

Poster Abstracts

POSTER SESSION I

Abstracts 1-22

1

N-Modified Creatine as a Treatment for SLC6A8-Related Autism Spectrum Disorders

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Creatine deficiency syndrome is caused by a mutation in the creatine transporter gene SLC6A8 and results in X-linked intellectual disability and autism spectrum disorder. SLC6A8 mutations are estimated to be as common as 1-5.4% of all X-Linked Intellectual Deficiencies, with this disorder affecting 42,000 patients in the USA and one million worldwide, but without comprehensive genetic testing, this number is likely underestimated. No treatment modality exists, as supplemented creatine cannot cross the blood brain barrier (BBB) or enter neuronal cells without a functioning transporter. Failure of creatine transport into the brain has significant impacts on energy metabolism, as creatine's phosphorylated form is an essential part of the high energy buffering system used to maintain ATP levels in brain tissues. This research introduces a new way to directly screen large numbers of creatine analog compounds in the simple, tractable zebrafish system, in a cost- and time-efficient manner, and has identified potential treatment modalities where none currently exist. We synthesized a new class of N-modified creatine analogs and using high throughput zebrafish screening have accelerated identification of creatine analogs that can cross the BBB, bypassing deficient SLC6A8 transporters. These analogs were biochemically assayed and showed 87-95% efficiency binding creatine kinase and producing phosphocreatine, compared to native creatine. 48-hour embryos were exposed to 125 uM creatine analogs for 24 hours. Brain tissue was isolated, and the amount of creatine analog in the brain was quantified by mass spectrometry. Analogs that both cross the BBB and interact with creatine kinase will be tested in SLC6A8 mutant mice before moving to preclinical trials. Other essential molecules in brain (or somatic) function with defective transporters will benefit from this approach, making our study applicable to the broader field of metabolic imbalance resulting from transporter defects.

2

Bioinformatic and genetic approaches to understanding the cnd-1 regulatory network

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NeuroD1, the vertebrate ortholog of *cnd-1*, is a basic-Helix-Loop-Helix protein involved in neuronal and pancreatic beta cell fate specification. NeuroD1 loss-of-function mutations have been implicated in human visual impairment, learning disabilities, deafness, and neonatal diabetes. In *C. elegans*, *cnd-1* is expressed in many unidentified head neurons and also in D-class motor neurons. Only three genes are known to act downstream of *cnd-1* in *C. elegans*; *unc-3*, *unc-4*, and *unc-30*. All three of these genes are transcription factors that are expressed in D-class motor neurons. However, the remaining downstream targets of *cnd-1* have not been identified. In order to investigate the regulatory role of *cnd-1*, we performed RNA-extractions from wild-type and *cnd-1* loss-of-function embryos, followed by RNAseq. We assembled a transcriptome outlining differentially expressed genes during embryogenesis and are currently following up by validating candidate *cnd-1* target genes using genetic approaches. In addition, NeuroD1 is known to function in a regulatory cascade with neurogenin and also to positively regulate itself in mammalian neuron specification. We seek to verify this relationship in *C. elegans* in order to better understand the regulatory context of our novel *cnd-1* target genes.

3

Drosophila Rbf regulates mitochondrial functions in developing muscles.

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The Retinoblastoma tumor suppressor (RB) regulates cell cycle progression through the suppression of E2F/DP heterodimeric transcription factors. Functional inactivation of RB causes tumor progression in a majority of human cancers. Having an understanding of the molecular machinery involved in both uncontrolled cell growth and cell differentiation is important for further development of treatment options for oncological patients. In *Drosophila*, homologues of mammalian RB, Rbf, and its targets, E2f1/Dp, are critical for cell proliferation and apoptosis. In addition, a conservative role of E2f1/Dp in mitochondrial functioning associated with cell differentiation and tissue development was described both in mammals and flies. Here we investigate whether mitochondrial effects of E2f1/Dp functioning in *Drosophila* developing muscles are directly controlled by Rbf. Using RNA interference approach and the UAS/GAL4 system, we reduced Rbf expression in developing muscles down to 27% of the control transcript levels. Such a deficit of Rbf expression resulted in a significant decrease in the expression of the mitochondrial genes accompanied by changes in mitochondrial morphology. Likewise to reported earlier phenotypes for Dp and E2f1 mutants in *Drosophila*, and Rb1 deficient mouse embryonic fibroblasts, Rbf mutants had thin or fragmented mitochondria. Functionally, these flies showed progressive flight impairments suggesting that mitochondria-enriched flight muscles are compromised. Our data suggest that Rbf is involved in the regulation of the mitochondrial program in muscles, which affects proper muscle maturation in flies. Taking into account the conservative function of Rbf, we will use *Drosophila* model to analyze molecular mechanism that enable Rbf to control mitochondrial genes.

4

Reprogramming Somatic Cells into iPSCs by Novel Cell Penetrating Peptide-Adaptors

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Human disorders and diseases have been captivating the minds of scientists and physicians as long as they have existed. This captivation has led to the discovery of new technology that may eliminate diseases such as degenerative diseases. Cellular reprogramming and the creation of induced pluripotent stem cells (iPSCs) are examples of this young technology. Many advances of this technology have been accomplished through the use of mRNA, viral vectors, and most recently cell penetrating peptides (CPP). Reprogramming by these mechanisms contain many pitfalls, however, including the introduction and possible integration of foreign DNA into cells, tumorigenic effects, and the inability of protein translocation as a result of inefficient CPPs. However, the creation of a novel CPP delivery mechanism has eliminated the need for mRNA, viral vectors, as well as inefficient CPPs. This novel CPP technology utilizes noncovalent bonds between calmodulin and its binding sequence in the presence of calcium, unlike previous CPPs that relied on covalent bonds. Using this CPP adaptor fusion protein, TAT-calmodulin (TAT-CaM) in association with reprogramming proteins, there should be an overall increase in reprogramming efficiency as well as decreased harm to the cells. Preliminary data shows that Oct4, the main transcription factor involved in pluripotency, contains an internal CPP and can enter cells without the aid of an additional CPP. Oct4 and Oct4 bound to TAT-CaM appear to induce pluripotency around the same efficiency. However, the other reprogramming proteins that will be used, Sox2 and Klf4, cannot enter cells without a CPP adapter and thus will require our TAT-CaM mediator to induce pluripotency. Combined, the three reprogramming factors fused with TAT-CaM are expected to yield a higher efficiency of reprogramming compared to previous methods.

5

Investigating the molecular control of the ecdysone response gene, E74, in *Drosophila* ovarian germ cells.

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Oogenesis is the process by which an egg develops from precursor cells in the ovary. This process has been widely studied; however, many of the molecular mechanisms that regulate oocyte development and growth remain unclear. The *Drosophila melanogaster* ovary is an exceptional model system for studying the mechanisms of oogenesis. As in humans, germ cells are surrounded by somatic cells which aid proper oocyte development and maturation. Steroid hormones largely drive this process, and in *Drosophila*, the predominant steroid hormone is ecdysone, similar to human estrogen. Ecdysone binds to a heterodimeric receptor which then functions as a transcription factor to promote gene expression. Other factors, including additional transcription factors and chromatin remodeling factors, likely refine this response. Ecdysone signaling is necessary for oogenesis via the regulation of many target genes. One target, Ecdysone-induced protein at 74EF (E74), is required for oogenesis and is highly expressed in ovarian germ cells; however, regulation of E74 expression in the ovary has not been well-studied. To investigate how E74 expression is regulated in the ovary, we used enhancer mapping to identify regions of the E74 locus critical for germline expression. Twenty-eight fly lines carrying pieces of the E74 gene locus fused to a minimal promoter and Gal4 were crossed with flies containing UAS-lacZ responder transgene. We identified two 200-bp regions within a large intron of the E74 locus that are sufficient to drive expression of a reporter. Together, these regions fully recapitulate the endogenous E74 expression pattern. We then identified several factors, including the chromatin binding factor Trl, as putative regulators of E74 expression at those sites. Trl expression partially overlaps with that of E74, suggesting that Trl may be an important modifier of ecdysone signaling in oogenesis. Future studies will characterize the roles of Trl in this process.

