Trans regulation of an odorant binding protein by a proto-Y chromosome affects male courtship in house fly

Pablo J Delclos†1,2, Kiran Adhikari†1,3, Alexander B Mai1, Oluwatomi Hassan1, Alexander A Oderhowho1,4, Vyshnika Sriskantharajah1,5, Tammie Trinh1,6, Richard P Meisel1

†Co-first author/equal contribution

Affiliation: 1Department of Biology & Biochemistry, University of Houston, Houston, TX, USA
2Current affiliation: Department of Natural Sciences, University of Houston-Downtown, Houston, TX, USA
3Current affiliation: Oxford Biomedica Solutions LLC, Bedford, MA
4Current affiliation: Department of Neurosurgery, Ochsner LSU Health Shreveport, Shreveport, LA, USA
5Current affiliation: University of Texas Health Science Center School of Biomedical Informatics, Houston, TX, USA
6Current affiliation: University of Texas School of Dentistry, Houston, TX, USA
Y chromosomes have male-limited inheritance, which favors the fixation of alleles that affect spermatogenesis, courtship, and other male-specific traits. Y-linked male-beneficial alleles can also have female-deleterious (sexually antagonistic) effects because they never experience direct selection in females. However, determining the mechanisms underlying these male-beneficial effects is challenging because it can require studying Y-linked alleles while they still segregate as polymorphism. We used a Y chromosome polymorphism in the house fly, *Musca domestica*, to address this challenge. Two common male-determining Y chromosomes (YM and III\textsuperscript{M}) segregate as stable polymorphisms in natural house fly populations, and they differentially affect multiple traits, including male courtship performance. We performed a meta-analysis of RNA-seq data and identified differentially expressed genes encoding odorant binding proteins (in the *Obp56h* family) as candidate causal agents in the courtship differences. The *Obp56h* genes are not found on either the YM or III\textsuperscript{M} chromosomes, suggesting that they must be regulated in trans by one of the house fly sex chromosomes. Using a network analysis and allele-specific expression measurements, we identified multiple genes on the house fly III\textsuperscript{M} chromosome that could serve as trans inhibitors of *Obp56h* gene expression. One of those genes is homologous to *D. melanogaster* CG2120, which encodes a transcription factor that binds both up- and downstream of *Obp56h*. We found that up-regulation of CG2120 in *D. melanogaster* nervous tissues reduces copulation latency, consistent with this transcription factor acting as a negative regulator of *Obp56h* expression. We propose the name *speed date* (spdt) for CG2120, with the house fly homolog named *Md-spdt*. The expression of *spdt* across *D. melanogaster* development and tissues suggests that evolution of higher expression in neurons may be constrained by pleiotropic or sexual antagonistic effects. We hypothesize that a cis-regulatory allele that increases expression of *Md-spdt* on the III\textsuperscript{M} chromosome exists because Y-linkage of this allele releases it from those constraints. This provides evidence for a molecular mechanism by which a Y-linked gene can evolve a male-beneficial function regardless of the negative effects on females.
INTRODUCTION

In species with genetic sex determination, a Y or W chromosome can have sex-limited inheritance (Bachtrog et al. 2014; Beukeboom and Perrin 2014). The male-limited inheritance of Y chromosomes is predicted to allow for the accumulation of alleles with male-specific beneficial effects (Rice 1996). These male-beneficial alleles can have female-deleterious (sexually antagonistic) effects because they are never exposed to direct selection in females (Charlesworth et al. 2005; Abbott et al. 2017).

One important way that sex-limited Y and W chromosomes appear to affect sex-specific traits is via trans regulation of genes elsewhere in the genome. For example, Y chromosome genotypes in Drosophila melanogaster have trans effects on gene expression throughout the genome, which modify a broad range of phenotypes, including immunity and chromatin (Lemos et al. 2008, 2010; Brown et al. 2020). D. melanogaster Y chromosome genotypes also have fitness consequences that depend on the genetic background, suggesting epistatic interactions between Y-linked alleles and the X or autosomes (Chippindale and Rice 2001). Similarly, in Poecilia spp., Y-linked alleles may affect sexually selected male pigmentation patterns by acting as trans regulators of autosomal gene expression (Morris et al. 2020; Kawamoto et al. 2021; Sandkam et al. 2021), although the specific mechanisms of these effects are not well understood.

Even though we are aware of sex-specific phenotypic and fitness effects of Y and W chromosomes, the mechanisms underlying these effects are not as well understood. Notably, we have a limited understanding of how Y and W chromosomes act as trans regulators of sex-specific and sexually antagonistic traits genome-wide. There are also very few sexually antagonistic alleles that have been genetically or molecular characterized on young sex chromosomes (cf. Roberts et al. 2009), which limits our ability to make generalizations about the molecular mechanisms underlying sexually antagonistic selection on Y and W chromosomes (Mank et al. 2014). To address these shortcomings, we sought to identify the genetic mechanism underlying how a young Y chromosome affects male courtship behavior in the house fly, Musca domestica.

House fly has a multifactorial sex determination system, in which multiple young proto-Y and proto-W chromosomes segregate as polymorphisms in natural populations (Hamm et al. 2015). The two most common proto-Y chromosomes (III^M and Y^M) are distributed along latitudinal clines on multiple continents (Franco et al. 1982; Tomita and Wada 1989; Hamm et al. 2005), and they affect thermal traits in ways that are consistent with their geographic distributions (Delclos et al. 2021). This polymorphism has remained stable in natural populations since at least the mid-20th century, suggesting that selection maintains multifactorial sex determination (Kozielska et al. 2008; Meisel et al. 2016).
Here, we focus on the effect of the house fly proto-Y chromosomes on male courtship performance. Males carrying the III\textsuperscript{M} chromosome (III\textsuperscript{M} males) outcompete males carrying the YM chromosome (YM males) for female mates (Hamm et al. 2009). In light of this and other aforementioned phenotypic differences between YM and III\textsuperscript{M} males, it is remarkable that the III\textsuperscript{M} and YM chromosomes carry nearly all of the same genes as their homologous proto-X chromosomes (III and X), and only a small number of allelic differences have been identified (Meisel et al. 2017; Son and Meisel 2021). These similarities between the proto-Y and proto-X chromosomes has led to the hypothesis that the phenotypic effects of the proto-Y chromosomes may be mediated by \textit{trans} effects of YM and III\textsuperscript{M} alleles on the expression of genes elsewhere in the genome (Adhikari et al. 2021). We aimed to identify the \textit{trans} regulatory allele(s) on the YM or III\textsuperscript{M} chromosomes that affect differences in courtship performance between the genotypes.

**METHODS**

\textbf{RNA-seq differential gene expression analysis}

We analyzed published RNA-seq data from \textit{M. domestica} male heads (NCBI Gene Expression Omnibus accessions GSE67065, GSE126685, and GSE126689, shown in Supplementary Table S1). The RNA-seq data include nine YM and fifteen III\textsuperscript{M} samples (Meisel et al. 2015; Son et al. 2019). We assigned RNA-Seq reads to house fly transcripts from genome assembly v2.0.2 and annotation release 102 (Scott et al. 2014) using kallisto in single-end read mode (Bray et al. 2016). All RNA-seq reads were single-end, and we set the average fragment length to 300 bp and standard deviation to 30 bp for all samples. Our expression analysis was performed at the level of transcripts (as opposed to genes), which should not affect our conclusions because all of our focal genes each produce one annotated transcript.

