My summer research project focuses on threetie basis of barriers that separate two lineages. Oftentimes, hybrids between tweckes suffer from reduced fitness, which is manifested in their reduced abilito reproduce and survive. Previsostudies have concluded that the cause of greatly reduced viability and feytilit hybrids may be attributed to abnormal amounts of gene product, a result of irregular gene expression. My summer project plans to address the mechanism by which geneseither under- or over-expressed in hybrids.

Our hypothesis for the incorrect gene expressin hybrids is that a divergence in the way DNA is packaged in the two parent species timbutes to the gene expression problem, such that DNA conformation in chromosomes of hylsrid flawed. The way in which chromatin, the DNA-protein complex in nuclei, is packageddely influences gene expression. Depending on whether an area of chromatin is loosely or tightackaged, genes may or may not be expressed and lead to proper appearance of threesoponding phenotype. One way to assess DNA packaging is to look at the expression of a transgengene that has been engineered and inserted at a particular location (in our case, on the boodlewosely and tightly packaged chromatin) of the genome. By looking at the levels of expression of the transgene purposefully inserted in hybrids we can assess whether the DNA conformation between two parent species is altered. My summer research applies this approach in the through system of a fruit fly, into which we have inserted a transgenwhitegene) that controls eve pigmentation in the fourth chromosome telomere. If the transgene is expressed in a fevore. It will be red, and if the transgene is silenced in a fly eye cell, the cell will be whitehus, a fly eye can have a variegated phenotype, patches of red and white, for its eye. These inarlevels of pigmentation will be compared between parent species and their hybrid progeny.

Our methods to test our hypothesis consist of a three-step approach. First, female Drosophila melanogasterirgins were collected. Then thesiegin females were crossed with different stocks of another fruit fly specie Brosophila simulan) smales. A one to one ratio of female virgins and males was used to perform sensand forty of each gender were put into vials and incubated at 20 degrees Celsius. Hybrid wiese then collected envy few days and frozen so that we could assess eye pigmentation. Once all hybrid flies were collected, pictures of their eyes were taken under a microscope and uplotate domputer. IMAGEJ, a software program, allowed us to analyze average pixel value (assueement of pigment) in the hybrid eyes. The darker the eye the lower the asset pixel value, and the lighter the eye the higher average pixel value. These values were then compared tax were pixel values of the parent species.

Thus far, we have completed crosses and eye pigment analyses for Australia and Malawi background hybrids. These results indicate thatily ruit flies have consistently less expression of the white transgene than the pure species parents (#i1). This supports the hypothesis that