6

Understanding animal polarity: Functional studies during early embryogenesis of the sea anemone

Nematostella vectensis

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How germinal layers are specified during early development of non-bilaterian animals is unclear. In bilaterian animals, rearrangements of the egg's cytoplasm and cortical domains polarize the embryo and direct proper partitioning of maternal determinants into distinct daughter cells often in relationship to a regular cleavage program. In some bilaterian animals, Lethal Giant Larvae (LGL) and PARTitioning-defective proteins (Par) are conserved components of cellular polarization during early embryogenesis. Par proteins and their role in establishing embryonic asymmetry have been widely studied in bilaterian development but not in more basally branching animals. Interestingly, the basally branching cnidarian sea anemone *Nematostella vectensis* shows a "random" cleavage pattern and it undergoes gastrulation at the animal (not vegetal) pole of the egg; begging the question of whether the same molecular mechanisms are conserved for specifying the site of gastrulation. We address this question by characterizing the localization and function of different components of the Par complex during early development of the sea anemone *N. vectensis*. The mRNAs of Par proteins are asymmetrically localized. However, Immunostaining using antibodies made against NvLGL and NvaPKC shows that these proteins distribute throughout the egg and embryo without any clear polarization confirming results obtained when we over expressed them using mRNA injections. In addition, the over expression of the full length and dominant negative version of some Par proteins affect cleavage divisions and gastrulation but do not have a clear effect on embryonic polarity. These data will provide a glimpse into the evolution of cell polarity and the organization of metazoan embryonic germ layer formation.

7

Investigation into the cellular origins of posterior regeneration in the annelid *Capitella teleta*

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Many animals can regenerate, although there is great diversity in relative regenerative capabilities. For example, some animals are able to regenerate their complete body from only a few thousand cells (e.g. planarians), while others can regenerate only a single cell or tissue type (e.g. vertebrates). One of the most intriguing questions in regeneration biology is the cellular source of new tissue that is formed. The polychaete annelid, *Capitella teleta*, displays robust posterior (but not anterior) regeneration following transverse amputation of body segments. However, the source, behavior and molecular characteristics of the cells that form new tissue during regeneration are largely unknown. We hypothesized that the putative primordial germ cell (PGC) niche in *C. teleta* is a source of multipotent progenitor cells. We postulate that following transverse amputation, cells from the PGC niche migrate to the site of injury, and contribute to the regeneration of somatic structures. We used the expression of *Cap1-vasa* as a marker of the PGC niche to examine the characteristics of this cell population, and its dynamics in the first days following transverse amputation in juveniles. *Cap1-vasa* also marks dispersed cells in the coelomic cavity of juveniles. We investigated the hypothesis that these cells are a migratory population originating from the PGC niche. To test whether there is cell migration into the wound site during *C. teleta* posterior regeneration, we used an indirect method involving incorporation of EdU. Finally, we assessed the relative capacity for posterior regeneration in juveniles with and without the PGC niche, by analyzing nerve extension, cell proliferation and number of regenerated segments as markers of relative regenerative capability. This work is the first study in *C. teleta* that addresses the potential source of cells contributing to regeneration of posterior segments, and establishes essential groundwork for future studies.

8

Epigenetic Contributions to Homologous Chromosome Recognition During Meiosis

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During meiosis, homologous chromosomes must correctly identify one another in order for proper alignment and recombination to occur. Improper pairing can lead to chromosomal rearrangements that can result in defective embryonic development. Currently, little is known about how homologs identify each other to the exclusion of the other chromosomes. There has been evidence supporting the role of DNA Double Strand Breaks (DSB) and strand invasion in homolog recognition. However, mutants lacking DSBs still have the ability to properly align homologs suggesting a DSB independent mechanism exists. In *C. elegans*, Pairing Centers (PC) initiate pairing between homologs. Despite their sequence specificity, some PCs are shared between non-homologous chromosomes further suggesting an additional mechanism for recognition. During meiosis distinct patterns of active transcription are produced on each chromosome and are associated with epigenetic modifications such as the methylation of Lysine 36 on Histone H3 (H3K36me). In humans, H3K36me is recognized by the chromodomain containing protein MRG15. Recently, pairing defects were observed in *C. elegans* lacking the MRG15 homolog, *mrg-1*. The specific role of MRG-1 in homolog pairing and recognition is unknown. Our hypothesis is that histone modifications that result from meiotic transcription, including H3K36me, provide an "epigenetic barcode" used to distinguish chromosomes during homolog searching, and is facilitated through the recognition of H3K36me by the chromodomain of MRG-1. We are examining the role of H3K36me in homolog recognition in germlines lacking *mes-4* and *met-1*, the histone methyltransferases responsible for H3K36me. Our recent data demonstrates that germ cells lacking H3K36me exhibit increased sterility and synaptic delay. Similar observations are seen in *mrg-1* mutants. These results suggest that epigenetic modifications such as H3K36me may play an important role in homologous chromosome recognition during meiosis.

SF1 and SF2 boundary functions are essential for Scr and ftz gene regulation

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Chromatin structure plays important roles in gene regulation. In particular, chromatin boundary elements (CBEs), which are genomic regions that interact with each other in the 3-D nuclear space to form chromatin loops, are known to inhibit or promote transcription by modulating access of enhancers to gene promoters. We have previously identified SF1 and SF2, two CBEs in the *Drosophila* homeotic gene (*Hox*) cluster, which determine the animal body segment identity. SF1 and SF2 flank and separate a non-*Hox* gene *fushi tarazu* (*ftz*), from the surrounding *Hox* genes *Sex comb reduced* (*Scr*) and *Antennapedia* (*Antp*). We hypothesized that the loop formed by SF1 and SF2 insulate *Scr* from the *ftz* early enhancers, while facilitating the late *Scr* enhancers to their promoter. To test this hypothesis, we created SF1 and SF2 knockouts using the CRISPR techniques and investigated the *Scr* and *ftz* regulation in these mutant animals. We found *Scr* to be ectopically expressed in the *ftz* pattern in the SF1 knockout animal during early embryogenesis, while in the late embryo the *Scr* expression is normal. Intriguingly, in the SF2 knockout, *ftz* is ectopically expressed in the *Scr* pattern in late embryos, while the early *ftz* expression is normal. Our evidence further indicates that these ectopic expressions contribute to developmental defects and increased lethality in the knockout animals. These results support our hypothesis that the CBEs organize the 3-D genomic architecture and play critical roles in tissue and stage specific gene regulation during animal development.

10

Retrotransposons in mammalian egg-to-embryo transition

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Retrotransposons profoundly impact mammalian gene expression, notably in germ cells. Previously, we reported LTR retrotransposons are massively upregulated during the egg-to-embryo transition in mice, and initiate synchronous, temporally-regulated expression of multiple genes (*Dev Cell* 7:597-606). Recent experimental work demonstrates co-option of retrotransposon promoters directly impacts mouse oocyte quality (*Cell* 155:807-816). We propose that transposons' activity during egg-to-embryo transition is important component of genome reprogramming network, including establishment of novel functional gene modules critical for early development. We hypothesize these networks may vary dramatically among different mammals.

We analyzed RNAseq data of total oocyte and preimplantation embryo transcriptomes to gain insight into transposon contribution to transcriptomes. We analyzed data for three mammalian species, laboratory mice, cows and humans. We programmed pipelines (1) to identify transposons expressed during egg-to-embryo transition, and (2) to discover and map "chimeric" gene transcripts containing alternative transposon-derived exons. Additionally, we applied Gene Ontology (GO)-based pathway enrichment tools to find specific gene network modules.

Our analysis demonstrated profound dissimilarities in classes of transposons expressed during oogenesis and early development among three species, and alternative promoters from transposons drive a large proportion of expressed genes (7%) in all three species. Remarkably, subsets of transposon-driven genes are significantly different among mammals. GO enrichment analysis revealed distinct modules among transposon-driven genes in each species. The profound dissimilarities among gene sets and modules indicate independent origins putatively shaped by natural vs artificial selection. Our findings underscore mammalian oogenesis as an evolutionary "playground" to select for "viable" modules in gene networks.

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Sex specific dynein requirement in *C. elegans* meiosis

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Homologous chromosome pairing and meiotic synapsis are essential processes that are required in both oogenesis and spermatogenesis to prevent aneuploidy and developmental defects in offspring. Despite the importance and high conservation of synapsis, not every aspect is the same between the two sexes. My preliminary results indicate that male and female *C. elegans* have different requirements for dynein components in regulating the initiation of synapsis. Dynein dependent forces have been proposed to test whether a potential homolog match is correct, and once a match has been established, synapsis (SYP) proteins are loaded between the homologs. Knockdown of the dynein light chain (DLC-1) at an elevated temperature results in abnormal SYP aggregate formation away from chromatin in females. Unexpectedly, DLC-1 depletion in males at the same temperature shows grossly normal synapsis. Even more surprisingly, mutants in the heavy chain and dynactin components of dynein also do not show SYP polycomplexes in female meiosis. This indicates that there is a previously undescribed function for DLC-1 in synapsis initiation. Understanding meiotic regulation, and sex specific differences in regulation using a genetically tractable organism will help us better understand natural and disease states in humans that lead to an increased incidence of aneuploidy and meiotically based infertility.