We tested for differentially expressed (DE) transcripts between males with a YM chromosome and males with a III\textsuperscript{M} chromosome using a combination of DESeq2 (Love et al. 2014), sva (Leek et al. 2012), and limma (Ritchie et al. 2015). We only included transcripts that passed an initial threshold filter of 0.5 counts per million in at least 4 samples. Read counts for each of those transcripts were normalized by variance stabilizing transformation in DESeq2. To remove batch effects across data sets, we used the sva package to identify and estimate surrogate variables that adjust for latent sources of variation (e.g., batch effects). To identify DE transcripts between YM and III\textsuperscript{M} males, we used the lmFit() function in limma to fit a linear model comprised of male type (YM vs III\textsuperscript{M}) and our surrogate variables as fixed effects, and read counts as the response variable. We then computed contrasts between male types and calculated test statistics using the eBayes() function. Transcripts below a false discovery rate (FDR) adjusted \( p \) value \( (p_{\text{adj}}) \) of 0.05 were categorized as DE (Benjamini and Hochberg 1995).
Weighted gene co-expression network analysis

We used weighted gene co-expression network analysis (WGCNA) to identify modules of house fly transcripts whose expression correlates with male type (YM or III\textsuperscript{M}) on normalized read count data that were adjusted for batch effects in sva (Langfelder and Horvath 2008). For all pairs of transcripts with variable expression across samples, we calculated Pearson’s correlation coefficient across all samples. We created an unsigned correlation matrix and adjusted the soft-threshold value ($\beta$) to which among-transcript covariances are exponentially raised to ensure a scale-free topology (this resulted in $\beta = 7$), thereby creating a weighted network of gene expression. An unsigned matrix allows us to identify connected transcripts whose expression is either positively or negatively correlated. Within this topological overlapping network (Li and Horvath 2007), transcripts were hierarchically clustered, and modules were identified based on the degree of similarity among transcripts. We used a merging threshold of 0.2, with a minimum module size of 30 and a mean connectivity threshold of greater than or equal to 0.7. We used the default parameters of WGCNA for the rest of the analyses. We then correlated module eigengene values for a given module across samples via Pearson’s correlation and identified modules differentially regulated between male types at FDR-adjusted $p < 0.05$.

To visualize WGCNA genetic covariance results among modules significantly associated with male type, we exported final co-expression networks to Cytoscape (Shannon et al. 2003). We attached information on log\(_2\) fold-change in expression between III\textsuperscript{M} and YM males, as well as chromosomal location, to the network as metadata so this information could be visualized. To identify transcripts that may have more central functions within and across our significant modules, we ranked transcripts in descending order based on intramodular connectivity ($k_{\text{Within}}$), calculated in WGCNA. Hub genes identified by intramodular connectivity are generally functionally important transcripts within a module (Langfelder et al. 2013).

We further analyzed among-transcript connections involving a family of odorant-binding protein genes (Obp56h). Specifically, to identify transcripts that may regulate or be regulated by genes within the family, we calculated a “connection score” $C_i$ for every transcript $i$ as follows:

$$C_i = \sum_{j=1}^{n} a_{ij} |F_j|,$$

where $a_{ij}$ represents the adjacency (the Pearson correlation coefficient raised to the soft-threshold power $\beta$) between transcript $i$ and Obp56h gene $j$, and $F_j$ represents the log\(_2\) fold-change in expression between III\textsuperscript{M} and YM males for Obp56h gene $j$. This weighted product ensured that connections with Obp56h genes that are more differentially expressed between male types were prioritized in calculating a transcript’s connection score. Transcripts were then ranked by $C_i$ to identify candidate genes that may be strongly tied to Obp56h expression. Transcripts with the 100 highest Obp56h connection scores were classified as “central genes”. We tested for chromosomal enrichment among these central genes using Fisher’s exact tests (comparing the
number of central and non-central genes on a focal chromosome with the number of central and non-central genes on all other chromosomes) to determine whether the expression of \textit{Obp56h} genes (which are all located on the \textit{M. domestica} chromosome V) might be involved in \textit{trans} regulation with genes located on the III\textsuperscript{M} proto-Y chromosome.

\textit{Gene ontology enrichment analysis}

To identify gene ontology (GO) classes and molecular pathways that are enriched among DE transcripts, across co-expression modules identified in WGCNA, or among central genes co-expressed with \textit{Obp56h} genes, we used the BiNGO plug-in within Cytoscape (Maere et al. 2005). We identified \textit{D. melanogaster} orthologs for each house fly gene within a given gene list via NCBI blastx best hits (with default parameters) and used the \textit{D. melanogaster} gene name as input (Adhikari et al. 2021). We identified GO terms that are significantly enriched in BiNGO for biological processes, cellular components, and molecular function.

\textit{Allele-specific expression analysis}

We tested for differential expression of house fly chromosome III genes between the allele on the III\textsuperscript{M} chromosome and the allele on the standard third (III) chromosome in III\textsuperscript{M} males. To do so, we followed methods as in previous studies (Meisel et al. 2017; Son and Meisel 2021), which used the Genome Analysis Toolkit (GATK) best practices workflow for single nucleotide polymorphism (SNP) calling to identify sequence variants in our RNA-Seq data (McKenna et al. 2010). We focused our analysis on libraries that were sequenced from head tissue of male house flies that comprise a CS genetic background (Meisel et al. 2015; Son et al. 2019; Adhikari et al. 2021). We used STAR (Dobin et al. 2013) to align reads from a total of 30 head libraries (15 III\textsuperscript{M} and 15 YM libraries) to the house fly reference genome (Musca_domestica-2.0.2). We then followed the same methods and applied the same parameters as we have done previously to identify SNPs and genotype individual strains (Meisel et al. 2017; Son and Meisel 2021). We performed separate joint genotyping for each house fly strain within a given experiment (a total of 4 III\textsuperscript{M} and 4 YM strain-by-experimental batch combinations).

We use the following approach to differentiate between III\textsuperscript{M} and standard chromosome III alleles. We first identified SNPs in the exonic regions of the top “hub” genes within a WGCNA module that mapped to house fly chromosome III. We selected SNPs in those genes that are heterozygous in III\textsuperscript{M} males and homozygous in YM males. We used the genotype of these SNPs in YM males (which possess two standard third chromosome alleles) to determine the standard chromosome III allele. The allele not found in YM genotypes was assigned to the III\textsuperscript{M} chromosome. We also identified positions where III\textsuperscript{M} males appear monoallelic for an allele not found in YM males. These positions that exhibit a complete bias for a III\textsuperscript{M} allele are suggestive of monoallelic expression of the III\textsuperscript{M} allele (i.e., no expression from the III allele).
We tested for differences in expression of \( IIIM \) and standard chromosome III alleles by following best practices for comparing allele-specific expression (Castel et al. 2015). First, for each strain-by-experimental batch combination, we calculated the normalized read depth at each variable site as the number of mapped reads at that site divided by the total number of mapped reads throughout the genome. At each variable site, we used Wilcoxon rank sum tests to make three different pairwise comparisons per site. First, we compared normalized read depths between \( IIIM \) and III alleles in \( IIIM \) males (\( IIIM-III \)). Second, we compared the read depths of the \( IIIM \) allele in \( IIIM \) males with the normalized read depth of both III alleles in YM males (\( IIIM-YM \)). Third, we compared the read depths of the III allele in \( IIIM \) males with the normalized read depth of both III alleles in \( Y^M \) males (\( III-Y^M \)). We set a threshold of significance at \( p < 0.05 \) for all comparisons.

**Drosophila melanogaster RNA-seq and microarray data analysis**

We analyzed RNA-seq results reported in a previous study (Shorter et al. 2016) to determine how knockdown of \( Obp56h \) affects gene expression in \( D. melanogaster \). Shorter et al. (2016) identified DE genes between \( Obp56h \) knockdown and control samples. This analysis was done separately in males and females, and in separate tissue samples within a given sex (head or the remaining body). We conducted GO enrichment analysis, as described above, on the list of DE genes in \( D. melanogaster \) male head tissue upon \( Obp56h \) knockdown.

We tested if an excess of DE genes (between \( Obp56h \) knockdown and controls) are found on the \( D. melanogaster \) X chromosome, which is homologous to house fly chromosome III (Foster et al. 1981; Weller and Foster 1993). This chromosome is known as Muller element A across flies (Meisel and Scott 2018; Schaeffer 2018). \( Obp56h \) is located on \( D. melanogaster \) chromosome 2R (Muller element C), which is homologous to house fly chromosome V. We used Fisher’s exact tests (comparing the number of X and non-X chromosome genes that are DE in a given tissue within a given sex with the number of X and non-X chromosome genes that are not DE) to determine whether \( Obp56h \) knockdown in \( D. melanogaster \) results in the disproportionate differential expression of X chromosome genes in male heads, male bodies, female heads, or female bodies.

We also tested if the same genes are DE between \( IIIM \) vs \( Y^M \) house flies and \( Obp56h \) knockdown vs control \( D. melanogaster \). Using NCBI blastx best hits, we identified 20 \( M. domestica \) transcripts that are orthologous to \( D. melanogaster \) genes that are DE upon knockdown of \( Obp56h \) (11 matches to upregulated \( D. melanogaster \) genes, and 9 matches to downregulated \( D. melanogaster \) genes). This list represents 40% of the 50 \( D. melanogaster \) DE genes. We compared the mean log\(_2\) fold-changes between \( Y^M \) and \( IIIM \) house fly males for those 20 genes to 10,000 random subsets of log\(_2\) fold-change values taken from our data (10,000 subsamples without replacement of 11 genes to test for an excess of positive log\(_2\) fold-change values, and 10,000 subsamples of 9 genes to test for an excess of negative log\(_2\) fold-change values; see Additional Files for R script). We assessed significance by calculating the proportion of
replicated subsamples that generated a mean log2 fold-change value more extreme than our observed mean.

