affect binding of POLR1D proteins to their cognate POLR1C proteins; however, the “human mutation” does. This suggests that while different amino acid mutations can both affect development, the molecular mechanism by which this occurs can be different. TCS is a disease of neural crest cells. We performed a preliminary RNAi screen in several cells types of the nervous system, and found that as in humans, dPOLR1D is required in the nervous system to complete development.

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**967C** A novel assay to study salivary gland dysfunction in a model of NGLY1 deficiency John Pleinis2, Kevin Hope1, Danielle DaCrema3, Clement Chow2, Aylin Rodan2 1) Department of Human Genetics, University of Utah School of Medicine, Salt Lake City, UT; 2) Department of Internal Medicine, Division of Nephrology and Hypertension, and Molecular Medicine Program, University of Utah School of Medicine, Salt Lake City, UT

NGLY1 deficiency is a rare disorder caused by loss of function mutations in the NGLY1 gene and often presents with developmental delay, liver dysfunction, and seizures. One of the most consistent and defining disease phenotypes is the reduced ability to make tears, sweat, and saliva. Therapies that could treat these secretory problems could have a profound effect on quality of life for these patients. To study salivary gland dysfunction in adult flies, we developed a “rock candy” assay, which measures the consumption of a solid blue sugar. Consumption, as measured by blue dye, is correlated with salivary gland function. Ablation of the salivary gland results in little to no consumption of the blue sugar. To model NGLY1-induced salivary gland dysfunction, we used RNAi to knockdown NGLY1 (Pgl) in flies) expression in the salivary glands of adult flies and found that this reduces consumption by nearly 50%. We previously showed that
the Na-K-Cl cotransporter, NKCC1 (Ncc69 in flies), is an enzymatic target of NGLY1 and a modifier of NGLY1 deficiency in flies. NKCC1 regulates ion flux in secretory cells like those in the salivary gland. We generated salivary gland-specific knockdown of NKCC1 and found a severe consumption defect. To test whether the defects we observed in either single knockdown was due to a genetic interaction, we generated salivary gland-specific double knockdown. The double knockdown had a severe consumption defect similar to what we observed in the single NKCC1 knockdown, suggesting that the defects observed are due to a genetic interaction between NGLY1 and NKCC1. We also demonstrate that drugs that alter NGLY1 or NKCC1 activity can rescue the consumption phenotype in NGLY1 knockdown. Together, these data indicate that the fly is a powerful model for studying salivary gland dysfunction in NGLY1 deficiency and may help develop life changing therapies.

968A  

De novo variant in MRTF-B is associated with intellectual disability, minor dysmorphic features, expressive language delay, impulse control issues, and fine motor delay. Jonathan Andrews1, Oguz Kanca1, David Li-Kroeger1, Cyndi Tifft2, Ellen Macnamara3, Shinya Yamamoto1, Hugo Bellen1, May Malicdan1, Michael Wangler1 1) Baylor College of Medicine; 2) NIH, Bethesda, MD; 3) NIH/NHGRI, Arlington, VA; 4) NIH/NHGRI, Rockville, MD

Myocardin-related transcription factor B (MRTF-B) is a member of a family of genes which serve to potentiate serum response factor (SRF)-dependent transcription and is highly conserved in both vertebrate and invertebrate model organisms. The MRTF-B protein is not currently associated with a human disease but has been shown to be highly expressed in all human tissues save the lung. Here we report a proband with a de novo variant in the second RPEL domain of MRTF-B with intellectual disability, minor dysmorphic features, expressive language delay, impulse control issues, and fine motor delay. We have generated a fly model of human MRTF-B, using a UAS construct carrying either the human reference or variant cDNA. Expression of the UAS-human variant cDNA under the control of the Nubbin-Gal4 driver was sufficient to induce significant morphological changes in the wing, while the expression of the human reference cDNA produced only minor changes in the posterior crossvein. Expression of Drosophila Mrtf using the Nubbin-Gal4 driver produced a similar change in crossvein length as was observed with the human reference. In Drosophila, the SRF ortholog is known as blistered (bs) and is known to suppress wing vein formation and promote the development of intervein cells. As exogenous co-expression of bs and Mrtf has been previously shown to significantly alter wing morphology, we also expressed our human reference and variant cDNA lines concurrently with a UAS-bs line. In these experiments we found that wing morphology was highly disrupted when bs and the reference human cDNA were co-expressed, while the co-expression of human variant cDNA and bs was lethal. Based on these findings, we sought and identified two additional probes with variants within or near the second RPEL domain. Expression of these additional variants using Nubbin-Gal4 produced changes in the posterior crossvein analogous to the changes observed when our human reference line was expressed. Taken together, these findings suggest that different residue changes within the RPEL domain of MRTF-B can have drastically different morphological effects in the fly wing, and our initial de novo variant may underlie a novel disorder.

969B  

Identification of gene expression changes in response to vitamin A deprivation Deepshe Dewett, Maryam Labaf, Kanh Lam-Knath, Kourosh Zarrinhalam, Jens Rister UMass Boston

Chronic vitamin A deficiency (VAD) leads to blindness, delayed growth, impaired immunity and is the leading cause of preventable childhood blindness. In contrast, VAD in flies does not result in developmental defects or lethality. But similar to mammals, flies deprived of vitamin A have defects in the visual system, like damaged photoreceptors and loss of light-sensing pigments. However, a gaping hole remains in our understanding of the molecular response to VAD. To identify genes and signaling pathways that respond to VAD, we performed RNA sequencing using Drosophila heads. We identified 68 differentially expressed genes between vitamin A-rich and deprived conditions. Out of these, 50 genes were upregulated, and 18 genes were downregulated on VAD. Among these are the genes that are responsible for the synthesis of the retinal chromophore (ninaB, ninaG, Pdh) and the termination of the phototransduction pathway (Arr1, Arr2, inac, stops). We also detected significant changes in the expression of genes that encode for specific aminoacyl-tRNA synthetases (GluProRS, IleRS, LeuRS, LysRS), major nutrient reservoir proteins (Lsp1alpha, Lsp1beta, and Lsp2), calcium buffers (Cpn, Cnx99a), and factors that mediate stress or immune responses (per, Cyp309a1, TotA, TotC, TotM, dnr1, Diedel).

A comparison between genes responding to VAD and previously identified blue light stress-responsive genes showed a largely nonoverlapping transcriptome response to these two types of environmental stresses. Only seven genes, with poorly understood functions, were similar between these two studies. Taken together, our study provides insights into the molecular mechanisms that respond to vitamin A deprivation.

970C  

Transcriptomic analysis in NF1: exploring drivers of diverse phenotypes Connor N. Broyles, Valentina Botero, Seth M. Tomchik Department of Neuroscience, The Scripps Research Institute, Scripps Florida, Jupiter, FL, USA.

Neurofibromatosis type 1 (NF1) is a chronic multisystem genetic disorder affecting approximately 1 in 3,500 humans. NF1 patients often experience multiple symptoms such as tumors, cognitive dysfunction, sleep disturbances, and metabolic changes. NF1 is caused by mutations in the NF1 gene, which encodes a protein called neurofibromin. One of the major
biochemical functions of neurofibromin is Ras GAP activity, which downregulates Ras signaling. Loss of function of the catalytic neurofibromin GAP-related domain, which is responsible for neurofibromin’s Ras GAP activity, is a major contributor to NF1 phenotypes such as increased energy expenditure. In addition, Ras-independent effects on other signaling pathways, such as cAMP/PKA, have been described for phenotypes like short stature and sleep disruption. The impacts on relatively pleiotropic cellular signaling pathways, as well as the systemic interactions among different cell types, raises the question of how loss of neurofibromin drives systemic and organismal pathophysiology in NF1. Downstream molecular pathways remain elusive. This study aims to identify key pathways involved in NF1 pathology using a transcriptomics approach.

Loss of neurofibromin may drive effects on organismal physiology via alterations in the transcription of downstream gene products. To gain insight into the transcriptional landscape in NF1, we have examined the effects of loss of neurofibromin on gene expression in Drosophila. RNA sequencing was used to identify differentially expressed genes (DEGs) in NF1 mutants (NF1flh) as compared to wild-type controls (w1118). RNA from heads and bodies was compared between mutants and controls to identify DEGs in neurally-enriched samples vs. the rest of the body. Analyses revealed several classes of dysregulated transcripts in NF1 mutants; specific for heads/bodies or shared in both. Gene ontology analysis revealed several enriched processes and pathways in NF1 mutants, with association to known human phenotypes, and follow-up genetic experiments are narrowing key pathways for NF1 effects on organismal phenotypes.

971A CryAB is a target protein of NUAK kinase activity to prevent protein aggregation in muscle tissue Ziwei Zhao, Erika Geisbrecht Kansas State University

NUAK belongs to the AMP-activated protein kinase (AMPK) family, which is comprised of conserved serine/threonine protein kinases known to regulate cell polarity or cell motility by controlling the assembly and/or disassembly of cytoskeletal proteins. Previous results in our lab have demonstrated that a mutation in Drosophila NUAK results in the degeneration of larval body wall muscles. One prominent feature in these muscles is the abnormal accumulation of select proteins, which displaces other components of the contractile sarcomere apparatus. To better understand how kinase activity contributes to NUAK function, we mutated important residues in the conserved kinase domain using CRISPR/Cas9 and transgenic mutagenesis approaches. Indeed, mutation of a key catalytic residue within the kinase domain resulted in earlier lethality and thinner muscles. These results were confirmed by expressing a salimius (Slis)-GFP fusion protein to visualize the dynamics of this process in vivo. Next we wanted to determine if we could separate the requirement for NUAK in embryonic muscle development from protein aggregation and muscle degeneration in larval contractile muscles. Compromising NUAK kinase activity in larval muscle using a post-embryonic driver still resulted in protein aggregation, thus suggesting that NUAK has independent functions in muscle development and muscle homeostasis. We have confirmed that a conserved threonine residue in the kinase domain of NUAK is essential to promote protein activity. To uncover proteins that may be substrates of NUAK kinase activity, we performed a yeast two-hybrid screen. I(2)elf, which encodes for CryAB, emerged as a top candidate since mutations in human CryAB/α-crystallin B cause a type of protein aggregate disease called Myofibrillar Myopathy. Our results thus far show that overexpression of NUAK increases CryAB phosphorylation and this post-translational modification can be reversed with phosphatase treatment. Taken together these data suggest that CryAB may be a target of NUAK phosphorylation in the prevention of protein aggregation.

972B Survival and motility of adult Drosophila melanogaster flies fed high-calorie diets during early development Mario Cayetano-Velazquez1, Irma Dueñas-García1, Luis Felipe Santos-Cruz1, Laura Castañeda-Partida1, Evelyn Hernández-Torres1, Elizabeth Soto-López1, Daniela Vargas-Sánchez1, Nefte Meni Yañez-Mendoza1, Maria Eugenia Heres-Pulido1, Noma Velazquez-Ulloa2 1) Lab. de Genética Toxicológica, FES Iztacala UNAM. Av. Los Barrios # 1. Los Reyes Iztacala, Tlalnepantla, Estado de México, CP 54090. ; 2) Lewis & Clark College

Excessive consumption of some macronutrients, such as lipids and carbohydrates can contribute to development of cardiovascular disease, neurodegenerative disease, obesity, and metabolic syndrome, among others. Drosophila melanogaster is an effective model organism to determine the effects of the consumption of hypercaloric diets during development because of the high conservation in the metabolic and signaling pathways between mammals and insects, in which many of the participating genes have orthologs in Drosophila. We evaluated the effect of different diets on survival and negative geotaxis in two Drosophila strains, Canton-S (CS, wildtype), and Oregon R(R)-flare (OR), which have increased Cyp450 enzyme levels and have been shown to be insecticide resistant. Larvae from these fly strains fed on either a Normal Diet (ND), a High Fructose Diet (FR; 17% fructose), a High Palmitic Acid Diet (AP, 3.2% palmitic acid), or a Mixed Diet (MX, 5% FR + 1% AP) until pupation. For the survival assays, ten recently emerged adult flies per vial, per treatment, per sex, per fly strain, were transferred to vials with ND food. Three vials per condition were collected per experiment, and five independent experiments were conducted. Flies were transferred to new vials and the number of survivors per vial was counted every 3 days for 54 days. To determine effects on survival, we estimated mean survival, time to 50% survival, maximum survival and performed a Kaplan-Meier survival analysis. We found that survival was significantly decreased by the hypercaloric diets. In the CS flies, all 3 high calorie diets decreased survival, while OR...
flies had decreased survival with the FR and AP diets, but not in the MX diet. In addition, the magnitude of the effect of the diets on survival was greater in the CS strain than in the OR strain, which suggests that increased P450 enzyme levels may confer protection. Next, we evaluated the effect of the different diets on negative geotaxis. For this assay, ten recently emerged flies per vial, per treatment, per sex, per fly strain, were transferred to vials and tested on a climbing assay. CS flies from the high calorie FR diet climbed the most, while OR flies that consumed the ND climbed the least. The high calorie treatments affect female and male flies of the CS and OR fly strains differently. We also noticed differences between these two strains, which support an involvement of the xenobiotic metabolism in the effects of high caloric diets.

973C Genotype-by-Sex-by-Exercise Studies Using Drosophila melanogaster: Comparing the Power Tower and the TreadWheel as Two Exercise Apparatuses Tolulope Kolapo¹, Jordan Albrecht², Michelle Tan³, Mckenzie Chamberlain⁴, Annie Backlund⁵, Alyssa Koehler⁶, Sean Shelley Tremblay⁷ 1) The University of Alabama, Tuscaloosa, AL; 2) The University of Alabama, Tuscaloosa, AL; 3) The University of Alabama, Tuscaloosa, AL; 4) The University of Alabama, Tuscaloosa, AL; 5) The University of Alabama, Tuscaloosa, AL; 6) The University of Alabama, Tuscaloosa, AL; 7) The University of Alabama, Tuscaloosa, AL

Exercise is a cost effective intervention strategy that has been recognized to play a role in prevention and corrective response to Metabolic Syndrome (Mets). Mets is a combination of metabolic disorders that are increasingly prevalent in Western cultures. Metabolic disorders associated with MetS include elevated blood pressure, increased blood-sugar levels, abnormal cholesterol and increased triglyceride concentrations. To determine the effect of exercise on obesity and other metabolic disorders, daily exercise regimes can be simulated using Drosophila melanogaster. The negative geotaxis; an innate escape response where flies ascend the wall of a cylinder when tapped to the bottom is a characteristic climbing tendency of the fruit fly that can be taken advantage of to induce exercise. The TreadWheel and Power Tower are two apparatuses that have been used to induce exercise in flies under controlled settings. For both pieces of equipment, significant effects of exercise on diet, weight, climbing ability, aging and other phenotypes have been reported in fruit flies. Although both machines allow for regulated induction of exercise, each one provides its own set of distinct advantages and limitations. Currently, no direct comparison of the effectiveness of the two pieces of equipment has been conducted. It is therefore unclear whether the two devices differ in their ability to induce exercise or in their impact on metabolic phenotypes. Our project compares stress levels and metabolic response induced in flies by the two apparatuses, and also determines the most effective apparatus to achieve optimum exercise in adult flies by evaluating genetic and sex-specific interactions driving variation in metabolic phenotypes. This study determines the response of seven wild derived genetic lines from the Drosophila Genetic Reference Panel to exercise treatment on both apparatuses. Fitness traits such as longevity, fecundity and climbing speed are measured. Our results show that flies exercised on both apparatuses varied in their response by genotypes and sex, and flies exercised on the TreadWheel lived longer after exercise than flies exercised on the Power Tower.

974A Pupation as a critical hypoxia-sensitive stage in Drosophila melanogaster Tsering Stobdan, Nicholas Wen, Dan Zhou, Gabriel Haddad University of California San Diego

Hypoxia plays a critical role in multiple disease conditions. The highly conserved cellular and molecular mechanisms between flies and higher eukaryotes, including humans, have made Drosophila a valuable organism to identify hypoxia-sensitive cells and genes critical to hypoxic environment. Since the life cycle of Drosophila constitutes four morphologically distinct developmental stages, a stage specific identification of mechanisms involving hypoxia response becomes critical. Although the effect of hypoxia on the embryo, larvae or adult is frequently reported, the effect on pupae or pupation is less clear. In this study, we examined different stages of Drosophila development for its hypoxia sensitivity and identified pupae as a critical stage affecting their subsequent eclosion. We show that the continuous exposure to 4% O₂ from embryo to pupae is lethal. However, returning to normoxia at early, mid and late pupae stage would result in 99.6%, 66.1% and 54.4% eclosion, respectively. Similarly, returning to normoxia at 1st, 2nd or 3rd instar larvae resulted in >97% eclosion. In contrast, exposing the normoxia-grown larvae or pupae to 4% O₂ is lethal. Additionally, exposing early pupae for 3 days of 4% O₂ led to only 6.1% eclosion, while shorter exposure such as for 2 days or 1 day to 4% O₂ resulted in significantly higher eclosion of 66.7% and 96.4%. These results indicate that pupation is a very sensitive stage to hypoxia during Drosophila development. Overall, these results will be the basis for the identification of critical genes involved in hypoxia sensitivity or tolerance during development.

975B Gapvd1 regulates slit diaphragm formation in Drosophila but is otherwise dispensable for fly development. Konrad Lang, Helena Heinkele, Julian Milosavljevic, Lea Gerstner, Tobias Hermle Renal Division, Faculty of Medicine and Medical Center, University of Freiburg, Freiburg, Germany

Introduction:
Mutations in the gene GAPVD1 cause nephrotic syndrome in humans. GAPVD1 interacts with the early endosomal regulator RAB5 but the subcellular localization of GAPVD1 remains unclear. Silencing of Gapvd1 in the podocyte-like Drosophila nephrocytes resulted in mistrafficking of fly nephrin.
Methods: We generated conditional knockdowns and a stable genetic deletion of *Drosophila Gapvd1* by CRISPR/Cas9 and used microhomology-mediated end joining to introduce a c-terminal HA-tag into the genomic locus of *Gapvd1*. We performed a functional analysis of the novel fly models.

Results:
Using coexpression of tandem *Gapvd1*-gRNAs and Cas9 in the germline we generated twin frameshift mutations at the second and third exons of the *Drosophila Gapvd1* gene. Flies carrying these mutations were homozygous viable without any overt phenotype. In contrast, the podocyte-like nephrocytes of these animals showed a severely altered slit diaphragm architecture with mislocalization of fly nephrin and the orthologue of NEPH1 as well as a partial loss of both proteins from the surface. This phenotype was similar but considerably stronger than the phenotype observed when using RNAi-mediated silencing. This suggests that the homozygous frameshift mutations result in a null allele. The phenotype was confirmed by conditional CRISPR/Cas-mediated knock down using two independent gRNAs. Deletion of *Gapvd1* in the *Drosophila* model thus results in a phenotype that manifests exclusively in podocyte-like nephrocytes. This recapitulates the disease manifestation in human patients with *GAPVD1* mutations who presented exclusively with nephrotic syndrome, supporting a role for *Drosophila* as a model for this genetic disease. To study the subcellular localization of *Drosophila* Gapvd1, we introduced an HA-tag into the C-terminus of the Gapvd1 locus. Immunofluorescence of nephrocytes derived from the resultant knock-in lines showed co-localization of the Gapvd1-HA protein with *Drosophila* Rab5, supporting that *Gapvd1* primarily resides in early endosomes. We overexpressed human *GAPVD1* in nephrocytes, that equally localized in early endosomes. Gain-of-function of the human gene entailed reduced tracer endocytosis in nephrocytes, suggesting a dominant negative effect.

Conclusion: We established suitable new *Drosophila* models to study the function of Gapvd1 in nephrocytes as an invertebrate podocyte model and observed colocalization with Rab5 and a nephrocyte-restricted loss-of-function phenotype.

