Annotation of Contig19 in Drosophila eugracilis

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Heterochromatin domains are regions found in the genome where DNA is densely packaged. In most cases, genes in these regions are silenced, presumably due to limited access of the transcriptional machinery. The small fourth chromosome of Drosophila, also known as the dot chromosome or F element, is unusual in exhibiting many characteristics of heterochromatic domains, and at the same time maintaining some euchromatic properties. There are 80 genes actively transcribed on the F element. To examine the regulation of these genes and the evolution of this unusual domain, a comparative study is being undertaken by manually curating gene models and identifying motifs specific to the F element using phylogenetic footprinting for species diverged recently (10-15 Myr) from D. melanogaster.

This project focuses on contig19, a 40 kb region of the dot chromosome in D. eugracilis. The D. melanogaster genome has been used as a reference to annotate all features. A variety of bioinformatics tools and databases were used during annotation, including UCSC Genome Browser GEP Mirror, FlyBase, BLAST, ab initio gene predictors, TopHat, RNA-seq data, DHS data, RAMPAGE data, BG3 and S2 data on DH sites, and RNA PolII CHIP-seq data. Four genes on this contig, Zyx, PlexA, ATPsynbeta, and CaMKII, were annotated. Comparative analysis indicates that PlexA, ATPsynbeta, and CaMKII are highly conserved between D. eugracilis and D. melanogaster while Zyx is less conserved and might have a new isoform. The transcription start site, repetitious elements, synteny and potential regulatory elements of each gene are being examined in this project, looking for any special characteristics of genes within the F element. These data can help us understand how genes in densely packaged DNA can be expressed when typical genes are generally silenced in such a heterochromatic environment.