We additionally obtained gene expression measurements from microarray and RNA-seq data collected from various larval, pupal, and adult *D. melanogaster* tissue samples (Supplementary Table S2). Microarray data were sampled from FlyAtlas expression measurements of larval central nervous system, adult head, adult eye, and adult brain (Chintapalli et al. 2007). RNA-seq data were sampled from larval, pupal, or adult antennae (Shiao et al. 2013; Menuz et al. 2014; Pan et al. 2017; Mohapatra and Menuz 2019) and adult proboscis (May et al. 2019). We only included samples from wild-type flies, and we used the published expression estimates (e.g., microarray signal intensity, transcripts per million, reads per kilobase per million mapped reads) measured at either the gene or transcript level. When data from multiple replicates were available, we calculated the mean expression level across all replicates for each gene or transcript. The range and distributions of expression levels varied across tissue samples because the data were collected using different methodologies. To compare across tissue samples, we calculated a normalized expression level for each gene or transcript in each tissue sample by dividing by the mean value across all genes or transcripts in that tissue sample. Any gene or transcript with an expression value of 0, detected in <4 replicates of the microarray data, or that failed to pass threshold in an RNA-seq data set (i.e., status not OK) was excluded. We further extracted expression measurements from genes identified as hubs within the WGCNA co-expression network (see above). If a hub gene had multiple annotated transcripts, and expression was measured for transcripts, we calculated the gene expression level as the mean across all annotated transcripts.

We obtained measurements of the expression level of *CG2120* from FlyAtlas microarray and modENCODE RNA-seq data sampled at multiple developmental stages and across tissues (Chintapalli et al. 2007; Graveley et al. 2011; Brown et al. 2014). The FlyAtlas microarray data are the average signal intensity across all probes for the gene, and the modENCODE data are reported as reads per kilobase per million mapped reads (RPKM), both of which were obtained from FlyBase (Gelbart and Emmert 2013).

**Competitive courtship assays**

We performed competitive courtship experiments in which two different house fly males were combined with a single female, and we recorded the “winning” male (i.e., the one who mated with the female), similar to what was done previously (Hamm et al. 2009). In these experiments, we used the same two house fly strains as in Hamm et al. (2009): a III* strain called CS and a Y* strain called IsoCS. These two strains have a common genetic background (CS), and only differ in which proto-Y chromosome they carry. Both strains are represented in the RNA-seq data we analyzed (Meisel et al. 2015; Son et al. 2019), and IsoCS was also included in a previous RNA-seq study comparing the effects of proto-Y chromosome and temperature on gene
expression (Adhikari et al. 2021). Our experiment differed from previous work because we reared larvae from each strain at either 18°C and 29°C, whereas Hamm et al. (2009) worked with flies raised at 28°C. We used the same larval wheat bran diet as done previously, and we fed adults an *ad libitum* supply of water and an *ad libitum* 1:1 mixture of dry-milk:sugar. This is also the same diet and rearing protocol used for the flies in the RNA-seq datasets that we analyzed (Meisel et al. 2015; Son et al. 2019). Male flies were isolated from females within ~1 hour of eclosion, and each sex was kept separately to ensure that flies had not mated prior to the experiment.

We carried out two distinct competitive courtship experiments: 1) inter-strain competition between males with different genotypes (i.e., YM vs III^M^) that were reared at the same temperature (363 successful mating trials out of 490 total attempts across 27 experimental batches); and 2) intra-strain competition between males with the same genotype that were reared at different temperatures (104 successful mating trials out of 129 total attempts across 7 batches). When we competed flies with different genotypes raised at the same temperature, all males were aged 4-6 d post pupal emergence. When we compared flies with the same genotype raised at different temperatures, 29°C males were aged 4-5 d post emergence and 18°C males were aged 6-7 d post emergence. We aged flies from the colder temperature for more days than flies from the warmer temperature because developmental rate is positively correlated with developmental temperature in flies (Atkinson 1996). The ages we selected ensure that all males were physiologically capable of mating, while also sampling flies at similar physiological ages across experiments. Aging calculations are reported in Supporting Information 1.

The two males in each experiment were labeled using red and blue luminous powder (BioQuip) by shaking the flies in an 8 oz paper cup. The color assigned to males was switched in each successive batch (i.e., blue YM and red III^M^ in one batch, and then red YM and blue III^M^ in the next batch). In addition, we included the genotype or developmental temperature of the blue-colored male as a fixed effect in our statistical analysis (see below), which provides an additional control for color.

For each replicate of the competitive courtship assay, we placed the two different males in a 32 oz transparent plastic container, along with a single virgin female. Each plastic container also contained a 1:1 mixture of dry milk:sugar in a 1 oz paper cup and water in a glass scintillation vial plugged with a cotton roll. Virgin females from the LPR strain (Scott et al. 1996) raised at 25°C were used for all combinations of males. The LPR strain has a different genetic background than the males used in the assay, minimizing any effects of co-adaptation between females and a particular subset of males. All flies were transferred into the mating containers using an aspirator and without anesthesia. All matings were performed in a 25°C incubator because copulation latency is too long for experimentally tractable measurement at lower temperatures. The color (i.e., genotype) of the first male to mate was recorded, as well as the time to mate.
We used the glmer() function in the lme4 package in R (Bates et al. 2015) to test for the effects of genotype and temperature on male mating success. First, to test the effect of genotype, we constructed a logistic regression model as follows:

$$W \sim G_B + T + G_B \times T + b,$$

with developmental temperature ($T$), genotype of the blue male ($G_B$), and their interaction as fixed effects. Experimental batch ($b$) was modeled as a random effect, with the winning male ($W$: CS or IsoCS) as a response variable. We then assessed significance of fixed effects (type II sum of squares) using the Anova() function in the car package in R (Fox et al. 2013). To test for the effect of temperature on mating success, we similarly constructed a logistic regression model as follows:

$$W \sim G + T_B + G \times T_B + b,$$

with genotype ($G$), developmental temperature of the blue male ($T_B$), and their interaction as fixed effects, experimental batch ($b$) as a random effect, and the winning male ($W$: 18°C or 29°C) as a response variable. We then assessed significance of fixed effects (type II sum of squares) using the Anova() function in the car package in R (Fox et al. 2013).

**Single-choice courtship assays**

We performed experiments to measure copulation latency, or the amount of time elapsed before mating, according to male type ($Y^M$ or $IIIM^M$). In these experiments, we used the same IsoCS ($Y^M$) and CS ($IIIM^M$) strains as above and in Hamm et al. (2009). We also tested one other strain from each genotype. CSrab ($III^M$) was created by backcrossing the $III^M$ chromosome from the rspin strain isolated in New York onto the CS background (Shono and Scott 2003; Son et al. 2019). CSaY ($Y^M$) was created by backcrossing the $Y^M$ chromosome from the aabys genome reference strain onto the CS background (Scott et al. 2014; Meisel et al. 2015). Virgin females used in the assays were all from the LPR strain (Scott et al. 1996), which has a different genetic background than all males tested. In addition, we also assayed LPR males to determine how copulation latencies of $IIIM^M$ and $Y^M$ males compare to those of males from the same genetic background as the females.

We first attempted to test flies reared at the same temperatures as in our competitive courtship assays (18°C and 29°C), as well as at an intermediate developmental temperature (22°C). However, we did not generate enough flies at 18°C, and so we only have data for flies raised at 22°C and 29°C. Our results demonstrate that 22°C is a sufficiently low temperature to detect effects of both genotype and developmental temperature on courtship success (see below). All larvae from each male strain were reared in 32 oz plastic containers on the same wheat bran diet described above (Hamm et al. 2009). Upon emergence, unmated male and female progeny were separated and fed water and a 1:1 mixture of dry milk:sugar *ad libitum* until assays were...
conducted. Assays of males raised at 22°C were conducted 10-11 days after eclosion, while those of males raised at 29°C were conducted 6-7 days after eclosion. This ensures that males were assayed at similar physiological ages. Females were all raised at 25°C, and unmated females were aged 8-9 days after eclosion (see Supporting Information 1 for all accumulated degree day calculations).