976C An in vivo screen identifies small molecule modulators of the endoplasmic reticulum stress response

Kevin Hope, Alexys Berman, Clement Chow Human Genetics, University of Utah, Salt Lake City, UT

Misfolded protein accumulation in the endoplasmic reticulum (ER) induces ER stress. Cells respond to ER stress by initiating the unfolded protein response (UPR) that upregulates chaperone protein expression, increases the degradation of misfolded proteins, and inhibits protein translation. Failure to effectively manage ER stress and restore homeostasis results in cellular dysfunction and ultimately apoptosis, a process implicated in numerous human diseases such as retinal degeneration, Alzheimer’s disease (AD), Parkinson’s disease (PD), Huntington’s disease (HD), and amyotrophic lateral sclerosis (ALS), among others. Identifying small molecules that modulate ER stress may be effective therapeutics for human diseases caused by misfolded protein accumulation. Here, we used a *Drosophila* model of retinitis pigmentosa (RP) that expresses misfolded rhodopsin protein, Rh1\(^{G69D}\), in the developing eye. Rh1\(^{G69D}\) expression induces chronic ER stress and apoptosis, resulting in a degenerative eye phenotype. We took a drug repurposing approach and used the Prestwick Chemical Library, consisting of 1520 small molecules, the majority of which are FDA-approved, to identify compounds that modulate neuron cell death in Rh1\(^{G69D}\) expressing flies. We identified multiple classes of drugs that enhance or suppress the degenerative eye phenotype, including compounds acting through monoamine neurotransmitters, folate metabolism, sodium channels, and the renin/angiotensin pathway. Degeneration-enhancing compounds may reveal novel ER stress pathways, and compounds that suppress degeneration are potential therapeutic candidates for RP. We are using an RNAi approach to identify the mechanism of action for the top enhancers and suppressors. Additionally, we will present findings on whether compounds that rescue cell death in the RP model can also rescue disease-associated phenotypes in other *Drosophila* models of protein misfolding diseases, such as PD, HD, and ALS. This work identified potential therapeutic drugs for RP and possibly other human diseases that result from misfolded protein accumulation and ER stress.

977A Humanized *Drosophila* model of the Meier-Gorlin syndrome. Maxim Balasov, Katarina Akhmetova, Igor Chesnokov UAB

Meier-Gorlin syndrome (MGS) is a rare autosomal recessive disorder characterized by microtia, primordial dwarfism, small ears and skeletal abnormalities. Patients with MGS often carry mutations in the genes encoding the subunits of the Origin Recognition Complex (ORC), components of the pre-replicative complex (pre-RC) and replication machinery. Orc6 is an important component of ORC and has functions in both DNA replication and cytokinesis. A mutation in conserved C-terminal motif of Orc6, Y225 to S, is associated with MGS and impedes the interaction of Orc6 with the core ORC. Recently, a new mutation in Orc6, K23 to E, was identified however, it is localized in the N-terminal domain of the protein. In order to study the functions of new Orc6 mutation in live animal system we used human Orc6 gene or a hybrid Orc6-HD transgene (containing intact human N-terminal TFIIB like domain and *Drosophila* C-terminal) to rescue the orc6 deletion in *Drosophila*. The obtained flies survive to the adult stage and allow studies of MGS mutations in humanized animal model system. Using this approach we discovered that unlike previously identified Y225S MGS mutation in Orc6, the K23 to E substitution in the N-terminal TFIIB-like domain does not disrupt the interaction between Orc6 and the rest of the ORC. However, K23E MGS mutation results in a reduced DNA binding ability of the Orc6 protein.
The flies carrying MGS mutations are unable to fly and display growth and development defects. Overall, despite having different underlying molecular mechanisms both MGS mutations resulted in similar phenotypes, deficient pre-RC formation and reduced DNA replication.

Our studies revealed the importance of evolutionarily conserved and variable domains of Orc6 protein and allowed the studies of human protein functions and the analysis of the critical amino acids in live animal heterologous system as well as provided novel insights into the mechanisms underlying MGS pathology. We believe that hybrid approach not only open a broad avenue to study new Orc6 mutations for medical and general science purposes but might be useful in other humanized models.

978B  Mimicking human disease-causing mutations in Drosophila PLC-γ  Mariah Torcivia, Alli Raymond, Justin Thackeray  Clark University

Activating mutations of both PLC-γ1 and -γ2 are observed in a variety of human diseases. This includes somatic mutations in a variety of cancers, as well as germline changes underlying auto-immune disorders; some inherited, activating, PLC-γ mutations have also been shown to be protective in early-onset Alzheimer disease. Drosophila has a single PLC-γ homolog encoded by small wing (sl) and although many sl mutations have been isolated, all are either hypomorphic or loss of function alleles. We are therefore using CRISPR to recreate in Drosophila some of the most commonly observed mammalian PLC-γ activating mutations. These mutant lines should be useful both in understanding the role of increased PLC-γ signaling in human disease, as well as providing reagents to screen for drugs that block this activation. One successful edit made to date is viable, mimicking the PLCγ1 S345F mutation observed in the malignancies from many human T cell leukemia patients. Flies homozygous for an sl349F edited X-chromosome typically show normal eye development but, in a background with increased dosage of sl+, ommatidia in two independent lines frequently show abnormal photoreceptor development, in which R3 is replaced by two photoreceptor cells with smaller rhabdomeres. This phenotype is similar to that seen in loss of function mutations of the nuclear receptor encoded by seven-up, in which R1, R3, R4, or R6 cells are transformed into R7. We are in the process of isolating additional activating mutations, and further characterizing the defect in the sl349F mutant line.

979C  Understanding the Progressive Loss of Larval Muscle Fibers in Cachexia Tumor Model System with Focus on Myosin  Ellen Thompson, Grace Stegemoller, Logan McDowell, Mardelle Atkins  Sam Houston State University

Cachexia is a wasting syndrome common in late-stage cancer patients, whose symptoms include loss of body fat, weight loss, and muscle weakness/deterioration. To study the cachexic effects on muscles, our lab utilizes a Drosophila melanogaster tumor model which displays the syndrome's major symptoms. These tumor larvae fail to pupate and display observable physical wasting and symptoms of muscle loss over the period of 8-12 days AEL. In my project, I am distinguishing how the muscle sarcomere structure is changing due to cachexia. I observe a progressive loss of larval skeletal muscle strength was detailed over a five-day span (day 8-12 in development). I determined that there is a progressive loss of the larval skeletal muscle integrity, with late day 12 larvae having many broken muscle fibers. At a subcellular level, loss of sarcomere structure is increasingly common over the progression of the disease. Using immunohistochemistry, I observed that larval skeletal muscle fibers displayed major defects in actin, alpha-actinin, and myosin heavy chain localization towards the later stages of the disease, with fewer disruptions on day 8 or 10. I also observed that of these three molecules, myosin heavy chain shows the earliest and most pervasive phenotypes. From this, it is prophesied that this structural change in the sarcomere protein structure is related to the cause of the breakdown in the larval skeletal muscles. These results suggest that myosin heavy chain mislocalization is an early event in the loss of muscle function and integrity observed in these larvae, and our future studies will investigate if this is due to decreased protein production or increased turnover.

980A  miR-277 targets hid to ameliorate Aβ42-mediated neurodegeneration in Drosophila eye model of Alzheimer’s Disease  Projakta Deshpande1, Chao-Yi Chen2, Anuradha Chimata2, Catherine Yeates2, Chun-Hong Chen3,3, Madhuri Kango-Singh4,5,6, Amit Singh1,4,5,6,7  1) Department of Biology, University of Dayton, Dayton, OH; 2) Institution of Molecular and Cellular Biology, National Taiwan University, Taipei, Taiwan; 3) National Institute of Infectious Diseases and Vaccinology, National Health Research Institutes, Miaoli, Taiwan; 4) Premedical Program, University of Dayton, Dayton, OH; 5) Center for Tissue Regeneration & Engineering (TREND), University of Dayton, Dayton, OH; 6) Integrative Science and Engineering (ISE), University of Dayton, Dayton, OH; 7) Center for Genomic Advocacy (TCGA), Indiana State University, Terre Haute, IN

Alzheimer’s disease (AD), an age-related progressive neurodegenerative disorder, exhibits reduced cognitive functions with no cure to date. One of the reasons for AD is the extracellular accumulation of Amyloid-beta 42 (Aβ42) plaques. We misexpressed human Aβ42 in the developing retina of Drosophila, which exhibits AD-like neuropathology. Accumulation of Aβ42 plaque(s) triggers aberrant signaling resulting in neuronal cell death by unknown mechanism(s). We screened for microRNAs (miRNAs) which post-transcriptionally regulate expression of genes by degrading mRNA of the target genes. In a forward genetic screen with candidate miRNAs, we identified miR-277 as a genetic modifier of Aβ42-mediated neurodegeneration. Gain-of-function of miR-277 rescues Aβ42-mediated neurodegeneration whereas loss-of-function of miR-277 enhances Aβ42-mediated neurodegeneration. Moreover, misexpression of higher levels of miR-277 in
the GMR>Aβ42 background restores the retinal axonal targeting indicating functional rescue. Furthermore, we have identified head involution defective (hid) as one of the targets of miR-277 by Fly TargetScan and validated by luciferase assay and qPCR. The hid transcript levels are decreased by ~2.3-fold when miR-277 is misexpressed in the GMR>Aβ42 background in comparison to the GMR>Aβ42 fly model. Hence, here we provide a mechanism of how miR-277 modulates Aβ42-mediated neurodegeneration by regulating hid transcript levels and demonstrate its neuroprotective role in Aβ42-mediated neuropathology.

981B  A Screen to Identify Genetic Modifiers of Seizure Susceptibility in a Drosophila model of PIGA Deficiency Shayna Scott, Emerald Lane, Emily Coelho, Clement Chow University of Utah, Salt Lake City, UT

Phosphatidylinositol glycan class A (PIGA) deficiency is an X-linked condition classified as a congenital disorder of glycosylation (CDG). CDGs are rare genetic diseases that result from mutations in genes that affect the biosynthesis or addition of glycans to other macromolecules. PIGA deficiency shares many symptoms with other CDGs such as shortened lifespan, hypotonia, facial dysmorphism, and epileptic seizures. PIGA encodes an enzyme that catalyzes the first step of glycosylphosphatidylinositol (GPI) biosynthesis transferring a N-acetylglucosamine to phosphatidylinositol. GPI-anchored proteins (GPI-APs) are localized to the cell surface and involved in processes such as endocytosis, immunity, and signal transduction. Like many CDGs and rare diseases, variability in phenotypic severity is common in PIGA deficiency patients, though the reason behind this variability is unknown. Because these disorders are often congenital, background genetics likely plays a big role. It is necessary to study a large number of genetic backgrounds to fully understand what affects the phenotypic outcome of a disease. To do this, we performed a genetic screen that utilizes the genetic variation found in ~200 lines of the Drosophila Genetic Reference Panel (DGRP) to identify potential genetic modifiers that impact seizure susceptibility in our PIGA deficient Drosophila model. We found that seizure susceptibility associated with PIGA deficiency is highly dependent on genetic background, with seizure susceptibility between 0-60%. We also found that median seizure recovery time was also highly dependent on genetic background. There is a moderate correlation between seizure susceptibility and recovery time, suggesting only a partial overlap between the genetic architecture that underlies variability of either phenotype. We also found that there was a moderate correlation between the sexes for both phenotypes, suggesting some sex-specificity to the modifier genes. To identify modifiers of these phenotypes, we performed genome wide association analyses for each independently, and by sex. We will present functional characterization of the strongest genetic modifiers. Further investigation into genetic modifiers that impact the seizure susceptibility will provide insights into improved treatments and personalized therapies.

982C  Exploring the role of shaggy and dmyc in development of combination therapy against human neuronal tauopathies in Drosophila. Pragati, Surajit Sarkar Department of Genetics, University of Delhi, South Campus, Dhaula Kuan, New Delhi, India

Human neuronal tauopathies which include Alzheimer’s, Parkinson’s, Pick’s disease(s) etc. are group of neurodegenerative disorders characterized by aberrant tau hyperphosphorylation resulting in the formation of toxic tau oligomers, paired helical filaments (PHFs) and neurofibrillary tangles. These toxic aggregates were found in different specific regions of the brain resulting in various symptomatic features in patients. Even though several attempts are being made, the definite cause of these disorders remains unexplained certainly due to complex disease traits and limitations associated with the availability of human disease samples. We have reported earlier that targeted reduction of dmyc (a Drosophila homologue of cmyc proto-oncogene) constrains NFTs mediated tau pathogenesis. Further, in order to unravel the molecular insights, our findings suggest a pivotal role of shaggy (a Drosophila homologue of gsk3β, kinase) in conferring the dmyc mediated rescue against tauopathies. We observed that concurrent downregulation of both dmyc and shaggy confers an additive rescue in restricting tau hyperphosphorylation levels, neurofibrillary tangles formation, and in restoring heterochromatin loss to the physiological levels against tau-mediated toxicity in Drosophila. The subsequent analysis also showed that dmyc regulates shaggy activity via regulating protein phosphatase 2A (dPP2A) in a dose-dependent manner. Taken together, our study provides novel molecular insights about the role of shaggy in tau aetiology, which may aid in developing combinatorial drug(s) against the devastating human neuronal tauopathies.

Keywords: Drosophila; Tauopathies; dMyc; Shaggy; Neurofibrillary tangles

983A  Coevolution is pervasive between unrelated glycosylation pathways and points to potential disease modifiers Holly Thorpe, Nathan Clark, Clement Chow University of Utah Department of Human Genetics, Salt Lake City, Utah

Glycosylation is one of the most common post-translational modifications. Defects in glycan biogenesis pathways, such as N-linked glycosylation, O-linked glycosylation, and GPI anchor synthesis, lead to rare, multi-systemic disorders classified as Congenital Disorders of Glycosylation (CDG). CDGs typically present with seizures, hypotonia, and developmental delay, but display large clinical variability with symptoms affecting every system in the body. This variability suggests modifier genes affect the phenotypes. I am employing evolutionary approaches to identify modifier genes of CDGs. Evolutionary Rate Covariation (ERC) relies on the premise that proteins that interact physically or genetically or are functionally related coevolve at similar rates. ERC values are calculated using the correlation coefficient of evolutionary
rates of gene pairs in a species tree. I used ERC values to look genome wide for coevolution with CDG genes, specifically genes involved in GPI anchor synthesis.

There was enriched coevolution among GPI anchor synthesis proteins. Unexpectedly, there was also enriched coevolution between GPI anchor synthesis proteins and proteins in other glycosylation pathways, suggesting more overlap between the different pathways than appreciated. Gene Ontology analysis of top genes that coevolve with GPI anchor synthesis proteins showed enrichment in genes involved in RNA modification and mitochondrial gene expression, suggesting interactions between these processes and GPI anchor synthesis. Gene pairs with the highest coevolutionary scores included both HTT and PIGG and ATG7 and PIGG. HTT and ATG7 are associated with neurodegenerative disorders possible indicating overlap in pathophysiology between the disorders.

To functionally validate these exciting signals, I screened for genetic interactions using the Drosophila eye. Many GPI anchor synthesis genes are necessary for Drosophila eye development and knockdown of these genes leads to rough and disorganized eyes. By creating double knockdowns of GPI anchor synthesis genes and coevolving genes in the Drosophila eye, I identified genetic interactions between genes previously thought to be unrelated. Many of the strongest evolutionary signals validate as interactors in this in vivo analysis. Coevolution is an underutilized tool for identifying interactions between unrelated proteins. These connections could lead to better understanding of glycosylation pathways and potential treatments for CDGs.

984B  Drosophila eye model to study the role of NAT9 in Alzheimer’s Disease related Dementia (ADRD) Emily Snider1, Prajakta Deshpande2, Amit Singh1,2,3,4,5 1) Department of Biology, University of Dayton, Dayton, OH; 2) Premedical Program, University of Dayton, Dayton, OH; 3) Center for Tissue Regeneration & Engineering (TREND), University of Dayton, Dayton, OH; 4) Integrative Science and Engineering (ISE), University of Dayton, Dayton, OH; 5) Center for Genomic Advocacy (TCGA), Indiana State University, Terre Haute, IN

Alzheimer’s Disease (AD), an age-related progressive form of dementia, is characterized by a decline in cognitive function. Accumulation of the peptide amyloid beta (Aβ42) plaque is one of the characteristics of the disease. The accumulation of these Aβ42 plaques triggers the hyperphosphorylation of tau, a microtubule associated protein, thereby destabilizing the microtubules. This results in the intracellular accumulation of neurofibrillary tangles. We employed the Gal4/UAS system in Drosophila melanogaster to misexpress human Aβ42 within the developing fly retina, exhibiting AD-like neuropathology. Accumulation of Aβ42 plaque(s) triggers the aberrant activation of signaling pathways like the JNK pathway resulting in neuronal cell death by unknown mechanism(s). Using forward genetic screening, we identified N-acetyltransferase 9 (NAT9) as one of the genetic modifiers of GMR>Aβ42 reduced eye phenotype. The previous study suggests that NAT9 stabilizes microtubules by acetylation of tubulins, thereby inhibiting JNK signaling. This study aims to understand the role of NAT9 in Aβ42-mediated neurodegeneration. The gain-of-function of NAT9 in GMR>Aβ42 background suppresses the Aβ42-mediated neurodegeneration whereas loss-of-function of NAT9 in GMR>Aβ42 background enhances Aβ42-mediated neurodegeneration. The eye antennal imaginal discs of loss-of-function of NAT9 in GMR>Aβ42 background shows the activation of JNK pathway by increased pJNK levels. Hence, here we propose that NAT9 downregulates JNK signaling pathway which can ameliorate Aβ42-mediated neurodegeneration.

985V  Age-dependent Lamin Remodeling Induces Cardiac Dysfunction via Dysregulation of Cardiac Transcriptional Programs Natalie Kirkland1, Alexander Whitehead2, James Hocker3, Pranjali Beri1, Geo Vogler4, Bill Hum2, Mingyi Wang2, Edward Lakatta5, Bing Ren5, Rolf Bodmer2, Adam Engler1 1) University of California, San Diego; 2) Sanford Burnham Prebys Medical Discovery Institute; 3) National Institute on Aging

As we age, structural changes contribute to progressive decline in organ function, which in the heart acts through poorly characterized mechanisms. Utilizing the rapidly aging fruit fly model with its significant homology to the human cardiac proteome, we found that cardiomyocytes exhibit progressive loss of Lamin C (mammalian Lamin A/C homologue) with age. Unlike other tissues and laminopathies, we observe decreasing nuclear size, while nuclear stiffness increases. Premature genetic reduction of Lamin C phenocopies aging’s effects on the nucleus, and subsequently decreases heart contractility and sarcromere organization. Surprisingly, Lamin C reduction downregulates myogenic transcription factors and shows that maintenance of Lamin C sustains their expression and prevents age-dependent cardiac decline. Our findings are conserved in aged non-human primates and mice, demonstrating age-dependent nuclear remodeling is a major mechanism contributing to cardiac dysfunction.

986V  Loss of MECR, an enzyme for mitochondrial fatty acid synthesis, causes iron accumulation, upregulation of ceramides and neurodegeneration Debddeep Dutta1,2, Oguz Kanca1,2, Seul Kee Byeoun3, Jun Hyoung Park4, Paul C. Maragolli2, Zhongyuan Zuo1,2, Rishi V. Shridharan1, Guang Lin1,2, Matthew Wheeler3, Benny A. Kaipparettu1, Akhilesh Pandey1, Hugo J. Bellen1,2,5, Undiagnosed Diseases Network 1) Department of Molecular and Human Genetics, Baylor College of Medicine, Houston, TX; 2) Jan and Dan Duncan Neurological Research Institute, Texas Children’s Hospital, Houston, TX; 3) Department of Laboratory Medicine and Pathology, Mayo Clinic, Rochester, MN, US; 4) Division of Cardiovascular Medicine, Stanford University School of Medicine, Stanford, CA; 5) Department of Neuroscience, Baylor College of Medicine, Houston, TX
Fatty acid synthesis occurs in the cytoplasm as well as in the mitochondria of eukaryotic cells. Autosomal recessive mutations in MECR (Mitochondrial Enoyl CoA Reductase), which encodes an enzyme required for the last step of the mitochondrial fatty acid synthesis, cause a childhood-onset pediatric neurodegenerative disorder called MEPAN syndrome (MIM # 617282). To study the mechanism underlying the neurodegenerative phenotypes observed in MEPAN syndrome, we generated a severe loss-of-function CRIMIC (SA-T2AGAL4-polyA) allele of the fly ortholog of MECR, CG16935/mecr using CRISPR. Homozygous mutants die at larval stages and the lethality is rescued upon expression of the reference human MECR cDNA, showing functional conservation across species. Given the lethality, we performed RNAi-mediated knockdown experiments in neurons and observed progressive locomotor defects and vision loss. Upon successfully modelling the key disease phenotypes in our fly model, we used the fly and fibroblasts from patients to evaluate the mitochondrial function and morphology. A GFP-tagged allele of mecR expressed at the endogenous levels showed that the protein is localized to mitochondria. Transmission Electron Microscopy of Drosophila eye mutant clones and the MEPAN patient fibroblasts revealed reduced mitochondrial numbers and abnormal mitochondrial morphology. Additionally, mitochondrial function (mitochondrial membrane potential, ATP production, electron transport chain activity, and oxygen consumption rates) are all compromised due to loss of mecR/MECR. Lipidomic analyses of fly mutants and MEPAN patient fibroblasts revealed elevated ceramide levels. Hence, our next question was: how does loss of mecR increase ceramide levels? We found that loss of mecR affects the stability of mtACP, a multifunctional protein that helps in the assembly of Fe-S cluster biogenesis. Consistent with decreased mtACP levels, loss of mecR causes Fe-S cluster biogenesis defects and elevated iron levels in the larval brain. Excess iron results in the accumulation of ceramides leading to mitochondrial defects and neurodegeneration in MEPAN syndrome.