12

The Role of Matrix Metalloproteinases in *Xenopus leavis* Neural Crest EMT and Migration

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The neural crest is a unique cell population found in developing vertebrates, characterized by its multipotency and migratory capacity. During their migration, they first detach from the neuroepithelium through epithelial-to-mesenchymal transition (EMT), then remodel the extracellular matrix (ECM) along their path, resembling the behaviors of metastasizing cancer cells. To explore how neural crest cells remodel the ECM during their migration, we looked at an important matrix metalloproteinase, MMP14 (or MT1-MMP), which is highly expressed both in many malignant cancers and in migrating neural crest cells. As a transmembrane protease, MMP14 plays critical roles in activating other MMPs (such as MMP2), processing various matrix proteins, and mediate signaling events with other cell surface receptors. By loss-of-function approaches, we found that MMP14 is required for cranial neural crest cell migration into the branchial arches. This function in promoting neural crest migration requires both the proteolytic activity of MMP14 and its signaling activity through the PEX domain. To determine whether the surrounding tissues hinder the requirement of MMP14 in neural crest migration, we isolated the cranial neural crest tissue and cultured them in vitro. In culture, loss of MMP14 does not inhibit the spreading of neural crest cells, but cells cannot break apart from each other and migrate individually. So similar to different ADAM proteins, MMP14 may also regulate the expression or turnover of cell adhesion molecules, such as Cadherins.

13

Finding key causative genes in muscle wasting

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Systemic wasting of body mass is a hallmark of cancer. Muscle degeneration particularly presents significant complications in cancer treatments, as it precludes administration of chemotherapy and/or surgical interventions. In many cases, progressive muscle weakness can become a primary cause of mortality in cancer patients. Most commonly seen in sufferers of gastric and pancreatic cancers, tissue degeneration proceeds regardless of increased nutritional uptake. The exact mechanisms behind this wasting are not yet understood, but if discovered, may inform the process of clinical treatment development. When tumors are experimentally induced in the gut of *Drosophila melanogaster*, the flies exhibit targeted degeneration of flight muscles, while other muscle types remain relatively intact. To determine molecular factors that mediate such specific muscle degradation, we have analyzed the changes of gene expression in cancer-responsive muscles before and shortly after the onset of tumor. Based on our analysis, we have selected candidate genes whose activity significantly changes in the presence of experimental tumors. These genes belong to various regulatory networks controlling transcription, hormonal signaling, mitochondrial respiration, and proteolytic degradation. Using the advantage of the *Drosophila* model, we then modulate expression of our candidate genes in an on-or-off manner to recapitulate muscle degradation and death. Because of significant evolutionary conservation of the candidate genes, our study is aimed at revealing novel genetic components in muscle wasting across species.

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Generating mef2ca and mef2cb transgenic zebrafish lines using BAC-mediated recombination and CRISPR/Cas9-mediated integration

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Generating mef2ca and mef2cb transgenic zebrafish lines using BAC-mediated recombination and CRISPR/Cas9-mediated integration

The genes mef2ca and mef2cb (myocyte enhancer factor 2c a and b) are important for craniofacial, muscle, and heart development in zebrafish. The goal of this project is to generate transgenic zebrafish lines expressing fluorescent markers under the control of endogenous regulatory elements for mef2ca and mef2cb. These lines will allow us to study the pattern of expression of mef2ca and mef2cb in development and track the function of these factors in specific tissues and cells in embryos. Using BACs (bacterial artificial chromosomes) containing mef2ca and mef2cb, Phusion PCR products containing the fluorescent reporter genes EGFP and mCherry will be integrated near the start codon of mef2ca and mef2cb using homologous recombination. Likewise, CRISPR/Cas9 will be used to 'knock-in' a fluorescent transgene into mef2ca and mef2cb in vivo so that endogenous regulatory sequences drive transgene expression. These lines should provide insight into the dynamic expression of mef2ca and mef2cb in cells and tissues throughout development of the zebrafish.

15

The role of Wnt Inhibitory Factor-1 in the Wnt signaling network governing anterior-posterior patterning of sea urchin embryos

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The specification and patterning of the anterior-posterior (AP) axis in many metazoan embryos is dependent on a posterior-to-anterior gradient of Wnt signaling. In the sea urchin embryo, patterning of the anterior neuroectoderm (ANE) along the AP axis is dependent on a network involving three interconnected Wnt signal transduction pathways: Wnt/ β -catenin, Wnt-JNK, and Wnt/ Ca^{2+} . While much has been learned about the roles of each of these individual Wnt signaling branches in development and disease, our understanding is still limited and little is known about how they interact with one another in any context. Wnt inhibitory factor-1 (Wif-1), one of the least understood secreted Wnt signaling modulators, has been shown to bind Wnt ligands resulting in Wnt/ β -catenin signaling inhibition. In this study, we report that wif-1 is zygotically expressed in two different germ layers during early AP and dorsal-ventral (DV) patterning in the sea urchin embryo: the endomesoderm and the dorsal ectoderm. Pharmaceutical manipulations suggest that wif-1 expression is activated by Wnt/ β -catenin signaling in the endomesoderm and a mechanism dependent on Nodal signaling in the dorsal ectoderm. We show that perturbing Wif-1 function disrupts gastrulation and specifically perturbs the correct positioning of the ANE along the AP axis, possibly through the inhibition of the Wnt/ Ca^{2+} pathway. Together, these data suggest Wif-1 may represent an important and direct link between the gene regulatory networks that control AP and DV patterning in the early sea urchin embryo.

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The Eph receptor/ephrin pathway is required for AIY interneuron development and food-seeking behavior

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In order to survive, an organism must be able to receive, integrate, and respond to sensory stimuli. However, the cellular basis of sensory perception and response is difficult to study in complex animals such as humans, and is therefore poorly understood. The nematode *Caenorhabditis elegans* is a relatively simple organism yet displays many distinct behaviors, making it an ideal system to understand the relationship between gene function, cell shape, cell physiology, and behavioral output. Much of the thermosensory and chemosensory information that the nematode receives from its sensory neurons is processed via a pair of interneurons called AIYL and AIYR. In wildtype animals, the AIY cell bodies lie just posterior to the pharynx, and extend an anterior process that contacts its contralateral partner at the base of the nerve ring. The AIYL and R processes then diverge and extend around the nerve ring, ultimately making contact again on the dorsal side of the animal via a gap junction. We previously showed that the Eph receptor tyrosine kinase VAB-1 is required for AIY cell body placement and ventral AIYL/R contact. Conversely, the ephrin EFN-4 is required for dorsal AIYL/R connectivity. We have extended these studies and show that the AIYL/R ventral contact is mediated via the ephrin gene efn-1. In addition, we show that this connectivity requires both vab-1 kinase activity and also a non-kinase dependent vab-1 function. To integrate AIYL/R morphology and function with behavior, we are using WormLab software to image and analyze EphR/ephrin mutants both on and off food. Wildtype animals search for food using long "runs" interspersed with reversals and ~170-degree "omega" turns. We find that vab-1 mutants display a strong circling locomotion, both on and off food. We are currently investigating neuromuscular junction morphology in EphR and ephrin mutants to see if this correlates with dorsal versus ventral locomotion bias.

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Fin-folds and autopods share a conserved Shh-Gremlin-Fgf regulatory network

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The morphological transition from fins to limbs involved the loss of the fin-fold (dermal rays) and expansion/remodeling of the distal endoskeleton to form an autopod (hands/ feet) with digits. Recently, we've made observations of an autopodial-like pattern of HoxD expression in paddlefish fin-folds and functional studies in zebrafish demonstrating a role for 5'HoxA/D genes in dermal ray formation. These insights led us to ask if other components of autopod regulatory networks are also involved in fin-fold development. The gene regulatory networks that integrate limb bud outgrowth and patterning have been partially characterized in tetrapods, revealing molecular interactions between the posterior limb bud mesenchyme (i.e., the zone of polarizing activity, ZPA) and the distal limb bud ectoderm (the apical ectodermal ridge, AER). In this network, ZPA-derived Sonic Hedgehog (Shh) acts through LIM-homeodomain transcription factors (LHXs) to induce the BMP antagonist Gremlin. Gremlin, in turn prevents BMP inhibition of AER-derived Fgfs, which maintain ZPA-Shh, which are required for proper patterning of the digits. According to Thorogood's influential "clock" model, delay in conversion of the AER to a fin-fold prolongs the signaling influence of the AER on the endoskeletal mesenchyme, resulting in expansion of fin radials and a reduction of the dermoskeleton. Limbs, which lack a dermoskeleton, reflect the extreme of this hypothesis in that the AER (and its proliferative cues) persist through autopod formation. Herein, we test this hypothesis in the American paddlefish, *Polyodon spathula* through a survey of expression of Shh-Gremlin-Fgf transcriptional network components and use these data to evaluate predictions of clock-based models of appendage evolution.