We followed a similar protocol as in a previous experiment testing copulation latency in *D. melanogaster* (Shorter et al. 2016). Briefly, five males from a single strain were aspirated without anesthesia into an 8 oz container covered with a fine mesh cloth secured by rubber band. Five LPR females were similarly transferred into the container, marking the start of the courtship assay. The house flies were then observed every 10 minutes over the course of four hours.

Copulation latency was determined in two ways. First, we measured the amount of time elapsed between the start of an assay and each mating within a container, defined as a male remaining attached to a female for at least 1 minute (Hamm et al. 2009). Male house flies typically remain attached to females for >60 minutes (Bryant 1980), making it unlikely, although possible, for us to miss matings within 10 minute intervals. Individuals who did not mate were excluded from this analysis. Second, we used a binary variable noting whether each male mated during the 4 hour assay. Although we were unable to distinguish between individual males in this assay, we did not observe any males mate more than once within 4 hours in a pilot study conducted between one male and five females, suggesting that observed matings were by different males.

All trials were conducted at 22–23°C.

To determine the effects of male type on the amount of time taken to mate, we used the glmer() function in the lme4 package in R (Bates et al. 2015) to create a mixed effects model as follows:

\[
L \sim G + T + G \times T + b + s,
\]

including male genotype (G), developmental temperature (T), and their interaction as fixed effects, batch (b) and strain (s) as random effects, and our response variable as copulation latency (L) in minutes. For the binary measure of copulation latency, we used a binomial logistic regression of the same model, with whether a fly mated as our dependent variable. We then assessed significance of fixed effects (type II sum of squares) using the Anova() function in the car package in R (Fox et al. 2013). Pairwise comparisons between male types (II, YM, and LPR) were conducted using Z-tests of proportions.

Lastly, we conducted a survival analysis, with unmated males treated as right-censored observations since information about their mating latency was incomplete. Specifically, we used the Surv() function in the survival package in R (Fox and Carvalho 2012) to convert our copulation latency measurements (in minutes) to a right-censored survival object. We then created a Cox proportional hazards regression model using the same parameters as above, with copulation latency (in minutes) as our response variable. Due to limitations of the survival
package, we included batch and strain as fixed effects in our model. We assessed significance of fixed effects using analysis of deviance for a Cox model using the anova() function in the survival package, and we report hazard ratios (HRs) and their 95% confidence intervals for copulation latency effect sizes (Burke and Holwell 2021).

**Assaying copulation latency in CG2120 upregulated D. melanogaster**

We used CRISPR/dCas9 transcriptional activation (CRISPRa) to increase the expression of CG2120 in *D. melanogaster*. We used a strain that expresses single guide RNA (sgRNA) targeting CG2120 (BDSC ID 79962), from the Transgenic RNAi Project CRISPR Overexpression (TRiP-OE) VPR collection (Ewen-Campen et al. 2017). We crossed males with the sgRNA to females carrying a transgene encoding a deactivated Cas9 protein (dCas9) expressed under one of two Gal4 drivers (Gal4>dCas9). One Gal4>dCas9 strain expresses Gal4 under regulation of the *apolpp* promoter (BDSC ID 67043), which expresses in the fat body (Brankatschk and Eaton 2010; Van De Bor et al. 2015). The other Gal4>dCas9 strain regulates Gal4 under the *elav* promoter (BDSC ID 67038), which expresses in neurons (Sink et al. 2001; Zhang et al. 2002). We collected male progeny from the cross between the sgRNA strain and each of the Gal4>dCas9 strains to assay copulation latency. Control males were generated by substituting the CG2120 sgRNA strain with a strain carrying an sgRNA that targets the QUAS sequence from *Neurospora crassa* (BDSC stock 67539). All flies were raised at 25°C on a medium consisting of cornmeal, yeast, sugar, and agar.

We measured copulation latency in CG2120 upregulated males and control males. In each experiment, we used a single unmated male from one of the crosses described above. That male was combined with a single unmated female from either the CantonS or OregonR strain in a vial containing our standard medium. The flies were observed for up to 2.5 h at 25°C, and the time at which they began to copulate was recorded as our measurement of copulation latency. Fly pairs that did not copulate were excluded from the analysis. Experiments were performed in four batches for the *apolpp* driver and five batches for the *elav* driver.

We compared the fit of linear models in order to test for differences in copulation latency between CG2120 upregulated males and control males. We first constructed a model that captured all variables in our experiment,

\[
C \sim M + F + M \times F + b,
\]

which included the fixed effects of male genotype (M, which can be control or CG2120 upregulated), female strain (F, for CantonS or OregonR), and the interaction between male genotype and female strain, as well as the random effect of experimental block (b). We used the lmer() function in the lme4 package in R (Bates et al. 2015) to determine how each variable affects copulation latency (C). We compared that full model with one excluding the interaction term with a \( \chi^2 \) test in the anova() function in R (R Core Team 2019) in order to test if including
the interaction term offers a significantly better fit. If the interaction term did not significantly improve the fit of the model, we compared the model without the interaction term with one without any effect of the female strain (i.e., F excluded from the model). If there was not a significant difference in fit between the models with and without F, we modeled female strain as a random effect. To test for an effect of CG2120 upregulation on copulation latency, we compared the fit of the model with male genotype as the only fixed effect and two random effects (f and b) with a model that only had the random effects (i.e., no fixed effects) using the anova() function.

RESULTS

Differential expression of odorant binding protein genes between III^M and Y^M males

Our first goal was to analyze RNA-seq data in order to identify DE genes between III^M and Y^M male heads that could be responsible for previously observed differences in competitive courtship assays between the genotypes (Hamm et al. 2009). We confirmed that the gene expression profiles of III^M and Y^M male heads are minimally differentiated (Meisel et al. 2015; Son et al. 2019). There are only 40 DE genes between heads of III^M and Y^M adult males (21 upregulated in Y^M males, 19 upregulated in III^M, Supplementary Table S3). Gene ontology analysis revealed no significant biological process, molecular function, or cellular component terms enriched within the list of DE genes.

Within the list of DE genes, we identified one gene (LOC105261916) encoding an odorant binding protein (Obp) upregulated in Y^M males. House fly Obp genes can be grouped into families corresponding to their D. melanogaster orthologs (Scott et al. 2014). The DE Obp gene in our analysis is orthologous to Obp56h. The Obp56h family, as well as other Obp families, was greatly expanded within Muscidae (house fly and close relatives, including stable fly and horn fly) compared to D. melanogaster (Scott et al. 2014; Olafson and Saski 2020; Olafson et al. 2021). In addition to LOC105261916, seven of the remaining eight house fly Obp56h genes for which we obtained RNA-seq count data showed similar trends of greater expression in Y^M than III^M males, with three of these showing significant DE (p<0.05) before an FDR correction (Fig. 1). All but one of the Obp56h genes has higher expression in Y^M than III^M males (8/9, regardless of significance), which is significantly greater than the fraction of other genes with higher expression in Y^M males, regardless of significance, in the rest of the genome (Fisher’s exact test, p = 0.019).
Figure 1 - Neighbor-joining phylogenetic tree of the Obp56h gene family within M. domestica and D. melanogaster based on protein sequences constructed in MEGA X (Kumar et al. 2018). Amino acid sequences were aligned by MUSCLE (Edgar 2004). M. domestica Obp56h genes are identified based on gene IDs. The bootstrap consensus tree was inferred from 10,000 replicates. Branch lengths are scaled according to the number of amino acid substitutions per site. The phylogeny was arbitrarily rooted at D. melanogaster Obp56h. Graphs at the branch tips show batch-adjusted expression levels for each M. domestica Obp56h gene from each replicate (small circles). Large circles show the average across all replicates, with error bars denoting the standard error (unfilled stars: p < 0.05 before FDR correction for multiple comparisons; filled star: p < 0.05 after correction).

**III M confers a courtship advantage by reducing copulation latency**

Knockdown of Obp56h in D. melanogaster decreases male copulation latency, or the time it takes for a male to begin to mate with a female after they are first exposed to one another (Shorter et al. 2016). The Obp56h gene family is generally expressed higher in YM males relative to III M males (Fig. 1). A previous study identified a competitive advantage of III M over YM male house flies in successfully engaging a female in mating (Hamm et al. 2009), consistent with shorter copulation latency in III M males because of lower expression of Obp56h genes. We confirmed this result by performing competitive courtship assays in which we allowed males carrying III M or YM to compete for a female of an unrelated strain. Consistent with the previous results, we observed that III M males are more successful at mating than YM males when raised at 29°C (Fig. 2A), a temperature similar to the developmental temperature used by Hamm et al. (2009).

