

Washington University in St. Louis

Washington University Open Scholarship

Volume 12

Washington University
Undergraduate Research Digest

Spring 2017

Driving Gene Expression in the Heterochromatic Environment of the Fourth Chromosome of *D. melanogaster*

Jacob Cantrell

Washington University in St. Louis

Follow this and additional works at: https://openscholarship.wustl.edu/wuurd_vol12

Recommended Citation

Cantrell, Jacob, "Driving Gene Expression in the Heterochromatic Environment of the Fourth Chromosome of *D. melanogaster*" (2017). *Volume 12*. 25.

https://openscholarship.wustl.edu/wuurd_vol12/25

This Abstracts A-I is brought to you for free and open access by the Washington University Undergraduate Research Digest at Washington University Open Scholarship. It has been accepted for inclusion in Volume 12 by an authorized administrator of Washington University Open Scholarship. For more information, please contact digital@wumail.wustl.edu.

DRIVING GENE EXPRESSION IN THE HETEROCHROMATIC ENVIRONMENT OF THE FOURTH CHROMOSOME OF *D. MELANOGASTER*

Jacob Cantrell

Mentor: Sarah Elgin

Genomes of higher eukaryotes can be divided into two fundamental and dynamic subtypes: euchromatin and heterochromatin. The fourth chromosome of *Drosophila melanogaster* is of particular interest because, despite the fact that it appears to be entirely heterochromatic, the genes within this densely packaged chromosome are expressed. Genes that are active in a euchromatic environment are silenced when transposed to heterochromatin. Our aim is to identify heterochromatic gene regulatory elements that drive their active transcription.

Insertion of a euchromatic *hsp70-white* transgene, which exhibits a uniform red eye phenotype when present in euchromatin, into heterochromatic regions on the fourth chromosome results in sporadic silencing, or Position Effect Variegation (PEV). We replaced the *hsp70* promoter of *hsp70-white* with a genomic fragment of a highly expressed fourth chromosome gene, *Rad23*. The fragment includes the *Rad23* promoter region and ~1 kb of an upstream sequence. Insertion of the *Rad23-white* transgene into the same location on the fourth chromosome switched the *hsp70-white* PEV phenotype to a uniform red eye, suggesting that the *Rad23* fragment is sufficient to drive strong expression of the euchromatic *white* reporter.

A series of transgenic reporter constructs containing fragments of varying lengths of the *Rad23* promoter region were prepared by molecular cloning and injected into embryos of the *Drosophila melanogaster* test line. Results show that the promoter region of *Rad23* containing 100 base pairs upstream of the TSS is sufficient to achieve full expression of *white* in a heterochromatic environment. However, a smaller fragment (only 50 bp upstream of the TSS) resulted in a fly with a completely white eye, indicating the loss of a key promoter element. Additional experiments to narrow down what elements of the noncoding regulatory region drive the expression of the reporter gene, and what constructs produce a PEV phenotype, are underway.