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Abnormal cardiac patterning and development in akirin mutant embryos

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Akirin is a highly conserved nuclear transcription co-activator that is essential for proper Twist-regulated gene expression during the embryonic myogenesis program. While Akirin has previously been shown to co-regulate the patterning of the skeletal musculature, recent studies have implicated Akirin as a crucial regulator of the tinman locus during specification and patterning of the cardiomyoblasts, the muscle cells that will form the dorsal vessel or heart. akirin mutants display a highly disorganized dorsal vessel, marked by missing cardiomyoblasts, and highly aberrant morphology. We are currently employing fixed embryo and live-imaging techniques to image heart formation in akirin mutant embryos, as well as dorsal vessel contraction in akirin mutants. Our results indicate that akirin mutant hearts are patterned abnormally from the onset of cardiac specification, and that the migration of cardiomyoblasts appears to be negatively affected as a result of a loss of Akirin. Given that Akirin is a highly conserved protein among metazoans, it is likely that these results provide a novel mechanism for cardiac specification and patterning that is similarly conserved from insects to mammals.

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Receptor ALK2, but not ALK3, mediates the regulatory role of BMP signaling in taste organ formation in a mesenchyme-specific manner

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Taste organs, among many epithelial appendages such as feathers, hair follicles, and teeth, require mesenchymal-epithelial interactions for proper development. Among the many molecules and pathways, regulatory roles of bone morphogenetic proteins (BMPs) are important. However, little is known about the involvement of receptor(s) mediating BMP signaling in taste organ development. In the present study, we used mouse models with constitutively activated (ca) BMP receptors in a mesenchyme-specific manner driven by Wnt1-Cre to examine the roles of type I BMP receptors ACVR1 (ALK2) and BMPR1A (ALK3) in the development of tongue and taste papillae. Wnt1-Cre driven caAlk2 and caAlk3 mutants at embryonic day (E) 10.5-11.5 had no observable morphological differences in the embryos compared to littermate controls. At E12.5-P1, caAlk2/Wnt1-Cre tongues were smaller, misshapen, and missing the pharyngeal region compared to littermate controls. In contrast, no obvious change was seen in caAlk3/Wnt1-Cre mice. We found that in caALK2/Wnt1-Cre tongues, fungiform papillae and early taste buds formed in the smaller tongue at E18.5. Immunoreactions on sections using the cell markers E-Cadherin, Vimentin and Desmin showed that both the epithelium, mesenchyme and muscles were disorganized in caAlk2/Wnt1-Cre mouse tongue at E14.5 and E18.5 compared to the littermate control and caAlk3/Wnt1-Cre mutants. Furthermore, caAlk2 and caAlk3 driven by K14-Cre (epithelium-specific) did not lead to an apparent phenotypic changes in taste organs. Our data indicate that BMP signaling plays an important roles in the development of taste organs in a receptor- and tissue-specific manner.

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Gastrointestinal Motility Issues Associated with Autism Spectrum Disorders

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Autism Spectrum Disorders (ASDs) are currently estimated to affect 1-2.6% of children world-wide. Along with the behavioral and developmental issues associated with ASD, gastrointestinal (GI) distress is a commonly reported but poorly understood co-occurring symptom. As a first step towards determining the mechanisms behind ASD related GI disorders, we are using a zebrafish ASD model to gain insight on how genetic variants with high autism relatedness impact GI function. We focus on the high-confidence ASD gene SHANK3; deletions that include SHANK3 are causal for Phelan-McDermid Syndrome (PMS), a form of ASD with GI distress reported in nearly 50% of cases.

The parallels between zebrafish and human GI physiology provide a basis for understanding mechanisms that underlie ASD associated GI distress. As with many zebrafish orthologs of human genes, the human SHANK3 gene is duplicated in Zebrafish. Retention of the two gene copies has been shown to reflect sub-functionalization; consistent with this, shank3a is expressed at higher levels in the brain than shank3b at early developmental (in situ and qPCR), while shank3b is expressed in a more ventral domain corresponding to the upper GI tract. Using a shank3 loss-of-function zebrafish I have compared changes in GI motility between shank3ab mutants and WT fish; tracking and quantifying differences in the frequency of peristaltic movements within the GI tract. Data collected thus far shows that shank3 zebrafish mutants display a significantly slower frequency of peristaltic contractions compared to their wild type counterparts. This is a highly penetrant phenotype since a single shank3b mutant allele is sufficient to significantly reduce the peristaltic frequencies. Our motility findings may relate to GI distress

common to ASD in people based on unpublished clinical studies showing delayed digestive transit times in PMS patients

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mPing as a tool for activation tagging in zebrafish

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The goal of this project is to demonstrate the successful *in vivo* transposition of the mobile element mPing, from *Oryza sativa* (rice), in zebrafish. mPing is a 430-bp, class II miniature inverted-repeat transposable element (MITE), which is mobilized by two enzymes: ORF1, which contains a DNA recognition domain, and TPase, which contains a catalytic DDE domain. mPing, like many invertebrate transposons, has yet to be tested for activity in a vertebrate organism, yet may serve as an effective tool for transposon mutagenesis in vertebrates, such as zebrafish. A single iTol2 expression vector, containing beta actin promoter-driven mCherry (interrupted by an mPing derivative called mmPing20x), will be co-injected with mRNAs for Tol2 transposase and ORF1-T2A-TPase (Pong transposase). The expression vector also contains a *cmc2:EGFP* transgenesis marker labelling cardiac cells, to verify transgenesis. The rate of successful transposition will be determined in injected F0 fish by measuring the ratio of mCherry-positive fish to the number of fish with cardiac EGFP expression. We will establish whether it is possible to remobilize the mmPing20x element in subsequent generations via injection of ORF1-T2A-TPase mRNA into individuals carrying the transposable element. The results of this study will form the basis for future research to use mmPing20x containing a *Xenopus*-derived EF2 α enhancer as an activation tag in zebrafish as a tool for gene discovery.

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Identification of Akirin interacting partners during embryonic myogenesis.

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The specification and differentiation of muscle precursor cells, or myoblasts, by the action of the Twist mesodermal and muscle transcription regulator is a key event in the formation of the *Drosophila* larval musculature. Myoblast population dynamics are tightly controlled by gene expression moderated by this myogenic transcription factor to determine somatic cell fates. Despite the primary importance of myoblast mechanics for building and patterning the musculature, the identities of many molecular players involved in this process remain unknown. Recently we have determined that Akirin, a highly conserved nuclear protein, appears to play a critical role in the regulation of Twist-dependent gene expression during mesodermal specification and muscle development. We hypothesize that Akirin serves as a cofactor to promote interactions between regulatory transcription factors and chromatin remodeling activity to impact gene expression across varying targets. Using a genetic interaction screen in *Drosophila*, we have begun to identify Akirin interacting proteins that participate in the process of muscle specification, patterning, and development. Our screening method has identified a number of proteins that genetically interact with Akirin during muscle patterning in the embryo. Double heterozygous mutant embryos for akirin and one of these potential partners demonstrate a host of deranged or misshapen muscle phenotypes. Thus far we have uncovered a small number of predicted gene products that appear to be involved in general transcription initiation, as well as components of chromatin remodeling complexes. By generating an interactome of its potential partners, we will gain crucial insight into Akirin's mechanism of molecular action during myoblast specification and muscle patterning.

POSTER SESSION II

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Signed, Sealed, and Delivered: CPP Technology Delivers Active Protein Cargo

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TaT-CaM is a recently developed cell penetrating peptide-linker construct that has been shown to successfully penetrate cells with protein cargos of various sizes and properties. This technology uses the TaT cell penetrating peptide (CPP) covalently attached to calmodulin (CaM) which binds specifically to target peptides in high calcium. Advantages include tight but non-covalent binding between CaM on the TaT CPP and its calmodulin-binding-site-tagged cargo in high Ca²⁺ buffer conditions, followed by release of cargo from the CPP after delivery into cells in which cytoplasmic Ca²⁺ concentration is low. Cellular membrane penetration and localization of various proteins with specific trafficking signals has been demonstrated previously. Here we explored the ability of enzyme cargo delivered by the TaT-CaM system to retain activity after cell penetration. Confocal imaging of fluorescently labeled catalase demonstrated successful penetration of BHK cells. Imaging of cells transfected with peroxisomal marker CellLight® Peroxisome-GFP and subsequently incubated for as little as 20 minutes with the TaT-CaM-catalase complex (TaT-Cat) indicate that delivered catalase localizes to the peroxisome. Cell lysates of TaT-Cat treated cells have significantly more catalase activity than control cells, demonstrating successful delivery of active enzyme. We are exploring the effect of the delivered catalase on cells subjected to oxidative stress, in this case treatment with H₂O₂.