987V  Transcription related proteins modify TDP-43 mediated toxicity in a fly model of ALS Deepak Chhangani1, Swapnil Pandey2, Lorena de Mena3, Pedro Fernandez-Funez1, Diego Rincon Limas1, 2 1) Department of Neurology, University of Florida, Gainesville, FL; 2) Department of Neuroscience, University of Florida, Gainesville, FL

Tar DNA binding Protein-43 (TDP-43) is a major DNA/RNA binding protein involved in multiple cellular processes including transcriptional regulation, mRNA splicing and stress granules formation. Mutations in TDP-43, such as TDP-43

988V  Vexed mediates non-cell autonomous loss of dopaminergic neurons Jacinta Davis, Elizabeth Kolaski, Daniel Babcock Lehigh University

Parkinson’s Disease (PD) is the most prevalent movement disorder with about 10 million people worldwide affected. The hallmark of PD is the loss of dopaminergic (DA) neurons. Little is known as to why these neurons are vulnerable to PD. We performed a genome-wide screen using over 200 wild-caught Drosophila melanogaster lines from the Drosophila Genetic Reference Panel (DGRP) to identify genes involved in the maintenance of DA neurons. From this screen, we found the gene CG42339. CG42339 is a previously unannotated gene with a human ortholog. Mutations and non-cell autonomous knockdown of CG42339 caused DA neuron loss in the Drosophila brain at late adulthood. We also discovered a progressive locomotor defect in the mutations and knockdown of CG42339. We further examined CG42339 by investigating the mechanistic function this gene plays in the maintenance of DA neurons. We were able to rescue this loss of DA neurons in a CG42339 mutant by limiting the innate immune response and inhibiting nitric oxide signaling. CG42339 appears to be acting as a receptor for advanced glycation end-products (RAGE), therefore we named this gene vexed.


Alzheimer’s disease (AD) is a multifactorial neurodegenerative disorder characterized by dementia and cognitive decline due to progressive cerebral cortical atrophy. Brains of AD patients are characterized by the accumulation of microscopic extracellular amyloid-beta (Aβ) plaques and intracellular neurofibrillary tangles composed of hyperphosphorylated tau. The deposition of Aβ42, which is one of the fragments of amyloid precursor protein (APP), has been known to play a role in initiating the events leading to the formation of amyloid and subsequently hyperphosphorylation of tau. However, animal models expressing either Aβ42 or tau individually do not mimic the complexity of the human condition. Indeed,
recent evidence suggests that Aβ42 and pathological tau interact synergistically to modulate neurotoxicity in AD. To shed light on their concerted roles in AD pathogenesis and to discover pathways mediating Aβ42 and tau interactions, we generated transgenic flies co-expressing human Aβ42 fused to a signal peptide along with the longest wild-type tau isoform. Overexpression of Aβ42 or tau in Drosophila using the UAS-Gal4 system causes mild to the moderate rough eye. In comparison, co-expression of Aβ42 with tau causes severe roughening and reduction of the eye size. The level of neuronal cell death in eye tissues was also significantly enhanced in flies co-expressing Aβ42 and tau. To identify pathways mediating Aβ42+tau interactions, we are currently using the Aβ42+tau eye phenotype as platform to screen 1,500 UAS lines expressing a variety of human genes. We have identified few enhancers and suppressors not previously known to be involved in AD pathogenesis, which will be helpful to uncover new molecular pathways and potential therapeutic targets. This work is supported by NIH grant R21AG069050 to DERL.

990V  Intra- and extra-cellular functions of ALS-related ER protein VAP in Drosophila Kosuke Kamemura1; Misako Okumura1,2; Takahiro Chihara1,2 1) Program of Biomedical Science, Graduate School of Integrated Sciences for Life, Hiroshima University, 1-3-1 Kagamiyama, Higashi-Hiroshima, Hiroshima 739-8526, Japan; 2) Program of Basic Biology, Graduate School of Integrated Sciences for Life, Hiroshima University, 1-3-1 Kagamiyama, Higashi-Hiroshima, Hiroshima 739-8526, Japan

VAP is a type II integral transmembrane protein localized at the endoplasmic reticulum (ER), and functions as a tethering protein of the membrane contact sites between ER and various intracellular organelles. Recent studies revealed that VAP is cleaved, and its N-terminal MSP domain is secreted to the extracellular space in C. elegans, Drosophila and also in human. The secreted VAP MSP domain is known to bind to a variety of axon growth cone guidance receptors (Eph, Robo, Lar etc.), implying that VAP could function non-cell autonomously. In addition, mutations in human VAPB were known to cause neurodegenerative diseases such as amyotrophic lateral sclerosis 8 (ALS8). However, as the VAP loss-of-function exhibits severe lethality in early developmental stages, the physiological functions of VAP, especially the extracellular functions of VAP MSP domain are not well understood.

In order to investigate the physiological functions of VAP (Drosophila ortholog is Vap33), we utilized Drosophila olfactory projection neurons with a mosaic analysis with a repressible cell marker (MARCM), which allows us to analyze the function of a gene of interest in a single cell resolution in vivo. In vap33+/ MARCM clones, the dendrite but not axon of projection neurons exhibited the severe morphological defects, which are rescued by the cell-autonomous expression of vap33 cDNA, suggesting that VAP cell-autonomously and preferentially regulates dendrite morphology. Interestingly, subcellular localizations of Golgi apparatus and mitochondria were also severely affected. These results indicate the requirement of intracellular functions of Vap33 in neural development. Moreover, to examine the roles of the secreted Vap33 MSP domain, we searched the important sequence required for Vap33 MSP secretion and succeeded to generate the “unsecretable vap33 mutant” whose Vap33 MSP is not secreted but can function intracellularly. The unsecretable vap33 mutant exhibited lethality. Overexpressing the unsecretable Vap33 in the vap33+/ PN clones rescued the dendritic defects, implying that the intracellular functions of unsecretable Vap33 are likely to be normal. From these results, we propose that secreted MSP domain plays important extracellular functions to maintain organismal viability. We’d like to discuss how secreted MSP domain contributes the animal physiology, which may lead the clues for the therapeutic strategy of neurodegenerative diseases.

991V  Characterizing synaptic deficits at adult neuromuscular Junctions in a model of Amyotrophic Lateral Sclerosis Jessica Sidisky, Brandon Hocking, Nikki Huhulea, Sara Moran, Bali Connors, Daniel Babcock Lehigh

The hallmark of neurodegenerative diseases such as Alzheimer’s Disease (AD), Parkinson’s Disease (PD), and Amyotrophic Lateral Sclerosis (ALS) is the loss of neurons. However, recent evidence suggests that synapse dysfunction occurs long before clinical symptoms are manifested. Although PD and AD are the most prevalent neurodegenerative diseases, ALS has the most rapid onset of symptoms from the time of clinical diagnosis. Although there has been significant progress made in understanding the disease, we still lack effective cures or treatments. Here we introduce an adult model of ALS to characterize the progression of synaptic deficits that precede motor neuron loss. Using a tissue specific-Gal4 and the temperature sensitive Gal80 to express human and mutant fused in sarcoma (FUS), a gene responsible for a genetic form of ALS, in the flight motor neurons in a spatial and temporally controlled manner. Synaptic deficits are characterized using the flight behavioral assay as a direct read out of flight motor neuron function. We found that overexpression of mutant FUS demonstrates a progressive loss of flight ability. We further define these synaptic defects using synaptic markers to understand how and when neuromuscular junctions (NMJs) are impaired before motor neuron loss. We also characterize the adult ventral abdominal NMJs in the same manner using a motor neuron driver specific to these NMJs. With this model, we hope to elucidate the mechanisms that underly synapse loss in a structural and functional manner.

992V  Targeted downregulation of Hipp1 ameliorates tau-engendered deficits in Drosophila melanogaster SUNG YEON PARK1,3, JIEUN SEO2, Seulbee Lee2, Sang Jeong Kim1,2,3, Yang-Sook Chun1,2,3 1) Ischemic/Hypoxic Disease Institute, Seoul National University, College of Medicine, Seoul, Republic of Korea; 2) Department of Biomedical Sciences, Seoul National University College of Medicine, Seoul, Republic of Korea; 3) Department of Physiology, Seoul National University
Tauopathies, such as Alzheimer’s disease (AD), are neurodegenerative diseases characterized by the deposition of neurofibrillary tangles comprising hyperphosphorylated tau protein in the human brain. Given that abnormal epigenetic alterations in heterochromatin configuration have been documented in AD patients and transgenic animal models of AD, we investigated the roles of novel heterochromatin-associated interactors in tauopathies. We examined whether tissue-specific downregulation or loss-of-function alleles of heterochromatin-associated interactors can affect tau-induced neurotoxicity using transgenic flies via UAS-Gal4 binary system. Here, we found that knockdown of H1P and insulator partner protein (Hipp1) ameliorates tau-engendered eye defects, locomotion defects, reduced lifespan, weight loss, and neurodegeneration by preventing hyperphosphorylation of tau. Nonetheless, RNAi-mediated reduction of Hipp1 failed to restore tau-induced heterochromatin loosening; it accelerated abnormal overexpression of heterochromatic genes. Instead, knockdown of Hipp1 restored tau-troubled aberrant expression of putative insulator targets and aberrant insulator-mediated epigenetic alterations. HIPP1 may have a role as an insulator binding partner regarding to be implicated in tau-induced neurodegeneration. Moreover, knockdown of Hipp1 in flies overexpressing tau restored the aberrant expression of AD susceptibility genes, Amph and Sox102F. These results suggest that downregulation of Hipp1 expression may be a potential therapeutic target in neurodegenerative disorders; they also provide new insights regarding the roles of insulator proteins in tauopathies.

**993V** De novo missense mutations in E3 ubiquitin ligase RNFT2 lead to intellectual disability as evidenced by loss of function studies in Drosophila

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Characterized by various deficiencies in both intellectual ability and adaptive behavior, intellectual disabilities (IDs) affect 1% of the population worldwide. Mutations in over 400 genes have been implicated in ID, however the underlying molecular links between genotype and phenotype for most of these genes remain unknown. Whole exome sequencing (WES) of a large cohort of 404 Iranian consanguineous families with autosomal recessive intellectual disability (ARID) led to the identification of a novel likely pathogenic missense variant (cT1150C; p.C384R) in the RNFT2 gene. RNFT2 (RING finger transmembrane domain containing protein 2, also named as TMEM118), encodes a RING finger E3 ubiquitin ligase. Its fly ortholog dmel/CG13605 has the same type of RING domain (C3HC4 type) and is predicted to be a ubiquitin E3 ligase involved in the ER-regulated protein degradation pathway. How disruption of RNFT2 gives rise to ID phenotypes is not known. To address this question, we sought to investigate the neurobehavioral consequences of losing the RNFT2 homolog in flies and identify its expression in the fly brain. In parallel, we investigated its subcellular localization in stably transfected HeLa cells. These cells will be used to identify the substrates of RNFT2.

Towards this end we have generated a Gal4 line and show that CG13605 is expressed in a subset of Kenyon cells in the fly brain starting at larval stages. Downregulation of CG13605 in the mushroom body with OK107-Gal4 lead to neurodevelopmental phenotypes such as loss of alpha and alpha’ lobes. For a more detailed functional characterization, we generated null mutant flies using the CRISPR/Cas9 system. Loss of CG13605 led to phenotypes such as alpha lobe loss or beta lobe fusion. The observed defects in mushroom body development, which is the primary memory and learning center of flies, provide evidence for RNFT2 being an ID gene. Ongoing experiments including rescue experiments, analyses using patient-specific variants of RNFT2 as well as behavioral studies are expected to further support this hypothesis.

**994V** Homologues of the human disease-associated amyloidogenic proteins APP and TGFBI are required for physiological protein aggregation in Drosophila secondary cells

Clive Wilson, Preman J Singh, Adam Wells, Claudia C Mendes, S Mark Wainwright, Pauline P Marie, Ben Kroeger, Deborah C I Goberdhan University of Oxford

Amyloidogenesis, the aggregation of soluble proteins into insoluble fibrils, has multiple biological functions in both health and disease. It is critically important in neurodegenerative disorders; for example, aggregations of A-beta peptides, cleaved products of Amyloid Precursor Protein (APP), form plaques in Alzheimer’s Disease (AD). Mutated Transforming Growth Factor-Beta-Induced (TGFBI), an extracellular fibrillar protein, assembles into amyloid that leads to corneal dystrophies. Proteins also aggregate naturally, sometimes via amyloidogenesis, for example, when peptide hormones condense into insoluble, inert dense-core granules (DCGs), which are stored in secretory vesicles until release. Understanding the cell biology that drives these normal and pathological processes could reveal the key mechanisms that distinguish them, but visualising these events as they take place in cells and tissues is challenging. Here we describe the development of a new cellular model for DCG biogenesis, the Drosophila prostate-like secondary cell (SC). These cells have highly enlarged (5 micron diameter) DCG compartments, permitting DCG assembly to be followed by light and fluorescence microscopy in living tissue and in real-time. Remarkably, the rapid formation of DCGs requires the complementary activities of the fly homologues of TGFBI, called MFAS, and APP, called APPL. While DCGs cannot form
in the absence of MFAS, several smaller DCGs are often made in a single compartment in the absence of APPL, but they frequently cannot coalesce into a single large DCG, as happens in normal cells. Expressing a pathological, amyloidogenic form of human A-beta or TGFBI in SCs alters MFAS assembly in DCGs and can suppress DCG disassembly when secreted. Genetic manipulations that mirror changes seen in AD patients also modulate DCG biogenesis and disassembly. Interestingly, intraluminal vesicles (ILVs), which are secreted as exosomes, are formed in SC DCG compartments. We show they are also required for assembly of a large DCG, consistent with recent findings linking ILVs and exosomes to amyloidogenesis. Our studies, therefore, demonstrate that both APP and TGFBI have physiological, as well as pathological, modes of rapid protein aggregation, which can uniquely be distinguished by real-time fluorescence imaging in living SCs. We are employing this model to screen genetic modifiers of pathological versus physiological amyloidogenesis, which might highlight new ways to target amyloidogenic diseases.

995V  Novel dominant and recessive variants in human ROBO1 cause distinct neurodevelopmental defects through different mechanisms Yan Huang1,2, Mengqi Ma1,2, Xiao Mao3,4, Davut Pehlivan1,5,6, Oguz Kanca1,2, Gulsen Akay7, Tadahiro Mitani1, Shenzhao Lu1,2, Sukru Candan1, Bo Xiao8, James Lupski1,6, Hugo Bellen1,2 1) Department of Molecular and Human Genetics, Baylor College of Medicine, Houston, Texas, USA; 2) Jan and Dan Duncan Neurological Research Institute, Texas Children’s Hospital, Baylor College of Medicine, Houston, Texas, USA; 3) National Health Commission Key Laboratory of Birth Defects Research, Prevention and Treatment, Hunan Provincial Maternal and Child Health Care Hospital, Changsha, Hunan, China; 4) Department of Medical Genetics, Maternal and Child Health Hospital of Hunan Province, Changsha, Hunan, China; 5) Division of Neurology and Developmental Neuroscience, Department of Pediatrics, Baylor College of Medicine, Houston, Texas, USA; 6) Texas Children’s Hospital, Houston, Texas, USA; 7) Medical Genetics Section, Balikesir Ataturk Public Hospital, Balikesir, Turkey; 8) Neurology Department, Xiangya Hospital, Central South University, Changsha, Hunan, China

The Roundabout (Robo) receptors present on growth cones of neurons induce axon repulsion in response to the extracellular ligand Slit. The Robo family of proteins controls midline crossing of commissural neurons during development in vertebrate and invertebrate model organisms. Mono- and bi-allelic loss-of-function (LoF) of human ROBO1 (MIM: 602430) has been associated with a breadth of phenotypes, including neurodevelopmental defects such as strabismus, pituitary defects, intellectual impairment, as well as non-neuronal defects in heart and kidney, with reduced penetrance and highly variable expression. We identified two novel ROBO1 variants associated with distinct phenotypes. In family #1, we identified a biallelic missense (p.S1522L) variant in three affected siblings with a recessive trait isolated nystagmus. In family #2, we identified a de novo p.D422G variant in the proband who presented with a severe early-onset epileptic encephalopathy. To assess the perturbance to biological homeostasis for these variants, we performed functional assays in Drosophila. We generated a null allele of robo1 by inserting a CRIMIC T2A-GAL4 in an intron. To our surprise, transheterozygous robo1 null mutation (T2A-Gal4/Df) leads to reduced viability but not lethality. In contrast, overexpression of either human ROBO1 or fly robo1 is toxic but also reduces viability. Note that human ROBO1 is not able to replace fly robo1 when driven by the Gal4 insertion. The fly cDNA driven by the Gal4 rescues the midline crossing at 18°C. However, the p.D413G variant in the fly cDNA fails to rescue midline crossing suggesting that it is a LoF allele. The recessive ROBO1 variant cannot be tested as the p.S1522L is not conserved in fly robo1. We therefore turned to gain-of-function (GoF) assays using the T2A-GAL4 to drive the human reference cDNA which leads to toxicity. ROBO1 p.S1522L is less toxic than the reference human cDNA with respect to viability and midline crossing. In contrast, the dominant p.D413G variant leads to a highly aberrant protein distribution of fly Robo1 and creates novel defects in several assays. This suggests that it is a neomorphic allele. In summary, our studies expand the phenotypic spectrum associated with ROBO1 variant alleles and assesses the potential nature of the variants.

996V  Localization of transgenes for Drosophila models of myotonic dystrophy type 1 Andrea Waltrip, Noah Smith, Ginny Morriss University of Mary Washington, Fredericksburg, VA

Myotonic Dystrophy Type 1, DM1, is a multi-systemic disorder that results from expansion of CTG repeats in the DMPK gene in humans. Drosophila melanogaster has been established as a model organism for the study DM1, by the construct of multiple transgenic DM1 lines containing different numbers of CTG repeats (60, 250, and 480), expressed using the GAL4/UAS system. Expression of long-repeat transgenes ((CTG)250 and (CTG)480) has been shown to produce the phenotypes consistent with DM1, relative to control lines ((i)CTG)60. The precise chromosomal location of insertion of the transgenes has not been reported. We are using both classical genetic and molecular approaches to localize CTG-repeat transgene insertion in the genome. To narrow down location to a specific chromosome, genetic crosses using GAL4 drivers on different chromosomes are being used to drive expression of repeat expansions to assess phenotypic ratios of eye color traits, specific to the transgenes, and flight capability, which has been shown to be defective in DM1 flies. We expect to see different phenotypic ratios in the F1 progeny from crosses using drivers localized to different chromosomes. We will confirm our chromosome localization by fluorescent in situ hybridization (FISH) of polytene chromosome preparations, using probes specific for the transgene insertion. Chromosome specific probes will be used to verify the chromosomal location. Since the FISH will allow us to narrow down the location of transgene insertion to a more specific
region of the chromosome, we can target that specific region to more precisely determine the insertion site using PCR and sequencing. Preliminary results from the genetic analysis suggest that the (CTG)_{230} transgene is likely localized to chromosome 2 and that i(CTG)_{80} and i(CTG)_{10} are not likely localized to the chromosomes 1 or 2. Knowing the location of the transgenes can allow for more practical mating schemes to study various aspects of myotonic dystrophy disease mechanisms, but can also provide crucial information for understanding the expression of the transgenes, which may be influenced by nearby regulatory elements in the genome.

997V  Increased oxidative stress precedes activation of the seizure-exacerbating glial immune response in *prickle* mutants

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Epilepsy is a neurological disorder characterized by seizures and affects ~1% of the U.S. population. We have previously shown that mutations in *prickle* (*pk*) cause spontaneous myoclonic-like seizures and ataxia in *Drosophila*, similar to what is observed in humans carrying PRICKLE mutations. Moreover, a transcriptome analysis of *pk* mutant brains revealed a robust increase in glial-mediated innate innate immune response (GIIR), suggesting that the immune response might be connected to seizure progression. Using genetic, behavioral and immunohistochemical methods, we also showed that inhibiting the GIIR leads to a reduction of neurodegeneration which in turn suppresses the age-related exacerbation of seizures in *pk* mutants. However, while these experiments were the first to genetically demonstrate a link between a brain-derived innate immune system activation and epilepsy exacerbation, the link between *pk* neuronal dysfunction and activation of the GIIR remained unclear. Several studies have linked increased oxidative stress to numerous neurological disease models including epilepsy, and we observe significant upregulation of antioxidant genes in our transcriptome data, suggesting that oxidative stress might be the initiating factor for GIIR and eventual seizure exacerbation. Here, we show that larval, but not adult, *pk* mutant brains show increased neuronal oxidative stress, while only adult *pk* mutant brains show a significant increase in antioxidant and innate immune responses. Together, these data show that oxidative stress precedes the activation of GIIR and may be responsible for generating the signal to the glia, thus identifying a potential mechanism for GIIR-associated seizure generation in *pk* mutants. This work is supported by NIH/NINDS R01NS098590.