Two Obp56h genes are only upregulated in YM males at 29°C, but not at 18°C (Fig. S1), raising the possibility that the effect of III M/YM genotype on male courtship success may be temperature-dependent. In addition, III M males have greater heat tolerance than YM males (Delclos et al. 2021), further suggesting that the benefits of the III M chromosome may be limited to warm temperatures. We therefore tested if the differences in courtship performance between YM and III M males are sensitive to temperature. We found that III M males were more successful...
at mating than $Y^M$ males regardless of developmental temperature (ANOVA, $X^2 = 20.7, p = 6.53 \times 10^{-6}$, Fig. 2A). We then tested whether males reared at different developmental temperatures, but with the same genotype, have a difference in courtship success. We found that males reared at 18°C outcompete males reared at 29°C, regardless of genotype (ANOVA, $X^2 = 13.1, p = 2.93 \times 10^{-4}$, Fig. 2B). This demonstrates that there is an effect of developmental temperature on courtship success, but it appears to be independent of the genotype effect. More generally, our results suggest that the $III^M$ male courtship advantage is robust to developmental temperature, which differs from the prediction based on temperature-dependent $Obp56h$ expression (Fig. S1).

![Figure 2](image)

**Figure 2** - $III^M$ chromosome and developmental temperature affect male courtship success. A) Outcomes of competitive courtship assays between $III^M$ and $Y^M$ males reared at 18°C or 29°C. Data points represent experimental batches. Horizontal lines denote the median across all batches. B) Outcomes of competitive courtship assays conducted between males reared at 29°C and 18°C. Trials were conducted between males of the same proto-Y chromosome genotype ($III^M$ or $Y^M$). Each data point represents ten replicate trials within a single batch. C) Outcomes of single-choice courtship assays in males reared at 22°C. Data points refer to the percentage of males (five males within one replicate) that mated with females within 4 hours within each experimental trial. Horizontal lines denote means within male groups. All females used were from the LPR strain.

We next tested if the effects of developmental temperature and genotype on competitive mating advantage could be caused by differences in copulation latency. To measure copulation latency, we combined five males from a single strain raised at a single temperature with five females from the unrelated strain used in our competitive courtship assays. Developmental temperature had a significant effect on copulation latency (ANOVA, $X^2 = 15.3, p = 9.40 \times 10^{-5}$), with males reared at 22°C mating faster than those reared at 29°C (Fig. S3A). This result is consistent with increased courtship success of males raised at 18°C relative to those raised at 29°C in our competitive assays (Fig. 2B).

We conducted a survival analysis to test for differences in copulation latency between $Y^M$ and $III^M$. To those ends, we fit a Cox proportional hazards regression model, treating unmated males
as censored data (Burke and Holwell 2021), and we observed significant effects of both
developmental temperature (ANOVA, $X^2 = 22.5, p = 2.16 \times 10^{-6}$) and male genotype ($X^2 = 18.0,$
$p = 1.22 \times 10^{-4}$) on copulation latency. However, there was no significant interaction effect
between genotype and developmental temperature ($X^2 = 2.63, p = 0.277$). Consistent with our
hypothesis that $\text{IIIM}$ males have a reduced copulation latency relative to $\text{YM}$ males, at $22^\circ C$, $\text{IIIM}$
males mated 3.5 times faster than $\text{YM}$ males (Cox model: HR = 3.50, CI = 0.95 to 12.9, Fig.
S2A). We do not observe this trend at $29^\circ C$ (Fig. S2B), possibly due to very few successful
matings at the warmer developmental temperature.

To further address the limitations of censored data, we treated copulation latency as a binary
variable by calculating the proportion of the five males per trial that mated within the 4 hour
assay. We observed significant effects of male genotype (ANOVA, $X^2 = 10.2, p = 6.18 \times 10^{-3}$)
and developmental temperature (ANOVA, $X^2 = 11.0, p = 9.04 \times 10^{-4}$) on the proportion of males
that mated. The effect of developmental temperature was largely a result of very few matings for
males that developed at $29^\circ C$ relative to $22^\circ C$ (Fig. S3B), consistent with our finding that males
that developed at a lower temperature have higher courtship success (Fig. 2B). In the $22^\circ C$
treatment, a significantly greater proportion of $\text{IIIM}$ males mated within 4 hours than $\text{YM}$ (61.7% v.
28.3%; Z-test of proportions, $p = 1.21 \times 10^{-4}$; Fig. 2C). This is further evidence that $\text{IIIM}$ males
have reduced copulation latency, which is consistent with their previously documented
competitive advantage (Hamm et al. 2009), the competitive advantage that we observe (Fig. 2A),
and the reduced expression of $\text{Obp56h}$ genes (Fig. 1).

The $\text{YM}$ and $\text{IIIM}$ males we and others used in courtship experiments all share the CS genetic
background that comes from a $\text{IIIM}$ strain (Hamm et al. 2009). This raises the possibility that $\text{IIIM}$
males perform better because they have a proto-Y chromosome that is co-adapted to its genetic
background. To test this hypothesis, we measured copulation latency in $\text{YM}$ males from the same
strain (LPR) as the females in our experiments. We observed a greater proportion of $\text{IIIM}$ males
mating within 4 hours when compared to the LPR $\text{YM}$ males (Z-test of proportions, $p = 0.038$),
although the copulation latency in LPR males was highly variable (Fig. 2C). Therefore, the
reduced copulation latency conferred by the $\text{IIIM}$ chromosome overwhelms any potential effects
of coadaptation of the proto-Y chromosome to male genetic background or male-female co-
adaptation within strains. The reduced copulation latency of $\text{IIIM}$ males is only detectable when
house flies develop at $22^\circ C$, suggesting that it is either temperature-dependent or we lack the
resolution to detect it when males develop at warmer temperatures (because they take too long to
mate).

**House fly chromosome III genes and Drosophila X chromosome genes have correlated
expression with $\text{Obp56h}$ genes**

The $\text{Obp56h}$ gene cluster is found on house fly chromosome V, suggesting that it is regulated in
trans by genes on the $\text{YM}$ or $\text{IIIM}$ chromosome. Chromosome V is unlikely to differ between the
YM and III\textsuperscript{M} males in our experiments—the majority of males compared in the RNA-seq data and mating experiments have a common genetic background (including chromosome V) and differ only in whether they carry III\textsuperscript{M} or YM. Removing samples with a different genetic background did not affect the general difference in \textit{Obp56h} expression between III\textsuperscript{M} and YM males (see Supporting Information 2 for a summary of these results). We therefore aimed to identify genes on the YM or III\textsuperscript{M} chromosome that could be responsible for the differential expression of \textit{Obp56h} genes between YM and III\textsuperscript{M} males.

We first identified 27 co-expression modules across YM and III\textsuperscript{M} house fly male heads. We focus on one of these modules (containing 122 genes, Supplementary Table S4) because it is differentially expressed between YM and III\textsuperscript{M} males ($p_{\text{ADJ}} = 0.001$), and it contains three \textit{Obp56h} genes that are DE between III\textsuperscript{M} and YM males (LOC105261916, LOC101891822, and LOC101891651) (Fig. 3). GO analysis revealed significant enrichment ($p_{\text{ADJ}} < 0.05$) of 15 biological process terms including those related to immune system processes (GO:0032501), responses to stress (GO:0006950), and response to external stimuli (GO:0009605) within this module (Supplementary Table S5). We used the WGCNA measure of intramodular connectivity, kWithin, to identify hub genes within the module that likely have important roles in the regulation of gene expression. The top five hub genes are (with \textit{D. melanogaster} orthologs in parentheses): LOC101887703 (CG8745), LOC105262120 (CG10514), LOC101894501 (Md-Gr35), LOC101893264 (gd), and LOC101893651 (CG2120) (Fig. 3, S4).