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Teratogenic effects of ethanol during in vitro maturation alter gene networks that persist through later stages of development

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Although the deleterious developmental effects of chronic alcohol consumption during pregnancy are well known, on the contrary, the consequences of acute alcohol consumption near conception are controversial. During this developmental period changes in epigenetic modification occur genome-wide, thus ethanol exposure may interfere with formation of proper epigenetic marks. Given the pluripotency of the embryo such misprints may be inherited throughout the body potentially leading to pathologies seen in Fetal Alcohol Syndrome via cellular metabolism. The objective of this study is to identify genes whose expression is altered by ethanol exposure near conception using in vitro egg maturation (IVM) techniques. Changes in gene expression were detected by microarrays (Affymetrix 430 v2.0) and bioinformatics analysis (Gene Ontology, VLAD, MetaCyc etc) in GV, metaphase II oocytes, 2-cell, 8-cell and blastocysts exposed to ethanol (0, 0.05, 0.1% v/v) during oocyte maturation only (FGO stage exposed). These levels mimic blood alcohol concentration after moderate and high (legal intoxication) drinking. Ethanol exposure decreased genes regulating the initiation of translational (Eif2, Eif3, Eif4, and Eif5 family members) from oocytes throughout pre-implantation embryonic development, when typically de novo RNA translation is observed. Many maternal transcripts persisted at the 2-cell stage embryos exposed to ethanol compared to control also suggesting deficiency in mRNA turnover. Likewise, DNA methyltransferases and histone demethylases were also decreased after ethanol exposure from the egg-to-embryo transition. Many genes involved in cellular stress such as mitochondrion transport and heat shock proteins, were increased after ethanol exposure. In summary, ethanol exposure near conception can induce changes in gene expression that persistent to pre-implantation stage altering mitochondrial, translational control, and epigenetic regulators. Funding: Impact Assets, CT.

ZNF845, a C2H2 zinc-finger transcription factor, is required for cnidocyte (stinging cell) development in the sea anemone *Nematostella vectensis*

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Cnidocytes, (stinging cells) are one of the few clear examples of a truly novel cell type. Found only in cnidarians (corals, jellyfish, hydroids, etc), cnidocytes vary in both morphology and function across taxa and are therefore an important diagnostic feature of this group. Development of the explosive organelle (the cnidocyst) requires the expression of several cnidarian-specific structural proteins but expression of these novel genes seems to be regulated by conserved families of transcription factors. In the sea anemone *Nematostella vectensis*, cnidocytes differentiate from a SoxB2-expressing progenitor cell, which also gives rise to neurons. Previously, we have shown that transcription factors from two gene families which are conserved across metazoans (PaxA and Mef2) are required for terminal differentiation in two lineages of cnidocytes in *N. vectensis*. In this study, we show that ZNF845 (a C2H2 zinc finger transcription factor) is expressed throughout the ectoderm during embryogenesis, in a pattern consistent with cnidocyte development and that knockdown of ZNF845 results in loss of cnidocytes. We further demonstrate that ZNF845 is downstream of SoxB2 and upstream of PaxA/Mef2 as knockdown of the former, but not the latter, results in loss of ZNF845-expressing cells. Finally, we show that ZNF845 is not expressed in the SoxB2- or the PaxA/Mef2-expressing cells, suggesting ZNF845 specifically labels an intermediate, cnidocyte-specific, progenitor cell between the SoxB2-expressing "neural" progenitor cell and the differentiating cnidocyte. In the hydrozoan *Hydra vulgaris*, ZNF845 labels a population of hydrozoan-specific stem cells (I cells) which give rise to cnidocytes, neurons, gland cells, and gametes. These results suggest that ZNF845 may have had a role in specifying progenitor cell identity in the stem cnidarian but that its function may have become specific to cnidocyte progenitor cells in the lineage leading to anthozoans (corals and sea anemones).

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Wnt16-Fzl1/2/7-NFAT signaling antagonizes the restriction of the anterior-posterior neuroectoderm in the sea urchin embryo

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Anterior neuroectoderm (ANE) specification, positioning, and patterning is a crucial event in body plan formation in all deuterostomes. Studies from diverse metazoan embryos indicate that Wnt/ β -catenin signaling is essential for the specification and patterning of the neuroectoderm along the primary axis. In early development of the sea urchin embryo, ANE positioning depends on integrated information from the Wnt/ β -catenin, Wnt/JNK, and Wnt/PKC pathways, forming an interconnected Wnt signaling network. We have previously shown that Fzl1/2/7-PKC pathway antagonizes the down-regulation of the ANE GRN by Wnt1/Wnt8-Fzl5/8-JNK signaling in the anterior ectodermal half of early cleavage and blastula staged embryos, allowing for the proper positioning of the ANE territory around the anterior pole. Yet, the exact mechanism by which Fzl1/2/7-PKC signaling antagonizes Fzl5/8-JNK signaling during this process is still unclear. Hence, our research aims to better characterize the Fzl1/2/7 pathway and the GRN it activates to identify possible interactions between these different Wnt signaling branches. Using a candidate gene approach in combination with whole-transcriptome differential screens, we identified a candidate Wnt ligand, Wnt16, a potential transcriptional effector, NFAT, an intracellular signal transduction modulator, Siah1, and several transcription factors in the GRN activated by Fzl1/2/7 signaling. We use morpholino knockdown assays to demonstrate that these regulatory factors are necessary to antagonize the ANE restriction mechanism mediated by Fzl5/8-JNK signaling. Our results indicate that Wnt16 and NFAT are necessary for ANE specification and activation of the putative GRN activated by Fzl1/2/7 signaling. Together, our data suggest that Wnt16 activates the Fzl1/2/7 pathway during ANE positioning and that NFAT acts downstream of Fzl1/2/7-PKC signaling as its transcriptional effector necessary to antagonize Fzl5/8-JNK signaling mediated down regulation of ANE GRN.

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Exploring the Regulation of ftz-f1 Expression in the Drosophila Ovary

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Drosophila melanogaster oogenesis is regulated by the steroid hormone ecdysone. One known transcriptional target of ecdysone is ftz-f1, a nuclear hormone receptor involved in many vital biological processes, including tissue formation. To test whether ftz-f1 is an ecdysone target in oogenesis, our research sought to identify regulatory elements within the ftz-f1 gene locus that contain putative Ecdysone Receptor (EcR) binding sites and are sufficient to drive expression in the ovary. Flies containing transcriptional reporter constructs encoding small regions (tiles) of the ftz-f1 locus, a minimal promoter, and a Gal4 reporter were used to compare ovary cell expression patterns. We computationally identified potential binding sites for EcR in the ftz-f1 locus. A small region within the large intron of the ftz-f1 locus, containing an EcR binding site, is sufficient to drive expression in mid oogenesis nurse cells and oocytes. Additional tiles that contained EcR binding sites showed specific expression in either the late follicle cells or terminal filament cells. Multiple drivers of operculum expression exist within the ftz-f1 locus including tiles that do not have EcR binding sites. One region of the ftz-f1 locus which does not contain an EcR binding site was sufficient to drive expression in discrete populations of somatic cells. Individual tiles within this region correlated to niche cell expression including terminal filament cells, cap cells, and escort cells. Ongoing research aims to further analyze the ftz-f1 specific regions that correspond to discrete patterns of cells expression in order to learn more about the factors that could influence the regulation of ftz-f1 expression.

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Screening for Genetic Factors that Determine Muscle Specialization

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Here we study a very basic question of how similar tissues come to express different genes, which can be seen in muscles destined to perform different functions. In our model organism, the fruit fly (*Drosophila melanogaster*), the large flight muscles in the thorax contract very frequently and can work for hours without fatigue. In contrast, small muscles in the abdomen sparingly contract, and do not support extensive physical load. These two muscle types demonstrate significant differences at the morphological as well as molecular levels. Specifically, flight and abdominal muscles express distinct muscle genes that are important for the same function, muscle contraction. We have characterized a reporter system made on the basis of differentially expressed muscle genes, Act57B and Act88F. This system will be used in genetic screening to identify and potentially unravel important genetic factors controlling the selectivity of gene expression in muscle specialization. We hypothesize that members of chromatin remodeling complexes can be involved in controlling selective gene expression in different muscle types.