998V  Adding low levels of omega-3 and omega-6 fatty acids to the diet eliminates seizure-like activity and paralysis and alters gene expression in the bang-sensitive mutant *technical-knockout*

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There is an increasing amount of evidence that dietary and metabolic alterations can help reduce seizure susceptibility. The seizure-susceptible *Drosophila* bang-sensitive (BS) mutants represent a good model system for investigating the link between diet, metabolism, and seizure activity. The BS mutants exhibit seizure-like activity (SLA) and paralysis following a variety of stimuli including mechanical vortexing. Previous work demonstrated that feeding the BS mutants standard media supplemented with a commercial ketogenic dietary mix (10% w/v) could eliminate the seizure-like activity and paralysis seen in these mutants. Given the mix of lipids in the supplement, this study examined the ability of specific polyunsaturated omega fats to alleviate the seizure-paralysis phenotype in one of the BS mutants, *technical knockout* (*tko*). The *tko* flies were fed 10% (w/v) sucrose-only media supplemented with one of two omega-3 fatty acids, eicosapentaenoic acid and docosahexaenoic acid, or one omega-6 fatty acid, arachidonic acid. The omega fatty acids were added at a final concentration of 25 µg/ml of media and 1-2 day old *tko* male flies were reared in the media for 3 days prior to testing for bang sensitivity. All three of the omega fatty acids eliminated the SLA and paralysis normally seen in *tko* flies following mechanical shock. Following the 3-day exposure to the omega fatty acids, the flies were then switched back to standard media for ten days and retested for bang-sensitivity. Following this period of feeding on normal media, the flies still did not exhibit SLA and more than 90% of the flies did not exhibit paralysis. This indicates the ability of the omega fatty acids to rescue the phenotype was maintained even when the flies were subsequently removed from the diet.

To investigate changes in gene expression associated with the feeding of omega fatty acids, RNA was extracted from the heads of *tko* flies fed 1) docosahexaenoic acid (25 µg/ml), 2) arachidonic acid (25 µg/ml), and 3) control sugar-only media. The Affymetrix GeneChip *Drosophila* Gene 1.0 ST Array was then used to perform whole transcriptome analysis of these fly heads. When compared to flies on the control diet, the data demonstrated that a number of ribosomal proteins were highly upregulated in fly heads following both omega fatty acid supplementation diets. Given that previous researchers have seen alterations in ribosomal protein levels in dendrites and axons following cellular changes, the increases seen here may be associated with synaptogenesis and neural alterations that modify excitability of the *tko* nervous system.

999V  Keck1 inhibition of EGFR signaling: a Domain V mediated event

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#Dros22
The EGFR/ErbB family of receptor tyrosine kinases function in numerous developmental contexts and their dysregulation has been linked to various types of cancers. With receptor activation underlying their roles in cancer, molecules that inhibit EGFR/ErbB activity provide potential therapeutic avenues. One such inhibitory molecule is Kekkon1 (Kek1), a member of the Drosophila family of leucine-rich repeats (LRR) and immunoglobulin domain (Ig) containing transmembrane molecules. Kek1’s extracellular LRRs, along with Domain V of the Drosophila receptor, are critical for binding. It has also been reported that Kek1 is able bind in a cross-species fashion the mammalian ErbB receptors. However, while the extracellular regions of both mammalian and Drosophila receptors contain Domains I-IV, mammalian receptors lack Domain V. Therefore, to clarify the interaction of Kek1 with the receptors and elucidate the role of Domain V, we sought to further define the mechanism of binding and inhibition. Using an ELISA-based assay, a battery of wild-type, chimeras, and variants of the Drosophila and human receptors were assessed for interactions with Kek1. Our results demonstrate that Kek1 binds the Drosophila and not the human receptors, and that specificity occurs principally through Domain V, which is absent in the human receptors. Our results also implicate differences between Domain IV in the Drosophila and human receptors in further restricting Kek1’s interaction to the Drosophila receptor. Mechanistically, our results support a model in which Kek1 binds to Domain V on the dimerization and not the ligand-binding interface of the Drosophila receptor, thereby inhibiting receptor dimerization and consequent activation. Our work provides deeper insight to the structural requirements underlying Kek1's interaction with dEGFR and has clear implications for the proposed use of Kek1 as a therapeutic.

1000V RASopathy Drug Discovery Aimed at Treating Hypertrophic Cardiomyopathy Kimberly Stephens1, Jared Gatto2, Jianping Hu3, Céline Guichard2, Rupa Mirmira1, Tirtha Das1, Husnu Kaniskan1, Jian Jin1, Ross Cagan3, Bruce Gelb1 1) Mount Sinai School of Medicine; 2) Aix Marseille University; 3) University of Glasgow

Introduction: RASopathies are pleiomorphic genetic traits, predominantly resulting from gain of function in RAS/MAPK signaling. Hypertrophic cardiomyopathy (HCM) is a leading therapeutic target because of its association with early mortality in affected infants. MEK inhibitors have shown promise in pre-clinical studies and compassionate-usage in patients but are not curative, have dermatological side effects and are not uniformly efficacious. Developing RASopathy therapeutics that can be administered long-term while allowing homeostatic levels of RAS signaling is challenging. To address that, we used phenotype-driven, whole-organism chemical screening in Drosophila RASopathy models to drive therapeutic discovery.

Hypothesis: Using a fruit fly-based platform, we can identify novel small molecular therapeutics for RASopathies.

Methods: We generated transgenic Drosophila bearing pathogenic alleles associated with HCM and screened a chemical library for lead compounds based on rescuing lethality. We chemically evolved our best hit, M1, iteratively and assessed rescue efficacy. We assessed putative targets using similarity ensemble analysis (SEA).

Results: The parent M1 compound showed weak efficacy toward transgenic RAF1-S257L flies with 10 uM dosing in fly media. JH107-7 was the best M1 derivative for the RAF1 model, achieving 50% rescue with 1 uM dosing. Testing in other RASopathy fly models revealed nearly complete rescue in flies expressing PTPN11-N308D but not BRAF-W531C. M1 derivative JH93-178T rescued 45% of BRAF flies but was not effective for other fly models. To assess potential as cancer therapeutics, we tested our M1 derivatives against two RAS-driven colon cancer fly models for which no single drug has shown efficacy. For JH107-7 and JH93-178T, we nearly matched efficacy of combination chemotherapy with trametinib and zoledronic acid, a drug combination effective in flies and a patient. JH107-7’s top 10 target predictions included RAF1, BRAF and mTOR.

Conclusion: Two M1-logs show efficacy for RASopathies and cancer in transgenic fly models with likely in-animal dosing of 5-50 nM (1/200 of concentration in food). Preliminary data with human RASopathy-mutant induced pluripotent stem cell-based HCM modeling suggest efficacy in rescuing cellular hypertrophy. We are currently assessing M1 derivatives’ impact on signaling pathways.

1001V How polyploid cells become tumor and how fly deals with it? Xian-Feng Wang, Chun-Ming Lai, Chih-Hsuan Chang, Honggun Bao, Deeptiman Chatterjee, Yi-Chun Huang, Wu-Min Deng Tulane University School of Medicine

Polyploid cells are frequently observed in human cancers and are positively correlated with malignancy and poor prognosis. Genomic instability and polyploidization which caused by polyploid cells re-enter mitosis and depolyploidization have been well studied, the appropriate microenvironment by which polyploid cells re-enter mitosis has been elusive. Here, taking advantage of our polyploid Drosophila salivary gland imaginal ring tumor model which is driven by polyploid mitosis and depolyploidization and single cell RNA sequencing, we identified essential factors, including JNK, JAK/STAT and Hippo, which provide favorable soil for polyploid cells re-enter mitosis and become tumor. Besides tumor initiation factors, we found that tumor initiating polyploid cells show potential cell polarity defects which makes those cells easily lose cell polarity and become tumor. We further found the tumor initiating polyploid cells enrich cell death signal, such as hid, rpr and Atg8, leading to those “dangerous cells” being eliminated at metamorphosis during normal development. Escaping from cell death of those polyploid cells at the early development stages and tumor situation are largely depended on cell death inhibitor Diap1. Our findings reveal tumor microenvironments that provide a tumor formation soil for polyploid cell tumorigenesis.
1002V Towards understanding the mechanism of tumorigenesis caused by centrosome dysfunction Chaitali Khan, Nasser Rusan National Heart, Lung, and Blood Institute, National Institutes of Health, Bethesda, MD 20892

Centrosome dysfunction is prevalent in many types of cancers; however, the mechanistic understanding by which centrosome defects cause cancer is still not very clear. Some studies have shown that centrosome amplification causes cancer by inducing mitotic errors and genomic instability. Other studies link centrosome dysfunction and tumorigenesis to the activation of a cellular stress response. Centrosome dysfunction can also lead to cancer by disrupting asymmetric stem cell division, which leads to errors in daughter cell fate. In Drosophila, genetic perturbations that disrupt Drosophila neural stem cells (NSC) asymmetric division, including centrosome mutations, give rise to tumors. Some of these mutations form tumors in the larva brain while others give rise to tumors upon transplanting NSC into the adult fly abdomen by an allograft method. We are using the allograft method to investigate tumorigenesis caused by compromised centrosome function in asymmetric stem cell division. Specifically, we are testing how does abnormal centrosomes affect the stem cell polarity and cell fate specification, genomic instability and how does this causes tumorigenesis of NSCs. Additionally, we are investigating the role of cellular stress response triggered due to dysfunctional centrosomes and its cooperation with tumorigenic signaling pathways in a cell autonomous or non autonomous manner. I will be discussing our findings addressing these questions.

1003V Using Drosophila Models to Dissect Biology and Signaling Mechanisms in Rare Drug Resistant Variants of Lung Cancer Sereene Kurzum1, William Marsiglia2, Chana Werther-Hecht1, Bruce Gelb1, Arvin Dar2, Tirtha Das1,2,3 1) The Mindich Child Health and Development Institute, Dept. of Pediatrics, Department of Genetics and Genomic Science, Icahn School of Medicine at Mount Sinai; 2) Department of Oncological Sciences, Icahn School of Medicine at Mount Sinai; 3) Dept. of Cell, Developmental, and Regenerative Biology, Icahn School of Medicine at Mount Sinai, New York NY USA

Animal models and clinical data have vastly improved our understanding of gene variants that lead to human disease. In low occurrence cancers, rare gene variants arise in tumors of patients receiving targeted therapy. We had previously developed a Drosophila model for a rare lung cancer arising from a fusion oncogene KIF5B-RET and showed that this fusion protein assembles a complex signaling hub composed of multiple receptor tyrosine kinases (RTK’s). Our finding predicted that drugs designed to inhibit the RET-kinase activity alone may be insufficient to fully suppress oncogenic signaling from this hub. Subsequently, multiple clinical trials have shown poor response of RET-inhibitor drug treatment in patients harboring KIF5B-RET fusions in their lung tumors. Recently a few drugs, like loxo-292, designed to specifically and potently inhibit RET-kinase have been developed. In our fly model, treatment of KIF5B-RET with loxo-292 prevented phosphorylation of RET and some components of the hub like EGFR, but not other components like FGFR. This result indicates a potential mechanism of how drug resistant compensatory pathways could arise even after treatment with RET-kinase selective inhibitors. Moreover, recent clinical studies have shown that KIF5B-RET positive lung cancer patients develop rare KIF5B-RET-variants while undergoing treatment with RET-kinase selective drugs, and that these drug resistant variants reduce efficacy of the treatment. We developed one such model, using the human KIF5B-RETG810R, and found that this variant enhanced RET-kinase domain phosphorylation and enhanced activation of various RTK dependent pathways of the KIF5B-RET hub. Indeed, none of the studied RTK pathways were suppressed when KIF5B-RETG810R fly tissues were treated with loxo-292. In summary, our current findings using Drosophila models indicate that RET-selective drugs like loxo-292 may not fully suppress signaling of KIF5B-RET or KIF5B-RETG810R variant. Our ongoing studies predict potential molecular mechanisms of drug resistance to RET-selective inhibitors in various cancers and highlight the value of combining whole animal Drosophila models with chemical genetics to study human disease.

1004V Rbf/E2F1-mediated transition from steroid-dependent to steroid-independent ecdysone receptor signalling in Drosophila prostate-like secondary cells Aashika Sekar1,2, Mark Wainwright2, Aaron Leiblich2, Claudia Mendes2, Clive Wilson2 1) Centre for Tumour Biology, Barts Cancer Institute, London, United Kingdom; 2) Department of Physiology, Anatomy and Genetics, University of Oxford, United Kingdom

Castration-resistant prostate cancer (CRPC) is an incurable, androgen-independent form of prostate cancer that emerges in the hormone-depleted environment of androgen deprivation therapy (ADT). However, CRPC growth still frequently depends on Androgen Receptor (AR) signalling. Unusually, loss of the tumour suppressor gene, Retinoblastoma (Rb), and consequent activation of transcription factor E2F1 have been linked to late-stage tumour progression to CRPC, rather than early-stage events. We have previously shown that binucleate secondary cells (SCs) of the Drosophila melanogaster male accessory gland (AG) share several functional and signalling similarities with human prostate epithelial cells. Upon mating, SC growth regulation switches from a steroid-dependent to a steroid-independent form of Ecdysone Receptor (EcR) control that induces genome endoreplication. Here, we demonstrate that the Drosophila Rb homologue, Rbf, and E2F1, as well as cell cycle regulators, Cyclin D (CycD) and Cyclin E (CycE), are key mediators of SC growth and endoreplication both in virgin and mated males. Importantly, we show that the CycD/Rbf/E2F1 axis requires the EcR, but not ecdysone, to trigger CycE-dependent endoreplication and associated growth in SCs after mating, mirroring changes in CRPC. We also demonstrate that excess Rbf activity reversibly suppresses binucleation in adult SCs. Overall, our work reveals mechanistic parallels between the physiological switch to hormone-independent EcR signalling in SCs and the pathological switch seen in CRPC, and suggests that the latter may represent the dysregulation of a
Currently unidentified physiological process, which permits AR signalling when androgen levels are low.

1005V  **Tissue specific knockdown of Scribble induces tumor progression and metastasis in Drosophila**  Jyotsna Singh, Saripella Srikrishna Department of Biochemistry, Institute of Science, Banaras Hindu University, Varanasi, UP 221005, India

Cancer is one of the deadly diseases worldwide and the cancer-related deaths across the globe will rise to over 11 million by 2030 and 27 million by 2050. Metastasis is the major cause of cancer associated mortality in humans. Metastasis is the process by which an uncontrolled malignant growth of tumor cell leaves the primary tumor and invades into distant regions from its site of origin, leading to formation of secondary tumors. Epithelial cell polarity regulator gene, *Scribble*, implicated in variety of cancers, plays major role in cancer progression and metastasis. *Drosophila* pupa is a valuable metastasis model for studying live dynamics of tumor micro environment and metastasis events by combing endogenous fluorescence probes and fluorescence microscopy. Here, we have generated a metastatic cancer in *Drosophila* by knockdown of *scribble* in wing imaginal discs using UAS-GAL4 system. We have found that *scribble* knockdown induced overgrowth of wing imaginal tissue leading to metastatic secondary tumor formation and absolute lethality at early pupal stage. Tumor progression and metastasis phenotypes in *Drosophila* were scored for 84 hours after puparium formation (APF) and recorded the invasion of GFP labeled tumor cells. Tumor bearing pupae showed spreading of secondary tumors cells throughout from anterior to posterior regions of the pupae while wild type pupae showed GFP expression restricted to developing wing discs areas only. Further, we checked mRNA expression of metastatic biomarkers at different time point of pupal development. The up-regulation of MMPs and reduced expression of *E-cadherin* were confirmed the increased cell proliferation and metastasis in *Scrib* knockdown tumorous tissues of pupae. Further study is in progress in this direction.

**Keywords:** *Drosophila*, Cancer, tumor, *scribble*, Metastasis

1006V  **Septins regulate heart contractility through modulation of cardiomyocyte store-operated Ca²⁺ entry**  Benjamin Tripoli, Courtney Petersen, Jeremy Smyth SUUS - Bethesda, MD

Proper calcium handling in cardiomyocytes is paramount to maintaining cardiomyocyte contractility and heart function. Recent data from our lab and others in the field have established that store operated calcium entry (SOCE) is an essential component of cardiomyocyte calcium regulation. SOCE refers to the pathway in which endoplasmic/sarcoplasmic reticulum (ER/SR) calcium store depletion causes influx of calcium to replenish the stores. The pathway is mediated by STIM proteins, which act as calcium sensors in the SR/ER, and ORAI calcium influx channels in the plasma membrane. Both upregulation and suppression of SOCE in cardiomyocytes causes disruption of normal cardiac function, demonstrating that proper regulation of SOCE in cardiomyocytes is essential. However, the mechanisms responsible for the regulation of SOCE in cardiomyocytes is not yet fully understood. To this end, septin GTPases have emerged as regulators of the SOCE pathway in non-cardiac tissues, with septin 1, 2, or 4 suppression resulting in SOCE suppression and septin 7 (PNUT) suppression resulting in SOCE upregulation. Importantly, the role of septins in cardiomyocytes is nearly completely unknown. Through intravital imaging analysis of *Drosophila* heart contractility, we now show that cardiomyocyte specific RNAi-based depletion of septins 1, 2, or 4 results in dilated cardiomyopathy nearly identical to that caused by SOCE suppression. We further show that co-expression of septin 2 RNAi with a constitutively active Orai channel suppresses the septin 2 phenotype, supporting the hypothesis that septin 2 associated dilated cardiomyopathy is due to SOCE suppression. Interestingly, septin 7 suppression resulted in hypertrophic cardiomyopathy similar to that caused by SOCE upregulation, and micro-computerized tomography (microCT) analysis of heart size further confirmed the hypertrophic phenotype of septin 7 depleted hearts. This septin 7 phenotype was suppressed by Orai depletion, again supporting a role for SOCE dysregulation in the septin depletion-mediated cardiac phenotypes. We are currently developing tools to analyze Stim and Orai localization, sarcomere organization, and t-tubule architecture to further investigate septins and SOCE regulation in cardiomyocytes.

1007V  **Focus on the foci: Investigating the role of HDAC4 aggregation in neuronal development in Drosophila melanogaster**  Hannah Hawley, Helen Fitzsimons School of Natural Sciences, Massey University, New Zealand

Dysregulation of histone deacetylase 4 (HDAC4) expression and subcellular distribution has been observed in a number of neurodevelopmental and neurodegenerative diseases, and in our *Drosophila melanogaster* model, HDAC4 overexpression impairs neuronal development and long-term memory. Interestingly, this is associated with minimal transcriptional changes. Upon increased abundance in nuclei, we observe HDAC4 aggregation into punctate foci, and therefore hypothesise that neuronal dysfunction mediated by HDAC4 overexpression is a result of aggregate formation.

The glutamine-rich N-terminus of HDAC4 forms an alpha helix which assembles into an unstable tetramer. To investigate whether HDAC4 aggregates contribute to neurodevelopmental deficits, transgenic *Drosophila* were generated which express HDAC4 mutants harbouring structure-guided substitutions of key amino acids important in mediating tetramerization. Expression of these mutant HDAC4 constructs significantly reduced aggregate formation in neuronal nuclei (ANOVA, p < 0.01), and this correlated with a significant reduction in defects in axon morphogenesis (Fisher’s
exact, \( p < 0.01 \) and photoreceptor development (ANOVA, \( p < 0.01 \)), as compared to wild-type HDAC4. These data suggest HDAC4 aggregation is at least in part responsible for neurodevelopmental and neurodegenerative disease in which HDAC4 is aberrantly expressed, and warrants further studies into the composition of these aggregates as well as strategies to mitigate their formation.

**1008V** De novo variants in SUPT16H are associated with developmental delay, intellectual disability, epilepsy and facial dysmorphism

We identified two individuals from two families carrying de novo missense variants in SUPT16H. The patients exhibit global developmental delay, intellectual disability, epilepsy, facial dysmorphism and structural abnormalities of the brain. SUPT16H is conserved across species and encodes the large subunit of the FACT complex which functions as a nucleosome organizer during transcription. Bioinformatic predications from population genomics databases indicate that SUPT16H is highly constrained for loss-of-function variants in humans and both missense variants p.T171I and p.G808R have not been reported to date. We employed Drosophila to characterize these two variants. We show that loss of the fly orthologue, dre4, causes early lethality. Tissue-specific RNAi-mediated knockdown of dre4 in the eye or wing disc leads to loss of eye and wing, respectively. RNAi-mediated knockdown in the nervous system also causes severe defects. Expression of the human reference SUPT16H transgene partially rescued the loss-of-function phenotypes in wing, eye and nervous system. However, the p.T171I or p.G808R variants are much less efficient at rescuing these phenotypes, indicating that they are partial loss-of-function alleles. We show that dre4 functions in a cell-autonomous manner, rather than in an endocrine fashion dependent on ecdysteroid production as previously proposed. Altogether, our data indicate that de novo loss-of-function variants in SUPT16H are associated with neurological features.