**Figure 3** - Network visualization of the co-expression module that is differentially regulated between III\textsuperscript{M} and YM\textsuperscript{M} males. Each circle within the module is a gene, and \textit{Obp56h} genes are indicated with purple fill. Lines represent edge connections between genes. Genes labeled with “+” are within the top 100 most strongly connected to \textit{Obp56h} genes. Genes are ordered from top to bottom according to intramodular connectivity (kWithin), with genes of higher connectivity (i.e., hub genes) on top, and peripheral genes on the bottom. Borders around genes denote log\textsubscript{2} fold-change in expression between YM\textsuperscript{M} and III\textsuperscript{M} male.
heads, with darker blue borders denoting upregulation in Y^M, and darker red borders denoting upregulation in IIIM. Chromosomal locations in house fly (Mdom) and D. melanogaster (Dmel) are shown for the 5 hub genes and Obp56h.

The genes present in the focal co-expression module provide multiple lines of evidence that Obp56h gene expression is regulated by trans factors that map to chromosome III. First, the module is enriched for house fly chromosome III genes (31 chromosome III genes versus 38 genes assigned to other chromosomes, Fisher’s exact test \( p < 1 \times 10^{-5} \), with 53 genes not assigned to a chromosome) and for DE genes between Y^M and IIIM males (16 DE genes in this module versus 24 DE genes assigned to other modules, Fisher’s exact test \( p < 1 \times 10^{-5} \)). In addition, the co-expression module is enriched for Obp56h genes relative to other Obp genes—three Obp56h genes and no other Obp genes were assigned to this module (Fisher’s exact test, \( p = 5.1 \times 10^{-3} \), Fig. 3). Furthermore, there is a significant enrichment of chromosome III genes within the 100 genes whose expression covaries most with Obp56h gene expression (corresponding to the top 0.55% covarying genes; Supplementary Table S6); of the 100 genes with the highest Obp56h connection scores, 26 are on chromosome III (Fisher’s exact test \( p = 2.0 \times 10^{-4} \), Fig. 4A). This enrichment is robust to varying the threshold used to classify a gene as in the top covarying; considering genes with the top 1%, 5%, or 10% covarying expression also resulted in significant enrichment of chromosome III genes (Fisher’s exact test, all \( p < 0.05 \)). These results support the hypothesis that trans regulatory variants that differ between III^M and the standard chromosome III are responsible for the differential expression of Obp56h genes between IIIM and Y^M house fly males. We cannot perform the same analysis for the effect of Y^M because only 51 genes have been assigned to the house fly X/Y^M chromosome (Meisel and Scott 2018), limiting our power to detect an excess of genes.

**Figure 4** - Percent of genes on each chromosome within (A) the top 100 genes with the strongest correlated co-expression to the Obp56h family in house fly, and (B) genes differentially expressed (DE) between Obp56h knockdown and control D. melanogaster (black bars: males, grey bars: females). Asterisks indicate a significant difference between observed (bars) and expected (red lines) counts of genes on each chromosome compared to all other chromosomes (Fisher’s exact test, \( p < 0.05 \)).
Our network analysis does not ascribe directions to the edges connecting house fly genes, and it is therefore possible that Obp56h has trans regulatory effects on chromosome III expression. To test this hypothesis, we examined available RNA-seq data from an experiment comparing wild-type D. melanogaster with flies in which Obp56h had been knocked down (Shorter et al. 2016). Obp56h is on the right arm of the second chromosome in D. melanogaster (2R, or Muller element C), which is homologous to house fly chromosome V (Foster et al. 1981; Weller and Foster 1993). House fly chromosome III is homologous to the D. melanogaster X chromosome, which is known as Muller element A (Meisel and Scott 2018; Schaeffer 2018). The D. melanogaster males in the RNA-seq experiment all share the same X chromosome, and only differ in one copy of their second chromosome (which either carries a UAS-RNAi knockdown construct or does not). If Obp56h genes have trans regulatory effects on element A genes in males, we would expect an excess of DE D. melanogaster X chromosome genes in Obp56h knockdown flies. Indeed, we found that Obp56h knockdown in D. melanogaster resulted in an excess of X chromosome DE genes in male head (Fisher’s exact test, p = 0.011, Fig. 4B) and body (p = 0.038, Fig. S5), but not in either tissue sample in females (Fisher’s exact test, both p > 0.49). These results suggest that there is male-specific trans regulatory control of D. melanogaster X-linked genes by Obp56h.

We found multiple similarities between house fly and D. melanogaster that suggest the genes that regulate and/or are regulated by Obp56h are evolutionarily conserved between the two species. Specifically, genes that were downregulated upon knockdown of Obp56h in D. melanogaster have house fly orthologs that are more downregulated in III M male house flies (i.e., lower log2 fold-change) than expected by chance (p = 5.60 x 10^-3, Fig. S6A). However, genes that were upregulated upon Obp56h knockdown in D. melanogaster were not significantly differentially regulated between Y M and III M male genotypes, although the observed trend suggests that these genes may be more downregulated in III M males than expected (p = 0.103, Fig. S6B). The GO term “response to stress” (GO:0033554) is significantly enriched amongst genes with strong connection scores with Obp56h expression in M. domestica and in the list of DE genes in D. melanogaster upon Obp56h knockdown (Supplementary Table S7), providing additional evidence for conserved co-regulation. Altogether, our results suggest that there is evolutionarily conserved trans regulatory feedback loop involving Obp56h expression and Muller element A in Drosophila and house fly through similar molecular functions.

**Network analysis reveals candidate regulators of Obp56h expression**

The house fly co-expression module contains candidate genes and pathways through which Obp56h genes, and likely male copulation latency, are regulated. For example, a serine protease gene, LOC101893264, orthologous to D. melanogaster gd (Konrad et al. 1998), is among the top 5 hub genes within the co-expression module (Fig. 3). This gene is predicted to encode a positive
regulator of the Toll signaling pathway (LeMosy et al. 2001; Valanne et al. 2011), suggesting that the *M. domestica* ortholog of *gd* (*Md-gd*) could have an important gene regulatory function within the module via Toll signaling. *Md-gd* is located on chromosome III, and it is upregulated in **III** males (adj. *p* = 0.022). We tested if *Md-gd* is differentially regulated between the **III** chromosome and standard chromosome III by comparing expression in **III** males (i.e., heterozygotes for **III** and a standard chromosome III) with **Y** males that are homozygous for the standard chromosome III. Differential expression of the **III** and **III** chromosome alleles would implicate *Md-gd* as having a causal effect on *Obp56h* expression. We identified seven polymorphic sites where all RNA-seq reads were mapped to the **III** allele, while no reads were mapped to the standard chromosome III allele in **III** males (Fig. 5A). At all seven diagnostic SNPs in this gene, the **III** allele is significantly more highly expressed than the **III** allele in **III** males (all *p* = 0.021), and it is more highly expressed than both **III** alleles in **Y** males at six of seven sites (all *p* = 0.021). Higher expression of the **III** allele is consistent with cis regulatory divergence between the **III** and standard chromosome III being responsible for elevated *Md-gd* expression in **III** males. The lack of expression of the **III** allele in either **III** or **Y** males is consistent with monoallelic gene expression of the **III** allele, although further evidence is required to confirm this hypothesis (see Supporting Information 3 for detailed discussion).

**Figure 5** - Allele-specific expression (ASE) in A) *LOC101893264* (*Md-gd*), B) *LOC101893651* (*CG2120* ortholog), and C) *LOC101894501* (*Md-Gr35*). The x-axis depicts base pair positions (scaffold coordinates) of the informative single nucleotide polymorphisms (SNPs) that differ between **III** and standard chromosome III alleles. The y-axis and data points depict the read depth of a given allele normalized by the total mapped reads for a given strain-by-experimental batch group combination (FPM = fragments per million). Lines depict mean read depths at each diagnostic site for **III** (turquoise) and **III** (salmon) alleles in **III** males, and mean read depths at each site for **III** alleles in **Y** males (black). Tables under each graph mark significant differences (*: *p* < 0.05) in normalized read depths at each diagnostic site for each of three pairwise comparisons: **III** allele vs. **III** allele in **III** males (**III**-**III**), **III** allele in **III** males vs both **III** alleles in **Y** males (**III**-**Y**), **III** allele in **III** males vs. both **III** alleles in **Y** males (**III**-**Y**).
We identified similar evidence of monoallelic gene expression within another hub gene, \( \text{LOC101893651} \), which is orthologous to \( D. \ melanogaster \ CG2120 \) (Fig. 5B). \( \text{LOC101893651} \) is among the most central genes within the co-expression module (Fig. 3), and it is strongly upregulated in III\(^M\) males (log\(_2\) fold-change: 1.33, \( p_{\text{ADI}} = 0.033 \)). \( \text{LOC101893651} \) is found on house fly chromosome III and is predicted to encode a transcription factor. At all four diagnostic sites within \( \text{LOC101893651} \), the III\(^M\) allele is significantly more highly expressed than the III allele in III\(^M\) males (all \( p = 0.021 \)), as well as both III alleles in Y\(^M\) males (all \( p \leq 0.027 \)). Within the WGCNA module, \( \text{Obp56h} \) expression is most strongly correlated with \( \text{LOC101893651} \), suggesting that \( \text{LOC101893651} \) could encode the transcription factor that is directly responsible for the repression of \( \text{Obp56h} \) expression in III\(^M\) males. Consistent with this hypothesis, there is evidence that the protein encoded by \( \text{CG2120} \) binds both upstream and downstream of \( \text{Obp56h} \) in \( D. \ melanogaster \) (Kudron et al. 2018). Below, we describe experimental results that test the hypothesis that \( \text{LOC101893651} \) (CG2120) regulates the expression of \( \text{Obp56h} \) genes.