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Genetic suppressors of mutant separase may elucidate membrane trafficking role of *C. elegans* separase

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Successful cell division depends on coordinated regulation of chromosome segregation and cytokinesis. Chromosome segregation requires equal partitioning of sister chromatids duplicated in S-phase and held together by the cohesin complex during mitosis. Separase regulates multiple processes during mitotic exit and cytokinesis. The canonical role of separase, the cysteine protease, is to cleave the kleisin subunit of cohesin, allowing chromosome segregation during mitotic and meiotic anaphase. Separase has also been shown to have a non-canonical role in regulating cortical granule exocytosis (CGE) during meiotic division in *Caenorhabditis elegans* embryos. A temperature sensitive separase multination (*sep-1(e2406)*), which results in a single missense mutation (C450Y) within the N-terminal domain of SEP-1, does not localize to cortical granules and is unable to promote exocytosis but results in minimal chromosome segregation defects.

We have performed a genetic suppressor screen to identify separase regulators that rescue lethality of *sep-1(e2406)*. We found multiple intragenic suppressors that alter residues in the N-terminus of SEP-1. These residues are likely to affect structure stability and binding to other proteins. Consistent with previous observations, we have also identified a substantial number of *pph-5* mutant suppressors. These suppressor mutations are found both in the TPR and phosphatase domain of the highly conserved phosphatase PPH-5 and may provide insight into the phosphoregulation of separase function during exocytosis.

We have also identified multiple novel *sep-1(e2406)* suppressors which belong to independent complementation groups, greatly expanding the potential for elucidating separase regulation during membrane trafficking.

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Identification of novel kal-1 transcriptional regulators

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Kallmann Syndrome (KS) is a rare genetic condition that alters olfactory sensation and also hypothalamic neuron migration, which ultimately inhibits reproductive development. We hypothesize that transcription factors required for the expression of known KS genes may be KS loci in their own right. The human kal-1 gene is strongly conserved between vertebrates and invertebrates and when mutated, leads to X-linked KS. Because this gene is not found in rodents, we are using a *C. elegans* model of X-linked KS to identify transcriptional regulators of the kal-1 gene. Our reporter gene and loss-of-function analyses identify the bHLH gene *cnd-1* as a kal-1 transcriptional regulator. As such, we anticipate that a loss-of-function in *cnd-1* will affect ventral enclosure in the same pathway as kal-1. Finally, we have performed a deletion analysis of the kal-1 promoter to narrow down which regions are required for kal-1 transcription at various stages of development and in what specific cell type. These data are being corroborated at single-cell level using kal-1-GFP and histone mCherry co-lineaging data.

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Conserved targets of ISL1 in genital development and binding at shared limb-genital enhancers in chicks

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The early development of limbs and external genitalia involves expression of many of same genes. In addition, ChIP-seq experiments in mice have indicated that many enhancers are active in both tissues. The *Isl1* gene encodes a transcription factor that is required for the initiation of hindlimb buds in mice. Conditional *Isl1* knockouts also show severe impairment in development of the genital tubercle - the embryonic precursor to the penis and clitoris. Similarities in external genital development accross amniotes suggests derivation from their last common ancestor over 300 million years ago. Using ChIP-seq against ISL1 in the genital tubercles of mice and chicks, we reveal a set of conserved enhancers targeted by ISL1 in both species. These genital tubercle binding regions are significantly associated with genes involved in limb development. Furthermore, ChIP-seq against ISL1 in chick early hindlimb buds suggests extensive enhancer sharing between ISL1-expressing hindlimb and genital tubercle cells.

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Regulation of Stem Cell Lineages in *Drosophila* Testes by Notch Signaling

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Spermatogenesis in *Drosophila melanogaster* testes is predicated on the proper interaction between germline cells and their microenvironment somatic cyst cells. Germline stem cells (GSCs) and somatic cyst stem cells (CySCs) are located at the apical tip of the testis where they go through asymmetric division to produce new stem cells and daughter cells. The daughter cells for the GSCs called gonialblasts are fully enclosed by the CySCs daughter cells, the cyst cells. The gonialblasts undergo transit amplifying divisions to produce clusters of precursor cells called spermatogonia that eventually develop into spermatids to produce fertile sperm. Throughout this process, the surrounding cyst cells grow in size and co-differentiate with the enclosed germ cells. The Notch signaling pathway relies on the membrane-bound ligand expressed by the signal-sending cell binding to the Notch transmembrane surface receptor on the signal-receiving cell. This signaling event leads to activation of Notch target genes in the receptor-expressing cell. Notch signaling appears to play a role in the early stages of spermatogenesis. Antibodies against Notch signaling components localize to the apical tip of the testes. Knockdown of Delta in the germline leads to germline loss while hyperactivation of Notch signaling in cyst cells results in failure of cyst cells and germline to differentiate properly. These cyst cells simultaneously express both early and late stage molecular markers and the germline fail to develop into spermatids. We hypothesize that Delta signals from the germline to the Notch receptor on the encompassing somatic cyst cells to prevent germline and cyst differentiation. We propose that activation of Notch in cyst cells prevents their premature differentiation and subsequent germline differentiation.

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The ETS-1 transcription factor in *Xenopus* heart development: Implications for a multi-hit model for HLHS

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Hypoplastic left heart syndrome (HLHS) is one of the most severe human congenital heart defects (CHDs), affecting ~3% of all infants born with congenital heart disease. Although there is strong evidence implicating a genetic etiology, only a few diseasecausing genes have been identified. Progress towards understanding HLHS has been hindered by the lack of genetically engineered animal models. Previously, we have found that loss of transcription factor Ets-1 in *Xenopus* leads to an embryonic lethal cardiac phenotype reminiscent of HLHS: a hypoplastic outflow tract and a thickened ventricular chamber with diminished chamber volume. Here, we further characterized the cardiac phenotype in ETS-1 knockdown embryos. We demonstrate that the HLHS phenotype can be rescued by grafting wildtype cardiac mesoderm progenitor tissue during early stages of cardiac development. Furthermore, loss of ETS-1 causes dysregulation of genes involved in normal endocardial/myocardial signaling. Specifically, knockdown of ETS-1 results in an increase in BMP10, ErbB2 and Myocardin expression during early heart development. These results indicate that loss of ETS-1 causes an HLHS-like phenotype through a multi-hit model involving the cardiac neural crest and endocardium, suggesting the possibility of early intervention for the prevention of HLHS.

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Transposon expression signature in *Nematostella vectensis* development

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Previously we have discovered massive orchestrated upregulation of Long Terminal Repeat (LTR) transposable elements (TEs) during the egg-to-embryo transition in mice. To expand these observations, we focus on elucidating TE expression in oogenesis and early development of phylogenetically distant species across all Metazoa. Here, we report unprecedentedly detailed analysis of TE expression in starlet sea anemone (*Nematostella vectensis*), a representative of Cnidaria, a sister phylum of Bilateria.

N. vectensis is among the simplest model organisms whose cells are organized into tissues. It is an established model organism in evo-devo studies, such as comparison to the development of more complex Bilateria, and is useful in our understanding of early embryonic development. Repetitive sequences comprise approximately 30% of *N. vectensis* genome, most prominent are tandem repeats and DNA transposons. We report differential expression analysis of TEs during *N. vectensis* development using RNA-seq data. We found Kolobok-1_NV, DNA transposon belonging to enigmatic Kolobok superfamily of eukaryotic DNA TEs, as the highest expressed transposon. DNA TEs of Harbinger and piggyBac families, and several retrotransposons of Penelope family, are also among the highly expressed transcripts in early *N. vectensis* development, particularly at the gastrula and early planula stages. Our findings underscore the notion that eukaryotic species' genome variability depends upon unrestricted expression of recently "co-opted" TEs during early development. Funding: Impact Assets, Farmington, CT.

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Parathyroid Cell Fate Instability and Cell Cycle Length

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The parathyroid is the organ responsible for maintaining calcium homeostasis in the body. During mouse development, the parathyroid develops in tandem with the thymus in the 3rd pharyngeal pouch, where the dorsal cells differentiate into parathyroid and the ventral cells differentiate into thymus. Although the fate of thymus cells is quite stable, parathyroid cell fate has been shown to be unstable with parathyroid cells transdifferentiating to thymus cells at low frequencies during late fetal stages. It is unclear whether the parathyroid program is spontaneously shut off and the thymus program subsequently is activated or whether spontaneous activation of the thymus program turns off the parathyroid program. Preliminary data suggests that induction of the master thymus regulator, FoxN1, in parathyroid cells is sufficient to downregulate the parathyroid program; however, we still do not know if the endogenous thymus program has been activated in these cells. Proliferation has been linked to DNA methylation. The thymus program may remain in an unmethylated state and be available for activation due to low proliferation of parathyroid cells during late fetal development. One aim of this research is to determine the length of the cell cycle in the developing parathyroid. In order to devise a strategy to manipulate the cell cycle in parathyroid cells, we are exploring a transgenic mouse line that expresses CyclinD1 under the Keratin5 promoter in hopes that the parathyroid cell proliferation is increased in these mice during embryonic development.