**1009V** Drosophila models reveal nuclear shape and lamin localization patterns that differentiate clinically distinct laminopathies

Laminopathies are a collection of diseases caused by mutations in the human LMNA gene, which include muscular dystrophy and lipodystrophy, an adipose tissue disorder. The symptoms and severity of laminopathies vary depending on the genetic background of the individual. The LMNA gene encodes lamin A/C, intermediate filaments that form a meshwork underlying the inner nuclear membrane. Generating Drosophila models allows for analysis of specific mutations in a defined genetic background and the ability to express mutant lamins in specific tissues. Thus, we modeled two LMNA mutations that cause distinct clinical symptoms in Lamin C, the Drosophila orthologue of human LMNA. Both mutations cause a single amino acid substitution in the conserved Ig-like fold domain of lamins. Lamin A/C Arg482Trp causes lipodystrophy; lamin A/C Arg527Pro causes Emery-Dreifuss muscular dystrophy and lipodystrophy. Using the Gal4/UAS system, wild-type and mutant lamins were expressed in larval body wall muscles and cardiac tissue. When expressed in the larval body wall muscles, lamin Arg527Pro aggregated within the cytoplasm and caused reduced larval motility. By contrast, lamin Arg482Trp caused abnormally shaped nuclei and had no effect on larval motility. Surprisingly, both mutants caused death at the pupal stage. When expressed in cardiac tissue, lamin Arg527Pro caused a shortened adult lifespan. By contrast, flies with cardiac-specific expression of lamin Arg482Trp exhibited a lifespan similar to that of flies expressing wild-type lamin. Thus, the Drosophila models recapitulated aspects of the human disease, with only Arg527Pro affecting muscle function. In addition, the distinct intracellular localization patterns and differing effects on nuclear morphology provide an explanation for how mutations that affect the same lamin domain cause distinct clinical pathology.

**1010V** Generating Drosophila melanogaster isofemale lines tolerating extreme oxygen conditions

Hyoxic and oxidative stress are common pathological elements in many diseases. In order to understand the mechanisms underlying mechanisms regulating tolerance or susceptibility to hyoxic or oxidative stress-induced injuries, we developed two Drosophila melanogaster populations that can live perpetually in severe, normally lethal, hyoxic or hyperoxic environments through experimental evolution. From these populations, we generated a Panel of Low Oxygen Tolerant (PLOT) and a Panel of High Oxygen Tolerant (PHOT) isofemale lines (total 79 PLOT and 65 PHOT lines) to identify and study specific genetic mechanisms that are responsible for the hyoxia, or hyperoxia, tolerant trait. These isofemale lines showed significantly enhanced hyoxia or hyperoxia tolerance as compared to their wildtype parental and the wildtype DGRP isofemale lines. For example, 95% of the PLOT lines showed hyoxia tolerance with eclosion rate >50% at 5% \( O_2 \) level. In contrast, none of the parental lines and only 15% of the DGRP lines had similar eclosion rates under the same hyoxic condition. Furthermore, under the lethal condition with 4% \( O_2 \), the PLOT lines exhibited a wide range of eclosion rate from 1% to 80% with a clear pattern of Gaussian-Distribution. These PLOT and PHOT flies may provide...
us a unique opportunity to study the molecular basis of stress-directed evolution, such as the role of genetic variations in the protein coding regions and functional DNA elements in hypoxia, or hyperoxia, adaptation. In addition, these PLOT and PHOT lines can also be used to study phenotypic plasticity as well as genetic-epigenetic interactions in hypoxic or oxidative environments.

1011V  Effect of Circadian Rhythm Disruption on DNA Double Strand Break Repair Pathway Choice Lydia Bergerson, Caleb Fitzmaurice, Tyler Knudtson, Halle McCormick, Alder Yu University of Wisconsin - La Crosse

Long-term night shift work has been associated with increased risk of certain cancers, although this link is not consistently supported. The most obvious difference between shift workers and non shift workers is increased disruption of circadian rhythms in shift workers. If circadian rhythm disruption increases use of mutagenic DNA double strand break repair (DSBR) pathways such as non-homologous end joining (NHEJ), that would provide a potential mechanistic link between shift work and cancer initiation.

To test the hypothesis that circadian rhythm disruption increases use of NHEJ DSBR, we used the Rr3 system. The Rr3 system is based on a chromosomally integrated DsRed gene that is rendered nonfunctional by an inserted I-SceI site flanked by a short direct repeat. Rr3 containing flies are crossed to I-SceI expressing flies and DSBs are induced in the F1 generation premeiotic germline. Individual repair events are recovered and scored in the F2 generation by appropriate crosses. Repair by base pairing at the direct repeats (single strand annealing, SSA) reconstitutes expression of the DsRed gene. Repair by NHEJ does not reconstitute DsRed expression. We compared relative usage of SSA and NHEJ repair pathways between flies maintained on 12:12 and 8:8 light:dark schedules. Disruption of circadian rhythms by the 8:8 schedule was confirmed by actimetric analysis. We found no significant difference in NHEJ or SSA repair pathway usage between 12:12 and 8:8 flies. Current experiments are investigating the effect of circadian rhythm disruption on use of homologous recombination repair.

1012V  Elucidation of the role of IFT52 associated with a novel skeletal ciliopathy using in vitro and Drosophila systems Vishal Singh Guleria, Rahul Parit, Girisha M Katta, Priyanka Upadhyay Kasturba Medical College, Manipal, India

Primary cilia (PC) are tubular antenna like structures present on the mammalian cell surface transducing extracellular cues like hormones, growth factors and mechanical stimuli. Also, playing an important role in cellular development, differentiation and homeostasis. These microtubular structures are maintained by intraflagellar transport (IFT) machinery involved in bidirectional transport of ciliary proteins. IFT52 is a component of the IFT complex and associated with a novel skeletal ciliopathy, short-rib thoracic dysplasia 16 with or without polydactyly (MIM# 617102). Cellular mechanism of IFT52 function in modulating primary cilia biogenesis, signalling and its dependent regulation of osteogenesis and chondrogenesis is not known.

In our study we have characterized PC during osteogenic (OS) and chondrogenic (CH) differentiation in vitro and elucidated the molecular mechanism of IFT52 function in a PC dependent manner during osteogenesis and chondrogenesis in C3H10T1/2 and ATDC5 cell lines and Drosophila systems. OS and CH differentiation was induced in both cell lines and ascertained by assaying for known differentiation markers, calcium and glycosaminoglycan deposition in the extracellular matrix. Ift52 transcript levels were studied using qRT-PCR. Additionally, we characterized the IFT52 ortholog, Osm6 in Drosophila by IFT52 knockdown (KD) in embryonic neuroblasts by RNA interference using pan neuronal Gal4 drivers.

OS differentiation was evident at 7 days following induction via alizarin red (AR) staining to reveal extracellular matrix (ECM) mineralization. CH differentiation was first detected at 14 days post induction by sulfated proteoglycan deposition and ECM mineralization using alcian blue and AR staining. We found Ift52 transcript levels were upregulated at 14 and 21 days of OS differentiation, while in CH differentiation they were initially upregulated at 7 days and subsequently depressed. Ift52 KD resulted in reduced protein expression and significant reduction of PC prevalence, length and OS differentiation.

IFT52 KD in embryonic neuroblasts of Drosophila melanogaster resulted in significant loss of chordotonal function like locomotion and hearing in adults and larva. Also, IFT52 KD embryos showed structural ciliary defects like profound depletion of the Ich5 cilia in embryos. Our findings are anticipated to remit a comprehensive molecular characterization of the Ift52 gene and enhance our understanding of its allied human anomalies.

1013V  Genetic investigation of the Endolysosomal Network in a Drosophila model of Alzheimer’s disease Sher Li Tan1,2, Robert Richards1, Tim Sargeant2, Louise O’Keefe1,2 1) The University of Adelaide, Adelaide, South Australia, Australia; 2) South Australian Health and Medical Research Institute, Adelaide, South Australia, Australia

Introduction: Alzheimer’s disease (AD) is the most common form of dementia, which affects memory, behaviour and daily functioning of affected patients with its pathological characteristics consisting of amyloid-beta (AB) plaques and phosphorylated tau accumulation in the brain. The mechanisms underlying the disease remain unknown and no effective treatments are available to prevent disease progression. Although the genetic contribution in AD patients is not fully understood, genome-wide association studies (GWAS) have identified several loci associated with increased AD risk
in genes within the endolysosomal network (ELN). However, the fundamental mechanism of AD disease progression regarding ELN is not well defined. Therefore, this project aims to identify and characterise the contribution of individual ELN genes to neuronal dysfunction in a Drosophila model for AD.

**Method/Approach:** Co-expression of full length amyloid precursor protein and beta-site APP cleaving enzyme (APP+BACE) has been investigated in Drosophila to determine whether flies are able to model symptoms observed in AD patients. The advantage of this APP+BACE model is that it will ensure correct spatial localisation of APP and its proteolytic fragments within the ELN. Phenotypes generated form the basis for genetic modification analyses of individual ELN genes.

**Result:** Our findings show that co-expression of APP+BACE causes retinal degeneration in the adult eye and this can be modified by altered expression of Rab5 or Rab7 ELN genes. In addition, neuronal expression of APP+BACE in flies has led to progressive loss of motor functions, behavioural problems, and a reduced lifespan.

**Conclusion:** Using this system, we can characterise the genetic and molecular contribution of individual ELN genes to APP+BACE toxicity.

**1014V  Drosophila models of SNRNP200-retinitis pigmentosa exhibit retinal apoptosis and loss of photoreceptor function** Sara Mayer, Quinton Christensen, Arlene Drack, Lori Wallrath University of Iowa

Retinitis pigmentosa (RP) represents a collection of genetic eye disorders that cause gradual loss of photoreceptors. The initial phases of RP involve loss of rod photoreceptor function, which leads to loss of cone photoreceptor function and eventual blindness. RP affects 1:4,000 people worldwide and is either syndromic or non-syndromic. RP33 is a non-syndromic form of RP caused by mutations in the SNRNP200 gene, which encodes a core component of the spliceosome. SNRNP200 possesses helicase activity that plays a critical role in pre-mRNA splicing. SNRNP200 is expressed in nearly every cell type, yet defects are only observed in the retina. To understand the function of SNRNP200 in vision, we turned to Drosophila. The Drosophila orthologue of human SNRNP200 is lethal(3)72Ab (or Brr2), which we refer to as dSNRNP200. Human SNRNP200 and fly dSNRNP200 have 74% amino acid identity and 89% similarity. Null mutations in dSNRNP200 possesses helicase activity that plays a critical role in pre-mRNA splicing.

**Method/Approach:** Using this system, we can characterise the genetic and molecular contribution of individual ELN genes to APP+BACE toxicity.

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that it potentiates the mutagenic effect of 4NQO. In addition, it was observed that it does not have an antimutagenic effect either.

1016A Using expansion microscopy to examine parasegmental boundaries at nanoscale resolution Samia Parveen, Ian Millerschultz, Adam Pare University of Arkansas

Over the course of animal development, distinct pools of epithelial cells are maintained by compartment boundaries. For example, in the early Drosophila embryo, the formation of compartment boundaries at regular intervals along the anterior-posterior axis establishes the segmental body plan of the future animal. These so-called “parasegmental boundaries” (PSBs) are composed of highly stable cell-cell contacts that are under increased tension compared with typical contacts, and they resist the movement of cells between adjacent compartments. However, it is still unclear what differentiates PSB contacts from normal contacts at a molecular level. The force-generating cytoskeletal proteins F-actin and myosin are more highly enriched at PSBs compared with normal contacts, and evidence suggests that E-cadherin-based adherens junctions may also be differently arranged at PSBs. However, the fine details of these structures are below the resolution limit of standard confocal microscopy. To get past this resolution limit without the need for expensive super-resolution microscopes, we have adapted existing expansion microscopy (ExM) protocols for use in the early Drosophila embryo. ExM involves the isotropic physical expansion of a sample through chemical treatments to swell the embryo in a controlled manner, maintaining the relative positions of all molecules. We are using this technique to examine the ultrastructural distribution of adherens junction and cytoskeletal proteins at PSBs to see how they differ from normal contacts. We are also examining two known upstream regulators of PSB formation, Tartan and Ten-m, to better characterize their subcellular localization patterns in the early embryo. We expect these studies will give us a better understanding of how the three-dimensional arrangement of the cellular effectors of tension and adhesion—as well as the cell surface proteins that regulate them—gives PSBs their unique properties.

1017B Adaptation of the CRISPR-Sirius tool for imaging the genome in Drosophila ovaries Erica Berent, Haosheng Li, Oscar Bautista, Joseph Terry, Bowen Man, Nicole Crown Case Western Reserve University

Traditional fluorescence in situ hybridization (FISH) techniques are destructive to chromatin structure, and they are labor intensive when combined with immunofluorescence. In order to label and image specific loci without relying on FISH, we have adapted the genetically encoded CRISPR-Sirius system (Ma et al 2018) for use in the Drosophila ovary. This system consists of three components: a nuclease dead Cas9, a modified guide RNA containing MS2 stemloops, and an MCP-fluorescent protein fusion that binds the MS2 stemloops. To adapt this system into flies, we first used site directed mutagenesis to mutate the catalytic sites of Cas9 in the nanos::Cas9 vector (Port et al 2014). Second, we cloned eight copies of the MS2 step loop (Ma et al 2018) into the core guide RNA in pCFD3 (Port et al 2014) to generate pCFD3-Sirius. Lastly, we used an existing nanos::MCP-GFP fusion protein stock (BDSC#63821). To test our system, we cloned a guide RNA targeting the 359 bp repeat into our new pCFD3-Sirius vector. We then crossed the pCFD3-Sirius-359 repeat flies to nanos::GFP-MCP flies and immunostained the ovaries with an antibody against GFP. We will present data showing that our system is able to target repetitive sequences in the Drosophila genome and we will present our progress in targeting unique non-repetitive sequences.

1018C An improved organ explant culture method reveals stem cell lineage dynamics in the adult Drosophila intestine Marco Marchetti, Chenge Zhang, Bruce Edgar Huntsman Cancer Institute, University of Utah, Salt Lake City, UT

In recent years, live-imaging techniques have been developed for the adult midgut of Drosophila melanogaster that allow temporal characterization of key processes involved in stem cell and tissue homeostasis. However, current organ culture techniques are limited to imaging sessions of <16 hours, an interval too short to track dynamic processes such as damage responses and regeneration, which can unfold over several days. Therefore, we developed a new organ explant culture protocol capable of sustaining midguts ex vivo for up to 3 days. This was made possible by the formulation of a culture medium specifically designed for adult Drosophila tissues with an increased Na+/K+ ratio and trehalose concentration, and by placing midguts at an air-liquid interface for enhanced oxygenation. We show that midgut progenitor cells can respond to gut epithelium damage ex vivo, proliferating and differentiating to replace lost cells, but are quiescent in healthy intestines. Using ex vivo gene induction to promote stem cell proliferation, we demonstrate that intestinal stem lineages can be traced through multiple cell divisions using live imaging. Both asymmetric and symmetric divisions can be identified in the reconstructed lineages. We find that daughter cells of asymmetric divisions remain in close proximity of each other, while the progeny of symmetric divisions actively move apart, with implications for cell differentiation and tissue organization. We show that the same culture set-up is useful for imaging adult renal tubules and ovaries for up to 72 hours. By enabling both long-term imaging and real-time ex vivo gene manipulation, our simple culture protocol provides a powerful tool for studies of epithelial biology and cell lineage behavior.

1019A Graphene Enabled Optical Cardiac Control of Drosophila Abby Matt1, Hongwu Liang2, Matthew Fishman3, Andrey Komarov1, Xinyuan Zhang1, Jing Men1, Elena Gracheva1, Alex Savtchenko2, Chao Zhou1 1) Washington University in
Current methods of cardiac stimulation, such as electrical stimulation, ultrasound stimulation, and cardiac optogenetics have certain risks or drawbacks. The use of invasive surgical procedures to implant electrodes and electrochemical reactions caused by electrode-tissue contact may cause serious detrimental effects. In this study, we implement a stimulation technique in the fruit fly Drosophila melanogaster heart that uses graphene injection to optically modulate cardiac activity. Graphene is a single layer arrangement of carbon molecules with unique electrical properties. We prepare a homogenized graphene solution for injection into the heart tube with a combination of phosphate buffer saline (PBS) or artificial hemolymph (AH). PBS and AH were also injected as control solutions. From the graphene solution, optical stimulation excites graphene particles that coat the heart tube, which in turn creates electrical stimulation, changing the membrane potential of heart tube muscles. Through the use of an Optical Coherence Microscopy (OCM) system, custom red light stimulation, and a machine learning based model for analysis, we demonstrate non-contact cardiac stimulation after injecting graphene-based biointerfaces into the heart. Further, the analysis of heart mechanics, such as fractional shortening, indicates that there is no change to the way the heart moves when stimulated. This system is promising for studies of various gene-related cardiovascular diseases or for testing drugs. For example, it could enable studies of various cardiovascular differences in Drosophila caused by human ortholog gene mutations.

1020B An expanded toolkit for gene tagging using synthesized homology donor constructs for CRISPR mediated homologous recombination Oguz Kanca1,2, Jonathan Zirin3, Yanhui Hu4, Burak Tepe1,2, Debdeep Dutta1,2, Wen-Wen Lin1,2, Liwen Ma1,2, Ming Ge1,2, Zhongyuan Zuo1,2, Lu-Ping Liu3, Robert W. Levis4, Norbert Perrimon3,4, Hugo J. Bellen1,2 1) Department of Molecular and Human Genetics, Baylor College of Medicine, Houston, TX; 2) Duncan Neurological Research Institute, Texas Children Hospital, Houston, TX; 3) Department of Genetics Harvard Medical School, Boston, MA; 4) Department of Embryology, Carnegie Institution for Science, Baltimore, MD; 5) Department of Neuroscience, Baylor College of Medicine, Houston, TX; 6) Howard Hughes Medical Institute, Harvard Medical School, Boston, MA.

Gene traps and protein traps are valuable genetic tools to study gene function. The majority of these constructs are inserted in the genome as artificial exons, with a Splice Acceptor (SA)-effector-Splice Donor (SD) configuration. The artificial exon must be inserted in an intron between two coding exons to function as a gene or protein trap. About 50% of the Drosophila genes do not have a suitable coding intron. We developed methods to replace the coding region of these genes with a Kozak-GAL4-3XP3GFP cassette creating a null allele, while expressing GAL4. We have generated >100 KozakGAL4 alleles to date. Analysis of the expression patterns in 3rd instar larval brains shows that ~80% of the tested KozakGAL4 alleles can drive expression of UAS reporter genes with discernible patterns. We selected KozakGAL4 alleles that drive reporter expression in restricted patterns and used publicly available single cell RNA sequencing (scRNAseq) data to find other genes that are co-expressed with the targeted gene. T2AGAL4 alleles of such genes, identified by scRNAseq, drive expression of reporter genes in patterns overlapping with the KozakGAL4 targeted genes. scRNAseq data serve as an independent means to verify the accuracy of KozakGAL4 expression patterns for reporting the expression pattern of the targeted gene. Hence, KozakGAL4 alleles serve as an alternative approach to artificial exon-based methods for genes lacking a suitable intron.

We also designed and generated new homology donor vector backbones to facilitate the generation of homology donor constructs. We showed that in vivo linearization of the donor constructs improves the efficiency of incorporation of large constructs (>5kb) using short homology arms (200 bps). Short homology arms make it feasible to commercially synthesize homology donors and minimize the cloning steps in donor construct generation. We have designed new vector backbones that include both gene specific sgRNA(s) and an sgRNA to linearize the homology donor together with the homology arms. This strategy obviates cloning separate sgRNAs and ensures delivery of all the components for targeting a gene in a single plasmid. We show that using this strategy greatly facilitates cloning of the donor constructs and increases the successful transgenesis rate to 80%. These upgrades, together with the KozakGAL4 strategy, will enable efficient targeting of 80% of the conserved fly genes by the Gene Disruption Project.

1021C Nebulos without white: annotated long-read genome assembly and CRISPR/Cas9 genome engineering in D. nebulosa Christopher Sottolano1, Nicole Revaitis1, Anthony Geneva1,2, Nir Yakoby1,3 1) Center for Computational and Integrative Biology, Rutgers University, Camden, NJ; 2) Department of Biology, Rutgers University, Camden, NJ.