There is also a gustatory receptor gene (\( \text{LOC101894501} \)) that is a hub in the co-expression module and upregulated in III\(^M\) males (\( p_{\text{ADI}} = 0.037 \)). \( \text{LOC101894501} \) (also annotated as \( \text{Md-Gr35} \)) is a homolog of \( D. \ melanogaster \) Gr98c (Scott et al. 2014). We evaluated the relationship between \( \text{Obp56h} \) and the hub genes in the co-expression network module, and we found greater evidence that this module and \( \text{Obp56h} \) plays an important role in chemosensation (Supporting Information 4, Fig. S7). \( \text{Md-Gr35} \) contains 4 exonic SNPs differentiating the III\(^M\) and III chromosomes. Within each III\(^M\) strain in each RNA-seq experiment, we observed significantly greater expression of the III\(^M\) allele than the standard chromosome III allele at two of the four diagnostic SNP sites (Fig. 5C). The other two SNPs showed the same pattern of III\(^M\)-biased expression but were not significant (both \( p > 0.05 \)). The III\(^M\) allele in III\(^M\) males is also expressed higher than both III alleles in Y\(^M\) males, consistent with \( \text{cis} \) regulatory divergence between the III\(^M\) and standard chromosome III driving elevated \( \text{Md-Gr35} \) expression in III\(^M\) males. However, the standard chromosome III allele is expressed significantly higher in III\(^M\) males than Y\(^M\) males at two of the four diagnostic SNP sites (Fig. 5C); we observe the same pattern at the other two sites without significance (\( p > 0.05 \)). Higher expression of the III allele in III\(^M\) males than Y\(^M\) males suggests that \( \text{trans} \) regulators further increase the expression of \( \text{Md-Gr35} \) in III\(^M\) males. This combination of \( \text{cis} \) and \( \text{trans} \) regulatory effects on \( \text{Md-Gr35} \) expression are consistent with the \( \text{trans} \)-regulatory loop we hypothesize between \( \text{Obp56h} \) and chromosome III that regulates male copulation latency. Future experiments could determine whether Gr98c (Md-Gr35) and \( \text{Obp56h} \) do indeed interact and, if so, what pheromonal or other chemical compounds they detect.

In contrast to the aforementioned three hub genes, the most central gene within the module (\( \text{LOC101887703} \)) may be regulated by \( \text{Obp56h} \). \( \text{LOC101887703} \) is orthologous to \( D. \ melanogaster \) CG8745, which is predicted to encode an ethanolamine-phosphate phospho-lyase and is broadly expressed in many \( D. \ melanogaster \) tissues (Chintapalli et al. 2007). \( \text{LOC101887703} \) is upregulated in III\(^M\) males (log\(_2\) fold-change: 2.21, \( p_{\text{ADI}} = 0.016 \)), which could
have a causal effect on Obp56h DE, be caused by Obp56h DE, or neither. In both
*D. melanogaster* and house fly, Obp56h expression is significantly negatively correlated with the
expression of CG8745 or LOC101887703, respectively (Fig. 6). Directly comparing CG8745
expression between control and Obp56h-knockdown *D. melanogaster* yields qualitatively similar
results, with Obp56h-knockdown *D. melanogaster* flies showing greater expression of CG8745 than controls
(Welch’s *t*-test, *t* = -4.27, *p* = 5.53 x 10^-3). The negative correlation between CG8745 and
Obp56h expression in *D. melanogaster* suggests that Obp56h downregulation causes CG8745
upregulation because Obp56h expression was directly manipulated by RNAi in the experiment.
The same relationship between CG8745 and Obp56h expression in both house fly and
*D. melanogaster* provides evidence that similar molecular mechanisms may underlie the
hypothesized trans regulatory effect of Obp56h in both species.

![Figure 6](image.png)

**Figure 6** - Correlations of gene expression between Obp56h (house fly LOC105261916) and CG8745
(LOC101887703) in (A) house fly male head tissue, and (B) *D. melanogaster* male head tissue. Values
for *D. melanogaster* are from count data as reported in Shorter et al. (2016). Linear regression models
were used to determine 95% confidence intervals (shaded in grey) summarizing the effect of Obp56h
expression on CG8745 expression in each species.

**CG2120 expression affects courtship**

We investigated the expression of CG2120, one of the hub genes in the house fly co-expression
network, across *D. melanogaster* development and tissues. Expression of CG2120 oscillates over
the life history of *D. melanogaster*: increasing throughout embryonic development, decreasing
with each subsequent larval instar, and then increasing in pupa and in some adult tissues (Fig
7A-C). CG2120 is predominantly expressed in digestive tissues in adults, with much lower
expression in nervous tissues (Fig 7B-C). Notably, CG2120 is more highly expressed in male
than female heads (Fig 7B), consistent with a regulatory effect on male courtship behavior.

We tested the hypothesis that CG2120 affects copulation latency in *D. melanogaster* via negative
regulation of Obp56h expression. To that end, we used CRISPRa to upregulate expression of
CG2120 in fat body cells or neurons of *D. melanogaster* males, using an *apolpp* and *elav*
We tested if upregulation of \textit{CG2120} reduces copulation latency, as
might be expected if \textit{CG2120} negatively regulates \textit{Obp56h}. We used females from two different
strains in our assays (CantonS and OregonR). Neither female strain nor the interaction between
female strain and male genotype had a significant effect on copulation latency, regardless of the
tissues in which expression was induced (Supplementary Table S8). There was also not a
significant difference in copulation latency between control and \textit{CG2120} upregulated flies when
we activated expression in fat body cells (Supplementary Table S8, Fig. 7D). In contrast, there
was a significant difference between control and \textit{CG2120} upregulated males when expression
was activated in neurons using the \textit{elav} driver, with \textit{CG2120} upregulated males mating faster
than controls (Supplementary Table S8, Fig. 7D). This is consistent with \textit{CG2120} activation in
neurons negatively regulating \textit{Obp56h}, which then reduces copulation latency.

\textbf{Figure 7.} \textit{CG2120} is broadly expressed and affects copulation latency. Expression levels of \textit{CG2120} are
plotted using (A) RNA-seq data from developmental stages (Graveley et al. 2011), (B) RNA-seq data
from adult tissues (Brown et al. 2014), or (C) microarray data from adult tissues (Chintapalli et al. 2007).
Expression measurements are shown as RPKM values for RNA-seq data and signal intensity for
microarray data (Gelbart and Emmert 2013). Bars are colored based on tissue sample categories: embryo
(white), larva (light gray), pupa (dark gray), adult female (magenta), adult male (light blue), neuronal
(black), digestive tract (green), or other adult tissues (orange). (D) Copulation latency in control male
\textit{D. melanogaster} and males in which \textit{CG2120} was upregulated (\textit{CG2120 up}) in fat body cells (using an
aporpp Gal4 driver) or neurons (elav driver). Each dot is a single male for which copulation latency was
measured. Vertical bars show the estimated effect of male genotype (control or upregulated) from a linear
model in which male genotype is a fixed effect and female strain and batch are random effects. The
asterisk indicates a significant difference ($P<0.05$) between control and \textit{CG2120} upregulated for a
specific tissue driver.
Figure 8 - Hypothesized relationships between $Obp56h$ expression, proto-Y chromosome genotype, and male mating behavior based on house fly and $D.\ melanogaster$ gene expression data. A) Summary of evidence for an evolutionarily conserved $trans$ regulatory loop between $Obp56h$ and Muller Element A (house fly chromosome III, and $D.\ melanogaster$ X chromosome). Our hypothesis is based on differential expression between III$^M$ vs. Y$^M$ male house flies, $Obp56h$ knockdown vs. control $D.\ melanogaster$, and network connectivity of $Obp56h$ family gene expression within house fly. B) Summary of candidate genes implicated in conserved $trans$ regulatory loop. Three of the top five hub genes of module A are located on house fly chromosome III, are negatively correlated with $Obp56h$ expression, and exhibit either allele-specific expression (ASE) or show signs of monoallelic gene expression biased towards the III$^M$ allele. Similar correlations between expression measures of $Obp56h$ and CG8745 (LOC101887703) in $D.\ melanogaster$ and house fly male head tissue suggest that $Obp56h$ regulates CG8745, which is the primary hub gene in the WGCNA module that is differentially expressed between III$^M$ and Y$^M$ male house flies. Shared correlations between $Obp56h$ expression and copulation latency in both house fly and $D.\ melanogaster$ also suggest that $Obp56h$ regulates male fly mating behavior.