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Comprehensive Big Data Bioinformatics Detects Dynamic Changes in Transposons Expression and Epigenetic Regulators during Transformation

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It is widely recognized that all breast cancers start when some cells in an otherwise healthy tissue begin to look abnormal and ultimately result in full-blown cancer. However, in many cases, initial abnormal cells do not always follow the deadly path and cancer does not develop at all. Little is known as to why some patients diagnosed with atypia and ductal carcinoma in situ (DCIS) remain cancer-free while in others the disease progresses to invasive ductal carcinoma (IDC). To identify molecular signatures driving cell fate decisions at atypia and DCIS stages to transformation, we investigated expression of transposonable elements (TEs) as in human atypia, DCIS and IDC. While mutagenic role of TEs in cancer is well documented, we focus on the novel role of LTRs as potential drivers promoting cell de-differentiation during early stages of tumor development with promise as a prognostic tool. To investigate the contribution of TEs in atypia and DCIS, we created a TE Enrichment Set Analysis (TESA) to identify TEs in RNA sequencing datasets across four stages of breast cancer, normal, atypia, DCIS, IDC in humans (n= 8-23). After quality control steps to remove outliers, TEs, compared to transcripts, exhibit substantially less variation in their expression because the first principle component accounted for over 80% expression variation compared to 20% in transcript expression variation ($p < 0.05$). Eighty-eight TEs were detected as significant across stages by ANOVA ($\alpha = 0.05$, FDR 5%). The majority are LTRs (67%) with the remaining split into DNA TEs (18%), SINEs (11%) and unclassified (4%). Our TESA data complements and provides experimental support that early genomic changes are a mechanism underlying subsequent tumor development. Translational bioinformatics is a technique to identify prognostic molecules for impending invasive breast cancer from biopsies of pre-malignant atypia and ductal carcinoma in situ. Funding: Impact Assets, Hartford CT.

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Determining the development of the parietal eye in brown anoles.

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Circadian rhythmicity controls several physiological and behavioral responses in animals, and in people specifically, disturbances of circadian rhythms underlie mood disorders such as depression and seasonal affective disorder. The proper functioning of the circadian axis in humans and most other vertebrates is dependent on the pineal organ, a neuroendocrine gland that acts as the main synthesizer of melatonin, and the coordination of this hormone's production relies on perception of the photoperiod. However, the mechanisms governing this link between photoreception and melatonin production are not well understood. Because light is perceived in lizards through an additional parapineal structure that is not present in humans, the parietal eye, this extracranial organ offers a unique opportunity to study mechanisms of the pineal complex, as it is easily accessible and can be easily manipulated. As a first step toward better understanding these pineal processes, this project establishes *Anolis sagrei*, the brown anole lizard, as a new model organism by using a histological approach to characterize a timeline of parietal eye morphogenesis, providing a foundation for identifying molecular instrumentation mediating development. The expectation is that, because the parietal eye develops a cornea, lens, and retina similar to those of the lateral eye, the parietal eye and lateral eye will display similar molecular mechanisms of development that illuminate their relationships to the pineal gland.

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Molecular Genetic Analysis of Genes Involved in Tail Bud Development in a Hermaphroditic Vertebrate (mangrove killifish) and Phenotypic Validation in Medaka

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Kryptolebias marmoratus (mangrove killifish) is a hermaphroditic vertebrate that inbreeds to genomic isogeny amenable to efficient and robust forward mutagenesis. Two recessive mutants were previously identified by genetic screening as shorttail (*stl*) and balltail (*btl*) phenotypes during embryonic development. Total RNAseq of mutants, siblings, and their wild type progenitor (99.97% homozygous) uncovered ENU induced missense mutations in the homologous genes *noto* (*stl*) and *msgn1* (*btl*). In situ hybridization patterns of notochord (*col9a1b*), somites (*hsp90aa*), spinal cord (*sox3*), and tail bud (*spt*) native markers in these mutants, revealed suppression of notochord, somite, and spinal cord by *stl*, whereas *btl* suppresses somites, while expanding notochord expression in the developing tail. Neither mutant affected the expression of *spt*, suggesting these genes are specifically involved in regulating the formation of tail between trunk and terminal posterior axis margins. Further analysis of *noto* and *msgn1* expression amongst these mutants demonstrates *stl* suppresses *msgn1* in the developing tail. To validate the above genetic analysis, we injected morpholinos and analogous mutant mRNA alleles in one cell stage embryos, followed by rescue with wild-type mRNA to phenocopy our results in medaka. Likewise, tail bud cell fate was marked in wild-type versus morpholino injected medaka embryos with Kaede fluorescence to uncover a crucial role during tail bud development in the formation of axial (notochord) and paraxial mesoderm (somite). We propose a model whereby *noto* initially organizes stem cells and *msgn1* positively regulates the formation of somites while suppressing notochord in the developing tail.

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Nociception, and the Experience of Pain Signaling Due to Nerve Damage

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Neuropathic pain is difficult to repair and alleviate. Pain is experienced through the integration of neuronal circuits involved with nociceptive signaling. These harmful stimuli are encoded and processed through a specific group of mechanosensory TRPA1 channels. Those who suffer from neuropathy-based pain attempt to relieve pain symptoms with prescription analgesic medications. Most of these medications come with detrimental side effects including a triggering of the brain's reward and addiction pathways. Taking these drugs for an extended period of time results in dependency problems. In an effort to understand the neurophysiological changes associated with neuropathic pain sensation and to understand how these channels respond during pain stimuli, I will closely examine the effects of suprathreshold stimuli on the TRPA1 channel and manipulation with the putative neuropathic analgesic, THC (delta-9-tetrahydrocannabinol). TRPA1 channel activity will be assayed using the zebrafish model through electrophysiological recording, behavior recoding, and anatomic changes in TRPA1 neuron anatomy and connectivity. Specifically, I will use electrophysiological methods to record noxious signal modulation in zebrafish from the TRPA1 channel containing cranial nerve VIII while in the presence of THC.

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Application of Tol2-based Activation Tag Constructs for Zebrafish Mutagenesis

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Transposons are segments of DNA that can move from one region to another within the genome. The Tol2 transposon from Medaka fish has successfully been used for transgenesis, integrating foreign DNA, into a wide variety of vertebrates. Our goal is to develop Tol2 into a mutagenesis tool for gene discovery. Mutagenesis by transposon insertion, called transposon tagging, enables the discovery and analysis of gene function by causing mutations. Activation tagging, a type transposon tagging, is when a strong enhancer is positioned within the transposon. Activation tagging is used to learn about the function of genes by inducing overexpression. This is significant because many genes may otherwise be hard to study because of lethality or redundancy. Activation tagging has never been used for zebrafish, but it is commonly used for gene discovery in plants.

Zebrafish can serve as vertebrate development models, therefore activation tagging within zebrafish allows for the discovery of genes that are important for vertebrate development. A Tol2-based activation tag, with a h2afx promoter sequence inserted in the middle of Tol2 terminal inverted repeats (TIRs), was engineered using various molecular biology techniques (PCR, digestion, and sequence analysis). Additionally, a DNA construct encoding Tol2 transposase, which will allow transposition of the activation tag to occur, was produced. The integration of both constructs into zebrafish embryos is being performed to measure transposition rates and look for altered gene function. To develop more active constructs for zebrafish mutagenesis, yeast transposition studies are also being performed in order to identify methods to increase transposition rates.

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Elucidating the Role of Securin Regulating Separase during Cortical Granule Exocytosis

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Meiosis is a tightly regulated process leading to the production of haploid gametes. A key player in this process is separase (SEP-1). Known for its canonical role in chromosome segregation, studies suggest that SEP-1 has an additional function in vesicular trafficking during cell division. We hypothesize that cell cycle machinery known to control SEP-1 activity for chromosome segregation also controls its localization to the cortex and subsequent exocytic activity. Following spindle attachment and chromosome alignment during the meiotic M phase, the anaphase promoting complex (APC/C) is activated, resulting in the degradation of SEP-1 inhibitory chaperone securin (IFY-1) and entry into anaphase I. We have observed that SEP-1 localizes to cortical granules and regulates their exocytosis during anaphase I, which is necessary for eggshell formation. Before it appears on cortical granules, SEP-1 localizes to cytosolic filaments near the plasma membrane. We have shown that SEP-1 colocalizes with IFY-1, on filaments during prometaphase, and both disassociate from these structures during anaphase I. Inhibition of APC/C activity prevents SEP-1 and IFY-1 from leaving the filaments. These data suggest that degradation of IFY-1 may regulate SEP-1 localization to vesicles. To address whether IFY-1 degradation is required to allow SEP-1 translocation onto vesicles, we generated a non-degradable IFY-1 (IFY-1DM::GFP). IFY-1DM::GFP is not completely degraded following anaphase I onset, remaining on chromosomes and in the cytoplasm into anaphase II. Expectedly, IFY-1DM::GFP causes embryonic lethality. Interestingly, IFY-1DM::GFP causes polar body extrusion failure, which could be related to defects in cortical granule exocytosis. We will investigate how IFY-1DM::GFP affects SEP-1 localization to cortical granules. This will provide insight into how key regulatory components of the cell cycle control SEP-1 localization to promote timely cortical granule exocytosis during anaphase I.