The plethora of morphological diversity among Drosophila species presents an opportunity to study the mechanisms underlying the molecular evolution of tissue patterning and morphologies. One of the challenges in investigating these species is that unlike the tremendous number of molecular and genetics tools available for Drosophila melanogaster research, many other species do not have sequenced genomes. In addition, the available molecular tools to manipulate genes are very limited. Here, we focus on D. nebulosa due to the ease of rearing, its differences in tissue patterning and eggshell morphologies, and outstanding organisal behaviour when compared to D. melanogaster.

As a first step to develop the fly as a new genomic and genetic resource, we generated a high-quality annotated genome assembly of D. nebulosa, using PacBio long-read sequencing. We utilized the assembly to successfully disrupt the white gene via CRISPR/Cas9, but were unable to integrate the Cas9 gene using homology-directed repair.
Interestingly, unlike many other Drosophila species, D. nebulosa males null for the white gene did not appear to court females. We conclude, gene disruption via CRISPR/Cas9 genome engineering is a useful tool in D. nebulosa. However, a simple selectable marker which does not impair vision is needed to replace the commonly used white-eye phenotype.

1022A A nickase Cas9 gene-drive system promotes super-Mendelian inheritance in Drosophila melanogaster. Sara Sanz Juste, Victor Lopez del Amo, Valentino Gantz University of California, San Diego

CRISPR gene drive systems can potentially modify an entire population due to their self-propagation capacity, representing a powerful tool to combat public health issues such as mosquito-borne diseases. Gene drive elements bias Mendelian inheritance (50%) towards super-Mendelian rates (~100%), allowing them to spread through populations. Traditional gene drives can replace wildtype alleles by performing DNA double-strand breaks and subsequent homology-directed repair (HDR). When the HDR is not precise, mutations at the target site can generate resistant alleles, preventing a gene drive’s efficient spread.

We hypothesize that simultaneous paired-nicks targeting two adjacent DNA regions should generate a staggering double-strand break, and the subsequent DNA repair by HDR will promote the gene-drive super-Mendelian inheritance. Here, we developed a proof-of-concept nickase-based gene drive strategy that enables super-Mendelian inheritance of an engineered allele using Drosophila melanogaster. Additionally, we analyzed the quality of potential resistant alleles formed by our nickase gene-drive system and compared them to the traditional gene drive to explore potential advantages. In fact, we demonstrate for the first time that both nD10A and nH840A promote HDR in the germline using CRISPR gene drives, yielding comparable super-Mendelian inheritance rates to traditional gene drives. Furthermore, the double-nicking nature of this arrangement produced large deletions when using the nickase H840A combined with paired-gRNAs in a PAM-in configuration. This outcome provides potential advantages to target essential genes.

Indeed, our nickase gene-drive design should expand our alternatives for gene drive applications and other germline editing purposes in a broad range of organisms.

1023B High-Resolution Imaging Method with Standardized Conditions Facilitates Reproducible, Spatial, Quantitative Data Heidi Pipkin1, Adam Smiley2, Andrew Arsham1 1) Bemidji State University; 2) University of Minnesota

Collection of reproducible high quality quantitative data on whole flies is challenging, especially in training environments where data collection is distributed among many individuals. Common adjustable light sources for dissecting microscopes create glare, shadow, and irreproducible lighting conditions; microscope-mounted cameras are optimized for low noise and high sensitivity but are often low resolution and can only record 1-2 flies at a time; and only a small portion of the fly is in focus. To address these limitations, we assembled a high-resolution imaging method with fixed lighting, staging, and exposure conditions. Whole flies are frozen and imaged without additional fixation or mounting medium. A 3D printed stage standardizes fly placement and reduces preparation time. Batch collection and computer automation of focus stacking eliminates inter-image and inter-operator variability eliminating the need for complex, irreproducible, and ethically murky image manipulation to adjust color, glare, and similar artifacts. A 50MP full frame digital camera, computer-controlled stepper motor, and focus-stacking software generates a single image of dozens of flies with all ommatidia, bodies, wing veins, and bristles in focus. These standardized composite images facilitate a many-fold reduction in imaging time and precise quantitative comparison of phenotypes involving body and organ color, size, shape, and pattern. Semi-automated data extraction and analysis using ImageJ and R further facilitate reproducibility, code-sharing, and flexibility and reduce the confusion and workflow complexity created by intermediary data products (crops, filters, masks, format conversions, etc.). In addition to being accessible to resource-limited labs, these methods are classroom-friendly and support undergraduate learning outcomes in image analysis, data visualization, computational biology, and statistics.

1024C Developing a High Throughput Drug Induced Phenomics and Transcriptomic Assessment Robert Courville, Mahtab Jafari University of California, Irvine, Irvine, CA

Background: Drosophila melanogaster may serve as an excellent pre-clinical model system in drug discovery. Developing high throughput platform using Drosophila as a model system in drug discovery presents an efficient and cost-effective in vivo pre-clinical platform to screen and evaluate therapeutics, at the phenotypic and transcriptomic levels.

Methods: We developed two high throughput screening platforms to evaluate drug induced phenomic changes by two pharmaceuticals with distinct pharmacological properties in an outbred Drosophila population: caffeine (a stimulant) and haloperidol (a sedative). The evaluated phenotypes were fecundity and behavioral changes; locomotion, feeding patterns, and sleep. To evaluate fecundity, flies were divided into mating pairs and transferred into vials with removable caps containing the drug/yeast solution. After 24 hours of mating, digital images of vial caps were created using a scanner in groups of 40 by treatment, and the images were uploaded into ImageJ where a custom macro autonomously counted the eggs. To evaluate behavioral changes, we optimized the Activity Recording Cafe (ARC), a machine-vision
system that allows for automated behavioral measurements. Flies were sexed into groups of 30 and habituated in ARC chambers for 24 hours. Machine vision settings were calibrated for optimal frame tracking and data were collected for 24 hours autonomously. Behaviors were measured at days 14 and 20 from eclosion. Transcriptomic analysis will be done at the UC Irvine Genomics High-Throughput facility using heads, ovaries, and testes.

**Results:** Both drugs impacted fecundity in an age specific manner. Caffeine increased fecundity in days 19-28 (p<0.05) but had no impact on fecundity on days 14-18. Haloperidol decreased fecundity in days 14-18 (p<0.05) and had no impact on fecundity in other ages. Caffeine increased fly activity and feeding relative to controls (p<0.05), and decreased sleep (p<0.05). Haloperidol showed decreased activity at both ages (p<0.05), but no significant differences in feeding or sleep. All phenomic assays are currently being replicated. Transcriptomic analysis is in progress. The efficiency (time and cost) of these high-throughput assays will be compared with conventional assays and the data will be presented on the poster.

**Conclusion:** The phenomic data show that the high throughput screening platforms were sensitive enough to detect drug induced phenomic changes.

1025A A novel transposable element based authentication protocol for *Drosophila* cell lines Daniel Mariyappa1, Douglas Rusch1, Shunhua Han2, Arthur Luhur1, Danielle Overton1, David Miller1, Casey Bergman2, Andrew Zelhof2 1) Indiana University; 2) University of Georgia

*Drosophila* cell lines are used by researchers to investigate various cell biological phenomena. It is crucial to exercise good cell culture practice. Poor handling can lead to both inter- and intraspecies cross-contamination. Prolonged culturing can lead to introduction of large- and small-scale genomic changes. These factors, therefore, make it imperative that methods to authenticate *Drosophila* cell lines are developed to ensure reproducibility. Mammalian cell line authentication is reliant on short tandem repeat (STR) profiling, however the relatively low STR mutation rate in *D. melanogaster* at the individual level is likely to preclude the value of this technique. In contrast, transposable elements (TE) are highly polymorphic among individual flies and abundant in *Drosophila* cell lines. Therefore, we investigated the utility of TE insertions as markers to discriminate *Drosophila* cell lines derived from the same or different donor genotypes, divergent sub-lines of the same cell line, and from other insect cell lines. We developed a PCR-based next-generation sequencing protocol to cluster cell lines based on the genome-wide distribution of a limited number of diagnostic TE families. We determined the distribution of five TE families in S2R+, S2-DRSC, S2-DGRC, Kc167, ML-DmbG3-c2, mnb2, CME W1 Cl.8+, and OSS *Drosophila* cell lines. Two independent downstream analyses of the NGS data yielded similar clustering of these cell lines. Double-blind testing of the protocol reliably identified various *Drosophila* cell lines. In addition, our data indicate minimal changes with respect to the genome-wide distribution of these five TE families when cells are passaged for at least 50 times. The protocol developed can accurately identify and distinguish the numerous *Drosophila* cell lines available to the research community, thereby aiding reproducible *Drosophila* cell culture research.

1026B Genetic barcoding for single cell transcriptomics and population behavioral assays Jorge Blanco Mendana, Margaret Donovan, Benjamin Auch, Daryl Gohl University of Minnesota, Minneapolis, MN

Advances in single cell sequencing technologies have provided novel insights into the dynamics of gene expression throughout development, been used to characterize somatic variation and heterogeneity within tissues and are currently enabling the construction of transcriptomic cell atlases. However, despite these remarkable advances, linking anatomical information to transcriptomic data and positively identifying the cell types that correspond to gene expression clusters in single cell sequencing data sets remains a challenge. We are developing a straightforward genetic barcoding approach that takes advantage of the powerful genetic tools available in Drosophila to allow *in vivo* tagging of defined cell populations. This method, called Targeted Genetically-Encoded Multiplexing (TaG-EM), involves inserting a DNA barcode just upstream of the poly-adenylation site in a Gal4-inducible UAS-GFP construct, so that the barcode sequence can be read out during single cell sequencing, labeling a cell population of interest. By creating many such independently barcoded fly strains, TaG-EM will enable a number of potential applications that will improve the quality and information content of single cell transcriptomic data. Potential applications of this method include positive identification of cell types in cell atlas projects, barcoding of experimental timepoints, conditions, or replicates, and detection and elimination of multiplets through detection of multiple barcodes. Furthermore, we demonstrate that the barcodes from TaG-EM fly lines can be read out using next-generation sequencing to facilitate population-scale behavioral measurements. Thus, TaG-EM will enable many types of large-scale behavioral screens in addition to improving the ability to reliably annotate cell atlas data, expanding the scope of single cell transcriptomic experiments, and improving the robustness of such data by facilitating inclusion of replicates.

1027C RNA viral metagenomics of 100-year-old *Drosophila melanogaster* museum specimens Alexandra Keene, Mark Stenglein Colorado State University, Fort Collins, CO

Paleogenomic sequencing has enabled the reconstruction of extinct genomes and shed light on ancient host-pathogen interactions. But sequencing has focused for the most part on ancient DNA rather than ancient RNA, in part because
of a belief that RNA is fragile and short-lived. In this study, I experimentally assessed the stability of single and
double-stranded RNA over months in dried insects. I also optimized methods for non-destructive extraction of RNA
from pinned D. melanogaster museum specimens that had been collected over the past 120 years. I was particularly
interested in reconstructing the historical dynamics of D. melanogaster and galbut virus, which is an exceptionally
successful persistent virus in current fly populations. Galbut virus is a three-segmented partitivirus that is ubiquitous
in wild D. melanogaster populations: its average global prevalence is higher even than that of Wolbachia. This high
degree of success is attributable to galbut virus being biparentally vertically transmitted with near 100% efficiency, which
should drive the virus to 100% prevalence in wild populations. But multiple lines of evidence indicate that some flies are
resistant to infection and this virus and host may be co-existing in an uneasy equilibrium. To understand how long galbut
virus has been infecting D. melanogaster and how its sequence might have changed over the past century, I obtained
specimens ranging from 15-120 years from entomological collections across the United States. I obtained fragmented
but detectable RNA from these old specimens and was able to amplify galbut virus sequences using short-range RT-qPCR
from century-old flies. I am extending and confirming this finding using shotgun RNA sequencing to fully reconstruct the
virome of these historic specimens. This indicates that this currently successful virus has infected D. melanogaster since
the dawn of Drosophila research, and supports the conclusion that ancient dried insect specimens are a source of usable
RNA for paleogenomic studies.

1028A PECAn, a pipeline for image processing and statistical analysis of complex mosaic 3D tissues Michael
Baumgartner, Remi Logeay, Paul Langton, Alex Mastrogiannopoulos, Eugenia Piddini University of Bristol

Investigating organ biology requires sophisticated methodologies to induce genetically distinct clones within a tissue.
Microscopic analysis of such samples produces information-rich 3D images. However, the 3D nature and spatial
anisotropy of patches of cells makes sample analysis challenging and slow and limits the amount of information that
can be extracted manually. Here we have developed a pipeline for image processing and statistical data analysis which
automatically extracts sophisticated parameters from complex multi-genotype 3D images. The pipeline includes data
handling, machine-learning-enabled segmentation, multivariant statistical analysis, and graph generation. This enables
researchers to run rigorous analyses on images and videos at scale and in a fraction of the time, without requiring
programming skills. We demonstrate the power of this pipeline by applying it to the study of Minute cell competition.
We find an unappreciated sexual dimorphism in Minute competition and identify, by statistical regression analysis, tissue
parameters that model and predict competitive death.

1029B MARRVEL and ModelMatcher: Online resources to facilitate cross-disciplinary collaborations between
scientists, clinicians and beyond Shinya Yamamoto1,2, J Michael Harnish1,2, Lucian Li1,2,3, Seon-Young Kim1,2, Sanja Rogic1,2,5,
Guillaume Poirier-Morency1,4, Kym M Boycott4, Philip Hieter3, Paul Pavlidis1,2,5, Michael F Wanger1,2, Hugo J Bellen1,2,
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Canada

Progress in clinical genomic technologies such as whole-exome and whole-genome sequencing is leading to identification
of novel genetic causes of human diseases. In order to understand the functional consequences of disease-associated
genetic variants and to reveal the underlying pathogenic mechanisms, it is important to facilitate interdisciplinary
collaborations between clinicians and basic scientists who share interests in the same/orthologous genes. While
a number of national and regional efforts, such as the Undiagnosed Diseases Network (UDN) in the USA and Rare
Diseases Models and Mechanisms Network (RDMm) in Canada, have been established to facilitate such collaborations,
efforts to stimulate international collaborations have been lagging. To fill this gap, we have been developing MARRVEL
(https://marrvel.org) and ModelMatcher (https://www.modelmatcher.net), two informatic tools that allow scientists to
effectively gather information about a gene of interest in diverse species including human and to identify clinical
collaborators on a global scale, respectively. Using these tools, Drosophila researchers can quickly gather information
about the human and other model organism orthologs of their genes of interest and identify potential clinical and
scientific collaborators around the world. While there is a lot of value in pursuing fundamental scientific questions
that do not have obvious or immediate clinical value, an active involvement in a clinical project may help increase the
significance of a specific research project, and increase the impact of the basic scientific work.

1030C A toolkit to generate interconvertible overexpression Drosophila transgenes Luis Alberto Baena Lopez1, Franz
Wendler1, Sangbin Park2, Claire Hill1, Alessia Galasso1, Kathleen Chang2, Iman Awan1, Yulia Sudarikova1, Mar Bustamante-
Sequeiros1, Sichen Liu1, Ethan Seung1, Gabrielle Aisabonoko1, Seung Kim2 1) University of Oxford; 2) Department of
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Three independent binary systems for conditional gene expression (Gal4/UAS; LexA/LexAop; QF/QUAST) have greatly

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aided the genetic investigations using *Drosophila* as a model organism. However, each of these inducible gene expression systems requires the independent generation of non-interchangeable fly repositories. To alleviate this important logistic disadvantage and the need of triplicating transgene repositories for combinatorial experiments, we developed a modular vector that contains the regulatory elements for all three binary systems. This enables the separate or simultaneous overexpression of new transgenes through any of the available transcriptional factors (Gal4-, LexA- or QF), thus offering new experimental possibilities for *in vivo* cDNA overexpression (robust RNAi downregulation needs to be individually tested). Additionally, each transcriptional regulatory element (UAS, LexAop, QUAS) was flanked by two different but compatible recombination sites that facilitate their elimination from the genome with spatial and temporal control upon exposure to adequate recombinases. This allows the efficient generation of genetic mosaics within somatic cells as well as the elimination in the germline of specific gene-activating sequences. These unique features of our genetic toolkit provide unprecedented experimental possibilities and logistical advantages such as the generation of new fly lines with a subset of gene regulatory elements of interest upon excising the unwanted in a common precursor strain containing all of them.

1031A  Comprehensive Resource for the Drosophila 4th Chromosome *Michael Stinchfield*, Mary Jane O’Connor, Brandon Weasner, Bonnie Weasner, Justin Kumar, Kevin Cook, Michael O’Connor, Stuart Newfeld 1) School of Life Sciences, Arizona State University, Tempe, AZ; 2) Dept. Genetics, Cell Biology and Development, Univ. Minnesota, Minneapolis, MN; 3) Dept. Biology, Indiana University, Bloomington, IN

The 4th chromosome is the final frontier for genetic analysis in Drosophila. Small and devoid of recombination the 4th has long been ignored. Nevertheless it contains 105 genes (79 protein coding and 26 noncoding RNA). 74% of the protein coding genes have human homologs and 68% of these have a disease association. For example Eyeless belongs to the PAX/RAX family where inherited loss of PAX6 leads to Aniridia and somatic loss of RAX2 leads to age-related macular degeneration. A complete understanding of metazoan biology and thus human health requires the examination of these genes. To advance this effort ASU, IU and UMN will collaborate to create a resource facilitating the analyses of genes on the 4th. The resource will contain stocks divided into several collections. 1. FRT101F with a CRISPR mutation for loss of function and MARCM studies. 2. Conversion of protein coding genes with an existing MIMIC or CRIMIC to T2A.GAL4 and eGFP for protein traps and gain of function studies. 3. Gain of function stocks with a UAS.fly cDNA for each protein coding gene. 4. Gain of function stocks with a UAS.human cDNA for the two closest human homologs for each conserved protein coding gene. 5. Auxiliary chromosomes and balancers. Progress on all collections will be described. This resource will be made available through BDSC to all qualified investigators.

1032B  Multiscale, multi-perspective imaging assisted robotic microinjection of *Drosophila melanogaster* embryos *Andrew D. Alegria*, Amey S. Joshi, Kunpeng Liu, Jorge Blanco Mendana, Benjamin Auch, Daryl M. Gohl, Suhasa B. Kodandaramaiah 1,4,5 1) University of Minnesota Twin-Cities, Department of Mechanical Engineering, Minneapolis, MN; 2) University of Minnesota Twin-Cities, University of Minnesota Genomics Center, Minneapolis, MN; 3) University of Minnesota Twin-Cities, Department of Genetics, Cell Biology, and Development, Minneapolis, MN; 4) University of Minnesota Twin-Cities, Department of Biomedical Engineering, Minneapolis, MN; 5) University of Minnesota Twin-Cities, Department of Neuroscience, Minneapolis, MN

Microinjection is an important technique that has enabled transgenesis and targeted mutagenesis in *Drosophila*, as well as the wide array of methods and genetic toolkits which depend on transgenesis. However, microinjection remains a labor-intensive and highly specialized manual procedure, which makes it a critical bottleneck in the field and thus ripe for automation. Here, we present a computer-guided robot that automates the targeted microinjection of *Drosophila melanogaster* embryos. The robot uses a series of cameras and microscopes to image a petri-dish containing embryos at multiple magnifications and perspectives. This imaging is combined with a machine learning algorithm and computer vision algorithms to pinpoint a location on the embryo for targeted microinjection with microscopic precision. We demonstrate the ability of the robot to successfully microinject *Drosophila melanogaster* embryos. Results obtained indicate that the robot can significantly increase throughput as compared to manual microinjection since approximately in 1 hour the robot can inject upwards of 200 embryos. Since microinjections are performed in embryos on the petri-dish, the robot further eliminates the preparatory steps such as the dechorination and placing of the embryos on a glass slide prior to microinjection. Further, the robot is able to maintain survivability rates comparable to that of manual microinjection with survival rates upwards of 50%. We have demonstrated robust and reproducible transposon and PhiC31-mediated transgenesis with automated robotic microinjection of embryos directly on petri-dishes. This microinjection robot provides a means to achieve scalability as additional systems can be built at relatively low cost ($13,000) allowing parallelization of the injection process. In the future, we anticipate the robot can be used to perform high throughput microinjections and potentially enable new types of injection-intensive experiments.

1033C  Making Hox Gene-specific Drivers Using a Modified Trojan-exon Strategy *Fengqiu Diao*, Benjamin White  National Institute of Mental Health, NIH, Bethesda, MD
Hox family transcription factors are selectively expressed in distinct segments along the neuraxis of the *Drosophila* central nervous system. This regional specificity of expression, together with the functional importance of Hox gene-expressing neurons in motor behaviors, makes Hox genes attractive candidates for gene- and region-specific targeting methods. The “Trojan-exon” strategy has been broadly successful in producing gene-specific drivers, but producing such drivers for developmentally important transcription factors, such as Hox genes, has been problematic (Diao et al., 2015). This may be because these transcription factors are particularly sensitive to truncation of the protein translated from the allele containing the Trojan exon. Truncation necessarily lowers the level of transcription factor expression, but may also result in dominant-negative translation products that suppress transcriptional activity. To address these problems, we have developed a novel type of Trojan exon that incorporates split inteins. The protein fragments derived from a gene whose translation is normally interrupted by the Trojan exon will thus be re-ligated into a full-length active gene product, while at the same time allowing for expression of Gal4 or Split Gal4 components. Using this method, we have successfully made Gal4 drivers for the *labial* (*lab*), *Proboscipedia* (*Pb*), *deformed* (*dfd*), *Antenapedia* (*Antp*), *Ultrabithorax* (*Ubx*) and *Abdominal-A* (*Abd-A*) genes. In addition, we have made Split Gal4 hemidrivers for the *lab*, *dfd*, *Abd-A* and *Sex Combs-reduced* (*Scr*) genes. Gal4-driven UAS-reporter expression in the CNS is consistent with the known expression patterns of these genes. In addition, functional manipulation of previously characterized cell types yields the anticipated results. *Dfd* and other Hox genes play well-established roles in motor function, and we find that our *Dfd* - and *Scr*-specific hemidrivers express in glatamatergic neurons, including putative motor neurons. Consistent with this, suppression of neurons targeted by *Dfd*-Gal4 at the larval stage impairs feeding by preventing the mouth-hooks from being elevated. Conversely, stimulation of these neurons leaves the mouth-hooks constitutively elevated. In adults, stimulation of neurons in the *Dfd*-Gal4 pattern also alters feeding behavior by eliciting vigorous proboscis extension. Our results suggest that the new Hox-gene specific drivers express faithfully in their respective cell types and will be useful in developmental, physiological, and neural circuit-mapping studies.

**1034V Towards a novel method for cryopreservation via embryonic nuclear transplantation in *Drosophila* Troy Louwagie1,2, Grace Kringle1, Jorge Blanco Mendana1, Lindsey Gengelbach1, Benjamin Auch1, Daryl Gohl1,4, Allison Hubel1,2,5 1) University of Minnesota- Twin Cities, Department of Mechanical Engineering, Minneapolis, MN; 2) University of Minnesota-Twin Cities, ATP-Bio, Minneapolis, MN; 3) University of Minnesota-Twin Cities, University of Minnesota Genomics Center, Minneapolis, MN; 4) University of Minnesota-Twin Cities, Department of Genetics, Cell Biology, and Development, Minneapolis, MN; 5) University of Minnesota-Twin Cities, Department of Biomedical Engineering, Minneapolis, MN**

Cryopreservation of *Drosophila melanogaster* has been an area of interest since the 1990s when methods of embryo vitrification protocols were introduced by Stepnokus et al. (1990) and Mazur et al. (1992). However, there has still not been widespread adoption of these methods due to the difficulty of the protocols. Recent publications from Zhan et al. (2021) and Asaoka et al. (2021) have shown successful development of an alternative vitrification methods for cryopreservation of embryos and a novel cryopreservation method based on transplantation of cryopreserved primordial germ cells, respectively. We are testing an alternative method that involves the slow cooling of embryonic nuclei and regeneration of stocks by embryonic nuclear transplantation (ENT). Using confocal Raman spectroscopy is beneficial for label-free imaging of nuclei and the distribution of water and CPA. Osmolyte solutions consisting of sucrose, glycerol, and other buffering agents that aid in the isolation of Drosophila nuclei act as quality CPAs in the slow-cooling cryopreservation process. Studies at 4°C show that certain CPAs, such as glycerol, can permeate the nuclear membrane while other CPAs, like sucrose, cannot penetrate the nuclear membrane. Using a controlled rate temperature-cooling stage, we can image the nucleus at -50°C with varying cooling rates and nucleation temperatures. We are also using an RT-qPCR assay for *hsp70* induction to monitor nucleus health and viability throughout the isolation, cryopreservation, and recovery process. This functional assay will aid in determining the optimal parameters of *Drosophila* cryopreservation with the use of a differential evolution (DE) algorithm. The DE algorithm uses experimental observations (CPA composition, cooling rates, nucleation temperatures, and post thaw viability) to modify existing population vectors and predicts solutions that may be more favorable. This DE algorithm could also be applied to other Drosophila cryopreservation methods to optimize CPA and freezing parameters. Finally, we have been attempting to create clones or chimeric embryos derived from isolated nuclei using ENT. ENT has previously been demonstrated in *Drosophila* and avoids issues associated with the impermeability of *Drosophila* embryos to cryoprotectants. We report progress in developing this novel method which has the potential to provide additional options for the practical cryopreservation of Drosophila stocks.

**1035V Development of a fly model to probe the functions of inorganic polyphosphates Sunayana Sarkar1, Harsha Sharma1, Jayashree Suresh Ladke2, Rashna Bhandari3, Manish Jaiswal1 1) TATA Institute of Fundamental Research, Hyderabad, India; 2) Centre for DNA Fingerprinting and Diagnostics, Hyderabad, India**

Inorganic polyphosphate (polyP), which consists of chains of orthophosphate residues of varying chain lengths, is found in all living organisms alongside other phosphate-rich biomolecules such as ATP and inositol phosphates. The biological functions of polyP have been extensively studied in prokaryotes and unicellular eukaryotes, however, their functions in
metazoans are largely underexplored. In fact, polyP has been referred to as a molecular fossil. In recent years polyP has been shown to be implicated largely in blood clotting and bone mineralisation in mammals. Moreover, several other functions of polyP are predicted due to its presence in various cell types and its specific binding with numerous proteins. However, the major limitation in testing the functions of polyP in metazoans is the elusivity of the genes involved in polyP synthesis and turnover, which restricts modulation of polyP levels in vivo. We have developed a Drosophila model to study the functions of polyP. Here we show that polyP exists in flies and that its levels are developmentally regulated during embryogenesis. Further, through phylogenetic analysis, we characterised Prune as a putative exopolyphosphatase. Indeed, we observed significantly higher levels of polyP in prune mutants as compared to wildtype flies. Prune is a mitochondrial protein known to hydrolyze cAMP. Mutations in prune are linked to a variety of processes such as mitochondrial biogenesis, eye pigmentation and neurodegeneration. Based on our data, we surmise that prune mediated regulation of polyP may attribute to some of the prune phenotypes and will help understand polyP biology in metazoans.

1036V A split-Gal4 system that is repressible by Gal80 Ben Ewen-Campen1, Haojiang Luan1, Jun Xu1, Tanuj Thakkar1, Benjamin White1, Norbert Perrimon1,2 1) Harvard Medical School, Boston, MA; 2) HHMI; 3) National Institute of Mental Health, NIH, Bethesda, MD

Most of the Gal4 drivers that are commonly used in Drosophila research are not solely expressed in the cells- or tissues-of-interest, but also in additional, often unwanted cells or tissues. To overcome this relative lack of specificity, the split-Gal4 intersectional technique was developed, which has successfully been used to create extraordinarily specific driver lines, including many expressed in only a single cell-type. However, unlike native Gal4, existing split-Gal4 tools cannot be repressed by the Gal80 repressor or any other commonly-used repressor, and therefore cannot be temporally controlled. Here, we present a split-Gal4 system which is fully repressible by Gal80 in vivo. Based on a split-intein, this system drives transgene expression at levels indistinguishable from the original split-Gal4 system, and combines the cell-type specificity of split-Gal4 with the temporal control of native Gal4/Gal80. In addition to creating highly specific genetic driver lines, we show that this system can also be used to systematically map single cell RNAseq clusters to anatomy.

1037V Efficient allelic conversion by homologous chromosome-templated repair in Drosophila somatic tissues Annabel Guichard, Sitara Roy, Sara Juste, Marketta Sneider, Ankush Araudkar, Zhiqian Li, Ehan Bier Univ California, San Diego, CA

Recent advances in Cas9-based genome editing offer broad applications in research, agriculture and medicine. These techniques typically entail insertion of exogenous gene sequences provided by a donor template into specific genomic sites following Cas9-mediated DNA cleavage. Here, we demonstrate efficient somatic allelic conversion wherein cleavage-resistant sequences present on one chromosome correct a targeted cleavage-sensitive allele at the corresponding site on the homologous chromosome. Using a set of allelic combinations of the white locus of Drosophila, we show successful allelic conversion as revealed by quantifiable red eye clones produced by Cas9-dependent cleavage, or by its nickase variants Cas9D10A and Cas9H840A. Repair phenotypes elicited by Cas9 versus nickases differ substantially in several respects, including efficiency, production of cleavage-resistant alleles through the NHEJ mutagenic pathway and developmental timing. We show that allelic correction does not strictly require long-range chromosomal pairing, and that sequence homology alone can promote such repair from a non-homologous location in the genome. If generalized to human cells, this simple approach could be adapted to correct a wide array of dominant or trans-heterozygous genetic conditions offering novel gene therapy strategies.

1038V A method to estimate the frequency of chromosomal rearrangements induced by CRISPR/Cas9 multiplexing in Drosophila Bruce Reed, William Ng, Richard Do, Brittney Lato, Emily Baker, Mackenzie Vallee University of Waterloo, Waterloo ON, CANADA

Using CRISPR/Cas9 to simultaneously induce mutations in two or more target genes, commonly referred to as multiplexing, may result in chromosomal rearrangements such as inversions or translocations. While this may be undesirable in some contexts, the ability to recover chromosomal rearrangements targeted to specific sites in the genome is potentially a powerful tool. The frequency of chromosome rearrangements induced by CRISPR/Cas9 multiplexing, however, remains unknown in Drosophila. To estimate the frequency of chromosome rearrangements induced by multiplexing, we developed a self-selecting screening system using Drosophila stocks that carry an autosomal pericentric inversion in what is known as the autosynaptic form. All progeny of normal females crossed to males of these autosynaptic stocks are lethal due to excessive aneuploidy. If an inversion is induced within the female germline, however, and if this inversion is analogous to the inversion in the male autosynaptic line, then it is possible to recover progeny in which aneuploidy is reduced and viability is restored. Using this self-selection screening method, we were able to estimate the frequency of viable progeny from females having germline expression of active Cas9 together with ubiquitously expressed sgRNAs targeted to desired inversion breakpoints. Salivary gland polytene chromosome analysis, PCR, and sequencing confirmed the recovery of chromosome breakpoints induced between the two sgRNA target sites. Overall, we demonstrate that CRISPR/Cas9 multiplexing can induce chromosomal rearrangements in Drosophila.
In general, the generation of a pericentric inversion within a female germline using CRISPR/Cas9 multiplexing varied depending on the choice of sgRNAs and occurred less than once for every 100 germlines tested. This frequency was also found to be comparable to using FLP/FRT site specific recombination. In using this particular system, we conclude that the induction of chromosomal rearrangements associated with CRISPR/Cas9 multiplexing is not a high frequency event.

**1039V Identification of Drosophila new genes using machine learning** Gabriel Goldstein1,2, Maria Vibranovski1,3, Yong Zhang1 1) Universidade de São Paulo, USP, São Paulo, Brazil; 2) Chinese Academy of Sciences, CAS, Beijing, China; 3) Arizona State University, Tempe, USA

New genes are defined by their presence in a taxon and absence in sibling taxa. These genes have great biological importance and are involved in processes of high selective pressure, being expressed in tissues such as brain and testis. There are a variety of genetic mechanisms that can lead to the generation of new genes, such as duplications and retrotranspositions for example, but most new genes are derived from duplications. The exact functions of these genes in organisms are still being studied, but some work has already shown a relationship with the resolution of sexual conflicts, for example. Despite this, there are a number of biological characteristics that are known to be different between new and old genes. An example of this is the expression profile of these groups, since new genes are mostly expressed in male gametogenesis and old genes are broadly expressed. The main gene dating method for identifying new genes uses synten, which is the phenomenon of conservation of the order and gene content of a region in the genome that occurs in related species, and parsimony when comparing genomes of related species to date all genes of a focal species. Despite the accuracy of the method, it is extremely dependent on the assembly and annotation of the genome of interest, which limits its application to model species that have a manual and curated annotation. With these facts in mind, we propose in this work a method of identifying new genes that uses biological information to separate new and old genes through the use of machine learning. Machine learning algorithms are those able to change with experience and are able to identify complex patterns and identify classes from a variety of information. With this, we trained a model with the random forest algorithm in the model species Drosophila melanogaster and obtained 0.508 precision and 0.718 recall with generated data. In addition, we identified the 1523 new genes of D. pseudoobscura using the existing method so that we can use this species as a second control point for our method.

**1040V REDfly: The Regulatory Element Database for Drosophila and other insects** Soile V. E. Keränén1, Andrew E. Bruno2, Marc S. Halfon1,4, 1) None; 2) Center for Computational Research, University at Buffalo-State University of New York, Buffalo, NY; 3) Departments of Biochemistry, Biomedical Informatics, and Biological Sciences, University at Buffalo-State University of New York, Buffalo, NY; 4) NY State Center of Excellence in Bioinformatics & Life Sciences, Buffalo, NY; 5) Department of Molecular and Cellular Biology and Program in Cancer Genetics, Roswell Park Comprehensive Cancer Center, Buffalo, NY

REDfly is a comprehensive knowledgebase of experimentally-validated cis-regulatory modules (CRMs) and transcription factor binding sites (TFBSs) for Drosophila and other insects, including the malaria mosquitoes Anopheles gambiae and Aedes aegypti and the beetle Tribolium castaneum. The database contains data on more than 53,500 regulatory sequences and almost 2700 TFBSs or their variants. Our goal is to include all functionally tested or predicted sequences, both with and without observable regulatory activity, to maximize the utility of the knowledgebase for purposes ranging from detailing the regulatory structure of a single locus, to large-scale studies of the regulatory genome, to providing training and/or validation data for machine-learning analyses of gene regulation. CRM data in REDfly are broadly segregated into three categories depending on the experimental evidence underlying their identification: reporter gene analysis, deletion of sequences from the genome, or predictions based on genomic assays such as ChIP-seq, ATAC-seq, and computational modeling. Finer-tuned filtering based on evidence type is possible through the “advanced search” interface. An additional key REDfly feature is its extensive and searchable expression pattern annotation for each CRM’s activity using structured anatomy, staging, and biological process ontologies. Recently, REDfly has started to curate silencers, which inhibit gene transcription, in addition to the existing curation of both tested and predicted enhancers. Also new is a “species” page that provides summaries of the data included for each species curated by REDfly, with BLAT and in silico PCR interfaces for each annotated genome. All REDfly code has been made open source. Future plans include updating REDfly’s search and download features to better serve our users, and providing a customized JBrowse interface for improved visualization and integration of REDfly data. REDfly is freely accessible at http://redfly.ccr.buffalo.edu and can be followed on Twitter at @REDfly_database.

**1041V Proteomic mapping of organ secretomes using in vivo proximity labeling** Justin A. Bosch1, Pierre M. J. Beltran2, Cooper Cavers3, Thai LaGraff2, Randy Melanson2, Steven Carr3, Norbert Perrimon1,3 1) Department of Genetics, Blavatnik Institute, Harvard Medical School, Boston, MA; 2) Broad Institute, Cambridge, MA; 3) Howard Hughes Medical Institute

Secreted proteins regulate essential biological processes and are attractive for developing therapeutics. Due to their low abundance in extracellular fluids, discovering the complete set of secreted proteins (the “secretome”) - as well as their tissue of origin – can be extremely challenging. To address this, we developed a proteomic method to label proteins in specific organs in Drosophila larvae using the biotin ligase TurboID, and subsequently enrich and identify biotinylated
proteins in the hemolymph. Using this tool, we constructed a secretome map of 540 proteins from 10 major cell types (e.g. fat body, muscle). In addition to identifying most known blood proteins (e.g. Lsp2, apolpp), we discovered hundreds of uncharacterized proteins, in many cases derived from a single cell type. Furthermore, while most blood proteins originate from the fat body, we identify secreted proteins from less appreciated cell-types, such as oenocytes and glia. Using CRISPR/Cas9 knock-in flies to validate selected tissue-specific uncharacterized proteins, we found that most were expressed in the presumed tissue and some were detected in circulation by western blot. One protein, CG6867, the single fly homologue of human Olfactomedin proteins, is specifically expressed in somatic muscle and secreted into larval hemolymph. Interestingly, CG6867 accumulates on the basal surface of most tissues (e.g. imaginal discs), suggesting CG6867 may be a novel circulating basement membrane protein. Our secretome map will serve as a resource to investigate blood protein function, discover organ communication factors, and compare with homologues of human biomarkers.

1042V Importance of cell-cycle and cell-sex correction in single-cell analysis: unmasking novel target genes of the Hedgehog pathway Nicholas Everettts, Melanie Worley, Riku Yasutomi, Nir Yosef, Iswar Hariharan University of California, Berkeley

Single-cell transcriptomics is an unparalleled technique for obtaining a holistic understanding of biological systems, including tissue composition, organ development, and cellular differentiation. Many computational packages offer default end-to-end analysis for single-cell RNA sequencing (scRNAseq), performing common tasks such as dimensionality reduction, clustering, and differential expression testing. Furthermore, when processing multiple samples, there are a plethora of tools available for integrating datasets. While much attention has been given to the importance of batch correction, fewer studies examine biological covariates within datasets that can mask important signals and decrease statistical power. We examined the how strong biological signals can obscure interesting biology by examining the adult muscle precursor cells (AMPs) of the Drosophila wing-imaginal disc, a population of cells that develop into all adult flight muscles but exhibit high transcriptional homogeneity during larval development. We show that by using scRNAseq analysis tools with default settings, cell clusters are primarily split by two biological features: cell-cycle status and the biological sex of the donor fly. This stratification not only hinders the classification of AMPs into their canonical cell types, the “direct” and “indirect” AMP subpopulations, but also obscures the discovery of new biological signals and interactions by separating otherwise transcriptionally-similar cells. Cell-cycle and cell-sex signatures are observable within the latent space generated by dimensionality reduction, enabling post hoc correction by identifying the latent dimensions that correlate with markers of proliferation and biological sex. By removing these confounded latent dimensions from downstream processing, cell-type classification of AMPs is improved and the Hedgehog pathway is unmasked as an important signaling pathway for the proper development of adult flight muscles. With our analysis, we identified and validated two novel Hedgehog-signaling targets, Neurotactin and midline, within the AMPs. Additionally, confounding covariates can be supplied to the deep-learning single-cell software scvi-tools, taking advantage of its powerful integration models while suppressing unwanted biological signals. Our research highlights the importance of identifying and correcting for unnecessary biological signals in addition to technical effects, and provides easily-implemented solutions for removing signals such as cell cycle and cell sex.

1043V Characterization of shock wave effects in syncytial embryos of Drosophila melanogaster using fluorescent nanoparticles Daniel Tapia Merino1,2,3, Achim Max Loske Mehling1,3, Gabriel Ramos Ortiz4, Juan Rafael Riesgo Escovar2,5 1) Maestria en Ciencias (Neurobiologia), INB, Queretaro, Mexico; 2) Instituto de Neurobiologia (INB), Queretaro, Mexico; 3) Centro de Fisica Aplicada y Tecnologia Avanzada (CFATA), Queretaro, Mexico; 4) Centro de Investigaciones en Optica, A.C., Guanajuato, Mexico; 5) Universidad Nacional Autonoma de Mexico Campus Juriquilla, Queretaro, Mexico

We employed fluorescent nanoparticles as fiducial markers to evidence the insertion of material into the early syncytial embryo of Drosophila melanogaster, using underwater shock waves. These mechanical waves generate microjets that, in principle, perforate the outer layers of the embryos (chorion and vitelline membranes) as well as the embryonic plasma membrane. A convenient shock wave pressure profile was found, to guarantee the survival and growth of a reasonable percentage of the treated embryos to become viable, fertile adults. Our results reveal that fluorescent nanoparticles of different sizes and composition, added to the incubation medium, upon shock wave exposure, are able to penetrate the embryonic layers, and stay inside the fly tissues. These results could lead to diverse biological applications in a few simple steps.