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Characterization of a Piglet Model of Traumatic Brain Injury Utilizing Non-Invasive Magnetic Resonance Imaging and Histological Assessment

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Traumatic brain injury (TBI) is a major cause of death and disability in the United States, chiefly affecting children ages 0-4 years. TBI at such a young age may lead to long-term neurological deficits. Recent failures in translatable research suggest a more human-like animal model, like a piglet, is necessary for developing an effective therapy. Magnetic resonance imaging (MRI) and histological assessments are pertinent in the comprehensive understanding and treatment of TBI at the tissue and cellular levels. We hypothesized that controlled cortical impact (CCI) would result in a concussive piglet TBI model with substantial changes in lesion and hemisphere volume coupled with histological changes that persist over time. TBI was induced in six male piglets, with MRI scans conducted 24 hours and 12 weeks post-TBI. Histological changes were observed by quantifying NeuN+ neurons, GFAP+ astrocytes, and Iba1+ microglia in the cortical peri-lesion area through 12-weeks post-TBI. Lesion size was significantly reduced comparatively at 12 weeks with a significant change midline shift as compared to 1 day post TBI. There was a significant ($p < 0.01$) decline in NeuN+ neurons beginning 1-week post-TBI. GFAP+ astrocytes increased significantly ($p < 0.0001$) from normal starting 1 day post-TBI. Lastly, Iba1+ microglia increased significantly ($p < 0.05$) at each timepoint. The observed directional change in midline shift and decrease in lesion size can be attributed to attenuated swelling and significant brain atrophy. The noted histological changes suggest there was significant cell death and there was a significant upregulation in GFAP+ astrocytes and Iba1+ microglia, which suggests TBI leads to gliosis and an inflammatory response that mounts over time. The characterization of key cytoarchitectural changes in the CCI TBI piglet model will enable more robust and predictive assessments of novel therapeutics that will likely lead to more success in human clinical trials.

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Determining the role of *ldlrp1a* in zebrafish skeletal development

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A line of mutant zebrafish containing a jaw mutation named b1187 was discovered during a forward genetics screen. This mutation is characterized by fused joints and abnormal shaping in cartilage and bone in the craniofacial region of zebrafish. To find the gene behind the b1187 mutation multiple genes were sequenced. Although there were no differences between mutant and wild-type sibling cDNA, the phenotype was closely linked to the *ldlrp1a* gene locus (low density lipoprotein receptor adaptor protein 1a). This led to a reverse genetics approach using the CRISPR/Cas9 system to create a line of zebrafish with a mutated *ldlrp1a* gene. *ldlrp1a* is known to be involved in cholesterol signaling, however it may also have a role in skeletal development. A F0 generation containing an *ldlrp1a* mutation was generated and was then crossed with wild-type siblings to create three separate F1 generations. The F1 generations were screened using PCR and T7 endonuclease digest to identify approximately half of the offspring who were heterozygous mutants for the *ldlrp1a* gene. Fin clip samples were taken from all three individual heterozygous carriers and a wild-type zebrafish and these samples were sent for sequencing. Of the three heterozygote carriers, one appeared to have a favorable 7 base pair deletion. This sequenced fish was then crossed to a wild-type zebrafish to create an F2 generation. In-crosses between F1 mutant carriers and histological stains of offspring are in progress. If we observe jaw abnormalities resembling the b1187 mutation we could conclude that *ldlrp1a* is not only involved in cholesterol homeostasis, but may also be involved in craniofacial development.

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Casein Kinase 1 delta/epsilon mediates anterior-posterior axis formation in the sea urchin embryo, potentially through localized activation of Disheveled

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Wnt signaling plays a central role in establishing anterior-posterior (AP) polarity in metazoan embryos. A key cytoplasmic component mediating Wnt signaling is the Disheveled (Dvl) protein. In the sea urchin, Dvl is highly enriched and differentially post-translationally modified in a specialized vegetal cortical domain (VCD) of the egg, and the vegetal blastomeres that inherit the VCD during embryogenesis. Functional analysis has shown that localized Dvl activity mediates canonical Wnt signaling in vegetal blastomeres, but the molecular basis of Dvl asymmetric localization and activation remain unresolved. Therefore, identification and functional characterization of proteins interacting with Dvl (DIPs) in the VCD will help us better understand how Dvl partners regulate Dvl activity and Wnt signaling. By applying Dvl Co-immunoprecipitation coupled with mass spectrometry we have identified several potential Dvl-interacting-proteins (DIPs) from isolated egg cortices and 16-cell-stage micromeres. Casein Kinase 1 δ/ϵ (CK1 δ/ϵ), one of our newly identified DIP candidates, is highly enriched and co-localized with Dvl at the vegetal pole of the sea urchin embryo. Downregulation of CK1 δ/ϵ activity by overexpressing a dominant-negative form of CK1 δ/ϵ resulted in the downregulation of genes expressed in the endomesoderm and the anteriorization of embryos. However, overexpression of CK1 δ/ϵ by injecting synthesized CK1 δ/ϵ mRNA into fertilized eggs only induced slight upregulation of endomesoderm genes and mild posteriorization of embryos. Intriguingly, we found that co-overexpressing CK1 δ/ϵ and Dvl induced significantly higher levels of expression of endomesodermal genes compared to expression levels of these genes in embryos overexpressing Dvl or CK1 δ/ϵ only suggesting that CK1 δ/ϵ synergizes with Dvl to positively regulate Wnt signaling. This work establishes CK1 δ/ϵ as a critical regulator of Dvl activation and AP axis specification in sea urchin embryos.

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Interactions of Akirin and Muscles Wasted during myogenesis

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We have recently identified the highly conserved nuclear co-factor Akirin as an essential partner during the process of Twist-mediate gene activation during embryonic myogenesis. akirin mutants display multiple defects in muscle patterning, with missing, misattached, and/or duplicated muscles. Live imaging data indicates that the muscles that do form are morphologically thinner and weaker than those observed in wild-type sibling embryos, and that these muscles rapidly deteriorate prior to hatching. These akirin mutant phenotypes were reminiscent of muscles wasted (mute) mutants; the specification, positioning, and patterning of mute mutant muscles initially form, but rapidly degenerate as the embryo nears hatching. Despite these phenotypic similarities, a connection between the two had yet to be described. Using a combination of confocal-based live imaging of developing embryos, as well as analyzing whole-mount fixed embryos, we have confirmed a genetic link between these loci. akirin/mute double heterozygous mutant embryos display a profound disorganization of the embryonic muscle pattern, with severely degenerated muscles, large numbers of unfused myoblasts, and abnormal patterning and formation of muscle groups in pre-hatching embryos. While a direct interaction between these two gene products is currently under investigation, these data strongly indicate a potential interaction during the myogenic process.

Using CRISPR/Cas9 to study the role of *zmym2* and *zmym3* in zebrafish craniofacial development

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Potocki-Shaffer syndrome (PSS) is a rare contiguous gene-deletion caused by heterozygous interstitial microdeletions of chromosome region 11p11-p12 and is characterized by developmental defects that include intellectual disability and craniofacial anomalies. PSS is associated with mutations in genes encoding factors in the PHF21A protein complex, including KDM1A (lysine-specific histone demethylase 1A), ZMYM2 (zinc finger protein 198), and ZMYM3 (zinc finger protein 261) proteins. It is hypothesized that these protein complexes are involved in craniofacial development of zebrafish in a way that reflects their function in humans. Previously, F0 founder fish carrying mutations in *zmym2* and *zmym3* were generated at the 1-cell stage. Founders were screened by PCR and T7 endonuclease digest which identifies mutations in the DNA and were used to generate F1 lines. The F1 generation was screened by using tail fin DNA in PCR and T7 endonuclease digest. F1 zebrafish were sequenced and frameshift mutations were identified. Zebrafish with confirmed frameshifts will be out-crossed to produce an F2 generation. The F2 generation, of which 25% are expected to be homozygous mutants, will be studied at 7 days post fertilization for anatomical abnormalities in craniofacial development by using Alcian Blue and Alizarin Red histological stains for cartilage and bone. The work in this project will be used to identify the roles of *zmym2* and *zmym3* in zebrafish development, and how a loss of function of these factors may underlie the defects seen in PSS.