1044V OligoY: a pipeline for the design of repetitive oligopaint probes for the Y Chromosome Isabela Almeida1, Henry Bonilla Bruno1, Mara Pinheiro1, Amanda Luvisotto1, Antonio Carvalho2, Maria Vibranovski3 1) Universidade de Sao Paulo, USP, Sao Paulo/SP, Brazil; 2) Universidade Federal do Rio de Janeiro, UFRJ, Rio de Janeiro/RJ, Brazil

Studies with the Y chromosome offer a series of obstacles, one of them being the heterochromatic and highly repetitive state of this structure, leading to difficulties in assembling scaffolds and contigs and, thus, in a lack of final assembled sequences for it. The Y chromosome of Drosophila melanogaster, the model organism used in this research, has an estimated size of 41Mb of repeat-rich sequences, but only 10% of them are assembled in the most recent genome
release. In contrast, the protocol for designing probes used in full chromosome fluorescent labeling experiments (FISH Oligopaint) does not include repetitive sequences to avoid off-target hybridization. For this reason, there are currently less than 1500 oligopaint probes for this Y chromosome model, which is a value at least ten times smaller when compared to the one observed for other chromosomes of the same species. Furthermore, this amount is insufficient to carry out FISH Oligopaint assays efficiently. The main objective of this research is to develop a pipeline that allows the design of oligopaint probes for the Y chromosome off any species of interest. The final pipeline includes the use of open-source and existing tools in Bioinformatics, identification of sequences unique to the chromosome of interest, guarantees the user the autonomy to choose parameters and effectively uses repetitive sequences unique to the target chromosome to design probes, therefore maximizing overall efficiency of cytogenetic experiments. After extensive tests and validations in silico and in situ, it was verified that the application of the developed pipeline, OligoY, allows staining the Y chromosome without generating off-target signal, despite the use of repetitive sequences for oligopaint probe design. In addition, some of the OligoY pipeline strategies’ can be used in other chromosomes to address gaps in repeat-rich regions that otherwise would not be stained.

1045V A Bibliometric Analysis of Somatic Mutation and Recombination Tests of Drosophila melanogaster Ghada Tagorti, Bülent Kaya Akdeniz University, Antalya, Turkey.

Somatic mutation and recombination tests (SMARTs) are accurate and low-cost assays performed to assess the genotoxic potential of multiple pollutants by using Drosophila melanogaster as a model organism. The present study attempts to understand the current research situation and knowledge base in SMART assays research. To accomplish this, data were extracted from the Web of Science Core Collection from 1984 to 2020, to be analyzed by HistCite, Biblioshiny (RStudio), VOSViewer, and CiteSpace. Results have shown an increase in publications within the last ten years, with an annual rate of 14.4 articles. A total of 392 records involving 35 countries were published in 96 sources, mainly from Brazil (n = 106; 27.04%), Spain (n = 69; 17.60%), and Turkey (n = 66; 16.84%). The Federal University of Uberlândia ranked first in terms of publications (n = 39; 9.95%). The Mutation Research-Genetic Toxicology and Environmental Mutagenesis journal had the highest total global citation score per year. Based on document co-citation, five large research clusters were identified. The ‘nanoparticles’ cluster was the youngest research frontier. The top-cited documents focused on SMARTs protocols, statistical tests, and proposed strains to improve the scoring. With this study, the evolution of SMARTs research trends over years was demonstrated. However, international collaboration should be more encouraged and supported.

1046A The Genomics Education Partnership: Teaching and Research Opportunities Raffaella Diotti1, Cindy Arrigo2, Martin G. Burg3, Justin R. DiAngelo4, Sarah C. R. Elgin5, Christopher E. Ellison6, Christopher J. Jones7, Jennifer Kennell8, Judith Leatherman9, Wilson Leung9, David Lopatto10, Vida Mingo11, Laura K. Reed12, Chinmay P. Rele12, Katie M. Sandlin12, Jamie Siders13, Norma Velazquez-Ulloa Velazquez-Ulloa14, The Genomics Education Partnership 1) Bronx Community College, Bronx, NY; 2) New Jersey City University, Jersey City, NJ; 3) Grand Valley State University, Allendale, MI; 4) Penn State Berks, Reading, PA; 5) Washington University in St. Louis, St. Louis, MO; 6) Rutgers University, Piscataway, NJ; 7) Moravian University, Bethlehem, PA; 8) Vassar College, Poughkeepsie, NY; 9) University of Northern Colorado, Greeley, CO; 10) Grinnell College, Grinnell, IA; 11) Columbia College, Columbia, SC; 12) University of Alabama, Tuscaloosa, AL; 13) Ohio Northern University, Ada, OH; 14) Lewis & Clark College, Portland, OR

The Genomics Education Partnership (GEP; thegep.org) is a nationwide collaboration of faculty from 180+ institutions with the goal of increasing understanding of genetics and familiarity with bioinformatics in diverse student populations through the incorporation of Course-based Undergraduate Research Experiences (CURES) in the curriculum. Participating institutions range from community colleges, to PUIs, MSIs, HBCUs, and R1 universities. To facilitate both faculty and student access to curriculum materials and support active participation in the research, GEP provides a web-based platform with curated curriculum/training materials that can easily be incorporated into existing courses. During the COVID-19 pandemic, the accessible and immersive GEP curriculum and custom bioinformatics tools provide an inexpensive online framework for students to participate in research despite the lack of access to a traditional lab environment. During the 2020-2021 academic year, GEP reached over 3900 students. Through the GEP curriculum, students learn to annotate newly-sequenced eukaryotic genomes. They learn to leverage evidence from related informant species, experimental data (e.g., RNA-Seq), gene prediction algorithms, evolutionary conservation, and basic molecular biology rules to create a defensible gene model. The GEP research projects include investigation of venom evolution in parasitoid wasps, the evolution of insulin pathway genes across 27 Drosophila genomes, and expansion of the F element in four Drosophila species. We plan to expand the projects to additional species. Student gene models are reconciled and collated to generate a large dataset for evolutionary genomic studies, with student/faculty co-authors. In addition, GEP supports the publication of the gene models as microPublications, with students as lead authors. The GEP also engages in science education research. Our recent findings suggest that student perceptions of science, positive and negative, impact student learning outcomes. In addition to accessing the curriculum materials and tools, GEP faculty benefit from professional development opportunities and the support of a national network of like-minded colleagues. With the support of NSF and NIH, GEP is actively recruiting additional faculty members, particularly at MSIs and
community colleges, and both science and science education partners to collaborate on additional projects. Supported by NSF grants 1915544 and 1431407, and NIH R25GM130517.

1047B DrosoPHILA: a partnership between teachers and scientists that begins in the lab and continues in city schools Kaitlin Laws1, Anthony Natale2,3, Edward Waddell1-4, Jamie Shuda2,3, Greg Bashaw1 1) University of Pennsylvania Perelman School of Medicine; 2) University of Pennsylvania Institute for Regenerative Medicine; 3) University of Pennsylvania Netter Center for Community Partnerships; 4) Holy Family University

While exposure to laboratory activities often awakens student interest in science, many schools lack the resources to offer such experiences. Outreach sponsored by local colleges and universities can provide students with first-hand knowledge of the scientific process, reinforce learning objectives, and connect them to scientists in their community. Our program, DrosoPHILA, is a two-part high school curriculum built by teachers in the School District of Philadelphia (SDP) with input and support from members of the Bashaw lab and experienced outreach educators. In the first phase of this collaboration, three teachers from SDP conducted paid research in the Bashaw lab while helping to develop outreach modules. These modules, Flies on Ice and Roundabout We Go!, highlight the utility of Drosophila as a model organism for neurobiology research while reinforcing students’ understanding of the scientific method and quantitative reasoning. In Flies on Ice, students determine how the time flies spend anesthetized on ice affects their recovery. In Roundabout We Go!, students study larval crawling in mutants with aberrant nervous system development. Each module is scaffolded by grade-appropriate presentations, student-driven experimentation, and guided data analysis. Our team leads at least two fifty-minute class periods of instruction.

In phase two of our program, we invited six SDP teachers to a two-day professional development program in the Bashaw lab, where they participated in our modules. During the 2018-2019 school year, our team led visits to these teachers’ classrooms to guide students through both modules, reaching 252 students. In Fall 2019, we expanded our reach, visiting 14 classrooms (349 students). At the onset of the COVID-19 pandemic, we transitioned our program online and reached nearly 500 students by remote instruction. In Fall 2021, we resumed classroom visits, reaching more than 600 students across 21 classrooms. By surveying students before and after our visits, we were able to identify whether and how DrosoPHILA affected their knowledge and science identity. Our preliminary analysis indicates that our modules improve students’ understanding of what it is like to be a scientist. Furthermore, because we surveyed students receiving both remote and in-person instruction, we are positioned to ascertain differences in outcomes under these circumstances. The teacher-informed design of our modules, along with the broad use of Drosophila as a model organism, makes them suitable for school districts across the United States.

1048C Investigating the impacts of engaging undergraduates as developers of inclusive curriculum through a service-learning course Blake Riggs1, Maurina Aranda2, Jeffrey Schinske3, Laura Burrus1, Kimberly Tanner1 1) San Francisco State University; 2) Southern Illinois University Edwardsville; 3) Foothill College

Scientist Spotlights – curricular materials that employ the personal and professional stories of scientists from diverse backgrounds – have previously been shown to positively influence undergraduate students’ relatability to and perceptions of scientists. We hypothesized that engaging students in authoring Scientist Spotlights might produce curricular materials of similar impact, as well as provide a mechanism for student involvement as partners in science education reform. To test this idea and investigate the impact of student-authored Scientist Spotlights, we developed a service-learning course in which teams of biology students partnered with an instructor to develop and implement Scientist Spotlights in a biology course. Results revealed that exposure to 3 or 4 student-authored Scientist Spotlights significantly shifted peers’ perceptions of scientists in all partner courses. Interestingly, student-authored Spotlights shifted peers’ relatability to scientists similarly among both white students and students of color. Further, student authors themselves showed increases in their relatability to scientists. Finally, a department-wide survey demonstrated significant differences in students’ perceptions of scientist representation between courses with and without student-authored Spotlights. Results suggest that engaging students as authors of inclusive curricular materials and partners in reform is a promising approach to promoting inclusion and addressing representation in science.

1049A Extending the Fly-CURE into an Upper-Level Undergraduate Bioinformatics Course Kayla Bieser1, Richard Tillett2, Jacob Kagey3, Karen Bieluch4, Nikolaos Tsotakos5 1) Nevada State College; 2) University of Nevada Las Vegas; 3) University of Detroit Mercy; 4) University of Maine; 5) Penn State Harrisburg

The Fly-CURE is a Course-based Undergraduate Research Experience (CURE) in which undergraduate genetics students from multiple US institutions characterize and conduct deficiency mapping of mutants in Drosophila melanogaster. A FLP/FRT EMS screen was conducted in the Drosophila eye in the context of blocked apoptosis to screen for conditional mutants that alter control of cell growth and development. The goal is to identify and map the mutation to a genomic region and/or gene on 2R. However, the gene and molecular location of some mutants have proven difficult to map utilizing these complementation techniques. To identify the mutations, we conducted whole genome sequencing (WGS) and bioinformatics analysis utilizing stocks of heterozygous EMS mutared flies with an unknown mutational locus, but
in some instances with a known genetic region mapped. A bioinformatics workflow was successfully established and verified utilizing four EMS mutants identified in the initial screen and a control stock representing their pre-EMS genetic background. When combined with deficiency mapping data, this methodology successfully identified mutated SNPs. To broaden participation and interest in bioinformatics, a scaffolded curriculum with the Fly-CURE was created at Nevada State College, a 4-year HSI. A 400-level bioinformatics course was implemented with a virtual format in Fall 2020 utilizing GitHub for course content and an app was developed in the CyVerse Discovery Environment for terminal and data access. A modification of Data Carpentries Genomics curriculum was utilized to introduce bash, the Unix command line and genomics methods, followed by a tailored hands-on pipeline for student analysis of WGS data collected from mutant flies the students had previously studied in the Fly-CURE. To-date, 49 students have completed the course and analyzed the genomes of 14 EMS mutant Drosophila, with eight mutations having been conclusively determined via this methodology. Preliminary assessment data supports that most students had little to no prior experience in bioinformatics and made significant gains in their understanding of file structures, Unix tools, Next Gen Sequencing, and their understanding of bioinformatics and how it relates to studying real world problems. The scaffolding of curriculum from genetics into bioinformatics allowed for greater understanding and buy-in by the students. Future goals include expansion of the curriculum to other institutions.

1050B Comparative Effectiveness of Antioxidant and Lowered Carbohydrate Diets on Dysplastic Guts Sandra Illescas, Megan Hampton, Marie Page, Jessica Pacheco California State University Northridge

Drosophila melanogaster is a well-established model system within the biomedical research community, however, there currently exists a large gap in the field: the lack of a standardized diet. Standardization is critical for understanding the effects of dietary changes on stem cells in Drosophila melanogaster. This study sought to determine the effects that reducing carbohydrates and/or adding antioxidants to the diet could have on intestinal stem cells (ISCs) in the Drosophila melanogaster midgut. Our study analyzed morphological changes in ISCs that occurred as a result of the use of one of three dietary groups designed to work against oxidative stress in contrast with the control. By using four experimental groups, this study was able to consider the effects of reducing sources of oxidative stress, providing protective measures against oxidative stress, and the potential synergistic or antagonistic effects between these two dietary manipulations. Experimental groups had a reduction of carbohydrates, the inclusion of antioxidants, or the combination of both manipulations. This multimodal approach allowed the observation of morphological changes in the ISCs that could potentially only be a result of the overlap between the two variable changes. We hypothesized that a diet lower in oxidative stress could ameliorate premature dysplasia in the Drosophila melanogaster midguts. Intestinal samples were analyzed using whole mount fluorescent microscopy and CellProfiler to quantify changes in the number and/or morphology of ISCs between experimental groups. Data was collected using CellProfiler to measure the cell count and area of GFP stained cells, and normalized to DAPI stained cells. A one-way analysis of variance (ANOVA) with a 95% confidence interval showed a significant difference among groups (p=0.007), that could be attributed to the combination of lower carbohydrates and added antioxidants, as determined by a post hoc Tukey test. These initial results would indicate that the addition of antioxidants in combination with the reduction of carbohydrates appears to promote a higher density of progenitor cells in ISCs of Drosophila melanogaster, contrary to our original prediction. Our work demonstrates how even slight dietary manipulations can have detectable effects on ISC biology and will hopefully serve as a basis for future studies exploring the link between diet and intestinal homeostasis.

1051C Students who participate in Fly-CURE demonstrate gains in self-efficacy and belonging across a Research Coordination Network both before and during the COVID-19 pandemic Jacob Kagey1, Kayla Bieser2, Joyce Stamm3, Alysia Vrailas-Mortimer4, Fly-CURE 1) University of Detroit Mercy, Detroit, Michigan; 2) Nevada State College, Las Vegas, Nevada; 3) University of Evansville, Evansville, Indiana; 4) Illinois State University, Bloomington, Illinois

Fly-CURE is a consortium of fifteen institutions implementing a Drosophila centered research project within undergraduate genetics laboratory courses. In this CURE, students work to characterize and map novel EMS mutants isolated from a FLP/FRT screen. To date, students have mapped 19 mutants on chromosome 2R including alleles of Egfr, Shot, Hpo, and Shn. These findings have resulted in six micropublications with 368 student co-authors. To understand the impact of Fly-CURE on student learning and attitudes towards research, we have assessed students on their perceived changes in sense of self-efficacy and sense of belonging, as well as gains on genetics-based learning objectives. These results were segregated by whether students had a previous research experience prior to the Fly-CURE. Overall, students who participated in the Fly-CURE report statistically significant gains in science self-efficacy and sense of belonging. Further, those students without a prior research experience reported lower scores in science self-efficacy before the class but have levels of science self-efficacy indistinguishable from those with previous research experiences by the end of the Fly-CURE. During the academic transition resulting from the COVID-19 pandemic, Fly-CURE labs were taught using virtual, hybrid, and in-person modalities. Despite these changes in delivery methods, we see no statistical differences in student-reported gains before or during the pandemic. The Fly-CURE is an adaptable Drosophila based CURE that impacts student attitudes towards research and contribute to the understanding of the genetic regulation of developmental growth.
1052A  A Research-based laboratory course in Molecular Biology, Genetics, and Evolution  Eric Spana Duke University

A introductory level course at Duke University in Molecular Biology, Genetics, and Evolution required a lab component that spanned as much of these three subjects as possible. I have developed a lab that uses local, wild-caught fruit fly species (mostly Drosophila) as the starting point for a laboratory portion in the course. Students choose flies from the collection and use PCR and sequencing of the Cytochrome Oxidase I gene to identify the species via DNA barcoding analysis. After phylogenetic analysis of the collected species, students amplify a region of a second gene, the Tyrosine Hydroxylase gene (TH) using degenerate PCR primers. This PCR product is then cloned into a TA cloning vector. After transformation, mini-preps, and restriction digest analysis, students can identify which samples contain the appropriate cloned DNA. Sequencing and analysis of the plasmid insertions follows and a number of evolutionary analyses are performed on the sequence. The region of TH that is being sequenced covers both introns and exons, and there is a cis-regulatory element in one intron that drives expression in the larval brain in D. melanogaster. Alignment of the TH region sequences identified a few highly conserved regions within the intron that contains the cis-regulatory element. I have run this course at two different Universities so far: Duke University and Duke Kunshan University, and have identified ~22 species in Durham, NC and ~11 in Kunshan, China with an overlap of only three species and at least one unidentified species in each location. This laboratory gives students experience in many common molecular biology techniques: PCR, cloning, plasmid DNA isolation, restriction digests, agarose gels, and sequencing as well as analysis of their own data for gene structure, reading frames, codon usage, dN/dS ratio, MK test, phylogeny. An online version of this lab used extensive BLAST and gene annotation of 80+ sequenced genomes to arrive at a similar result.

1053V  Reproducibility for Everyone  April Clyburne-Sherin1, Nele Haelterman2, Ruchika Bajaj3, Reproducibility for Everyone Group 1) Reproducibility for Everyone, Brooklyn, NY; 2) The Scientist, Wilmington, DE; 3) UCSF, San Francisco, CA

Rigor and reproducibility are at the core of modern science. Many new initiatives and tools have been established to address barriers to reproducibility. While very welcome, these projects have led to a proliferation of online tools and resources which can be hard to sift through.

Reproducibility for Everyone (R4E) is a global, community-led reproducibility education initiative. R4E runs practical and accessible workshops to introduce the concept of reproducibility to researchers. We demonstrate reproducible tools and methods that can improve research by making it more efficient, transparent, and rigorous. Every R4E workshop is customized for the audience. For Drosophila 2022, R4E instructors will focus on aspects of reproducibility most relevant to Drosophila research and share resources and tools curated for the GSA community.

This workshop will introduce reproducible workflows and a range of tools along the themes of organization, documentation, analysis, and dissemination. After a brief introduction to the topic of reproducibility, the workshop will provide specific tips and tools useful in improving daily research workflows. The content will include modules such as data management, electronic lab notebooks, reproducible bioinformatics tools and methods, protocol and reagent sharing, data visualization, and version control. All modules include interactive learning, real-time participation, and active knowledge sharing. The methods and tools introduced help researchers share work with their future self, their immediate colleagues, and the wider scientific community.

Following this workshop, participants will be able to:
1. Apply a conceptual framework for reproducibility, replicability, and robustness of research.
2. Explore practical, accessible tools and methods for advancing the reproducibility of research.
3. Reuse and adapt the Reproducibility for Everyone modular curriculum to their own training and research needs.
4. Evaluate their reproducibility barriers and solutions through interactive knowledge sharing.

1054V  A semester-long genetics lab exploring gene families through comparative genomics and CRISPR-based gene editing  Jennifer Kennell Vassar College

Course-based undergraduate research experiences (CUREs) offer students opportunities to gain the benefits of conducting independent research in the context of a course laboratory. Recently my research lab has become interested in studying members of the Haloacid Dehalogenase (HAD) family of non-protein phosphatases. In particular, we began by characterizing in Drosophila possible orthologs of Phosphoglycolate Phosphatase (PGP), an enzyme that was recently identified as playing an evolutionarily conserved role in dephosphorylating toxic side products of glycolysis. Using the tools I have learned through participation in the Genomics Education Partnership (GEP) Pathways Project, I have developed a CURE for my intermediate-level Molecular Genetics class that involves annotation of possible orthologs of these phosphatases in various Drosophila species. Each semester students annotate a different family member and perform evolutionary analyses of the predicted gene and protein sequences. In addition, they also generate a mutation in the D. melanogaster ortholog using transgene based CRISPR-Cas9 gene editing; these mutant lines can then be used in future studies by independent research students in my laboratory. Through the gene annotation portion of the project
students get hands on experience with comparative genomics approaches while reinforcing their understanding of eukaryotic gene structure and evolution. And through the mutagenesis portion of the project they gain experience in setting up fly crosses, analyzing progeny, and conducting molecular analyses to identify possible mutations in the new fly lines they generate. The guide RNA transgenic lines necessary for the project are readily available through various stock centers or are fairly straightforward to generate and make excellent independent projects for undergraduates. This CURE is very flexible and can easily accommodate any gene or gene family of interest. The mutagenesis portion of the project can also easily be incorporated into the existing GEP Pathways Project studying the Insulin pathway.