DISCUSSION

We combined analysis of functional genomic data, behavioral experiments, and genetic manipulation to determine the regulatory architecture underlying variation in a courtship trait in flies (Fig 8). Our results provide evidence that $Obp56h$ expression affects copulation latency in house fly, consistent with its effects in $D.\ melanogaster$ (Shorter et al. 2016). Based on their locations on house fly chromosome III, positions as hub genes in a house fly co-expression module (Fig. 3), and divergent expression between the III$^M$ and standard III chromosomes (Fig. 5), we hypothesize that LOC101893651 ($CG2120$), Md-gd, and/or Md-Gr35 ($Gr98c$) negatively regulate $Obp56h$ expression (Fig. 8B). In addition, we hypothesize that $Obp56h$ negatively regulates CG8745 (LOC101887703) because knockdown of $Obp56h$ causes an increase in CG8745 expression in $D.\ melanogaster$ (Fig. 6B). Together, these results suggest that there is a $trans$ regulatory loop in which $Obp56h$ is both regulated by and regulates expression of genes on fly Muller element A (house fly chromosome III and $D.\ melanogaster$ chromosome X).
We have experimentally validated that CG2120 affects copulation latency in a way that is consistent with negative regulation of Obp56h in D. melanogaster neurons (Fig 7D). We propose the name speed date (spdt) for CG2120 because its expression reduces copulation latency, and we refer to the house fly ortholog (LOC101893651) as Md-spdt. Future manipulative experiments will help in further evaluating the direction of regulation of the other co-expressed genes.

**Sexual antagonism, pleiotropic constraints, and sexual selection**

Repression of Obp56h expression reduces copulation latency, which we show is associated with an advantage in male-male competition for female mates (Fig. 2). The apparently simple correlation between Obp56h expression and copulation latency suggests that sexual selection should favor downregulation of Obp56h expression. This raises an important question: why do some males not downregulate Obp56h expression to gain a sexually selected advantage? YM males do indeed express Obp56h at lower levels than YM males (Fig. 1), demonstrating that downregulation is possible. In addition, experimental repression of Obp56h in D. melanogaster males also reduces copulation latency relative to wild-type males (Shorter et al. 2016), further demonstrating the possibility for downregulation.

We hypothesize that the regulatory architecture associated with Obp56h expression creates evolutionary constraints that inhibit selection to reduce expression of Obp56h in YM males and D. melanogaster. Our analysis of coexpressed genes suggests that Obp56h exists within a complex regulatory network, including trans feedback, that is conserved between Drosophila and house fly (Fig. 8). This network architecture likely constrains the evolution of trans regulators of Obp56h and also creates downstream pleiotropic effects of changes to Obp56h expression. These pleiotropic effects likely create correlations between traits and weaken the response to selection (Lande and Arnold 1983).

Selective constraints on Obp56h expression may arise because the trans regulators are evolutionary constrained. We identified three candidate negative regulators of Obp56h (spdt, gd, and Gr98c), one or more of which may be constrained in their ability to increase in expression. Importantly, trans regulators are predicted to have pleiotropic effects, which could impede the response to selection on traits they affect (Carroll 2005). For example, gd is a serine protease involved in Toll signaling (Konrad et al. 1998; LeMosy et al. 2001). The Toll signaling pathway regulates cellular processes, including embryonic development and immune response to infection (Belvin and Anderson 1996; Valanne et al. 2011). Therefore, upregulation of gd could have pleiotropic effects that constrain the evolution of gd expression.

The expression profile of spdt (CG2120) suggests that its upregulation in head or neurons might be similarly constrained. Notably, spdt is expressed highly in embryos and in the adult digestive tract, but much lower at other developmental stages or tissues, including adult head (Fig. 7A–C).
The high expression in digestive tissues suggests that it is mechanistically possible for spdt to evolve higher expression in neurons, which would decrease copulation latency (Fig. 7D). We hypothesize that constraints on the regulators of spdt, or the genes it regulates, likely prevent its upregulation in neurons—specifically neurons in the head that affect copulation latency. Consistent with this prediction, there are 6,142 binding locations identified for Spdt from ChIP-seq data (Kudron et al. 2018), suggesting that upregulation of spdt would have downstream effects on the expression of a large portion of genes in the genome.

We additionally hypothesize that downregulation of Obp56h may have negative fitness effects that further prevent the evolution of reduced expression, even though it could be advantageous to males. One deleterious effect may arise from the interaction between Obp56h and Gr98c (Md-Gr35). Md-Gr35 is a hub gene in the house fly co-expression module containing Obp56h, and the expression levels of Md-Gr35 and Obp56h are negatively correlated (Fig. 3). In addition, we find evidence for both cis and trans effects on the upregulation of Md-Gr35 in III\textsuperscript{M} males (Fig. 5C), suggesting a possible feedback on Md-Gr35 from Obp56h downregulation. Obps most typically interact with chemoosensory receptors (odorant, ionotropic, and gustatory receptors) in the detection of chemical cues or signals, although they can have other functions as well (Zhou 2010; Benoit et al. 2017, Sun et al. 2018b). Md-Gr35 is the only chemoosensory receptor assigned to the co-expression module, suggesting that Md-Gr35 and Obp56h are each other’s interacting partners. Chemoosensory binding proteins and receptors are also known to co-regulate one another, and a negative correlation between the expression of a chemoosensory receptor and its interacting binding protein has previously been reported in a pair of genes that modulate male Drosophila mating behavior (Park et al. 2006). If Obp56h serves a sensory detection role in fly heads, then Gr98c (Md-Gr35) is a promising candidate gene with which it interacts. The interactions between these two genes may constrain selection on Obp56h expression levels.

One mechanism by which interactions between Obp56h, Gr98c (Md-Gr35), and other co-expressed genes may create constraints is through the production of cuticular hydrocarbons (CHCs). CHCs are lipid compounds used for both chemical communication and resistance to various environmental stressors, including desiccation (Chung and Carroll 2015). The expression of Obp56h is associated with CHC profiles in D. melanogaster (Shorter et al. 2016). In addition, LOC105262120, another hub gene in the network (Fig. 3), is homologous to CG10514, whose expression is correlated with CHC production in Drosophila serrata (McGraw et al. 2011). CHCs are often under strong sexual selection across insect systems, with individual or combinations of CHCs serving as important mating cues (Thomas and Simmons 2009, Berson et al. 2019a,b). The correlation of multiple hub genes with CHC profiles in Drosophila provides additional evidence that Obp56h expression, and the house fly co-expression module more generally, is related to male mating behavior, and possibly under sexual selection.
Pleiotropic or intersexual conflicts may arise over Obp56h expression because CHCs are important for protection against abiotic stressors (Arya et al. 2010; Blomquist and Bagnères 2010; Otte et al. 2018, Sun et al. 2018a), including desiccation resistance (Lockey 1988). Disrupted expression of individual genes responsible for CHC production in D. melanogaster can result in significant alterations to both mating behaviors and ecologically relevant phenotypes (Marcillac et al. 2005; Shorter et al. 2016), which could impede the response to sexual selection via trade-offs with natural selection (Rowe and Rundle 2021). This dual role of CHCs in mating and desiccation resistance suggests that sexual selection on Obp56h expression could be pleiotropically constrained by trade-offs with stress response. Our GO enrichment analysis on both the house fly and D. melanogaster RNA-seq data also revealed that Obp56h expression is correlated with the expression of genes involved in general stress responses, supporting this hypothesis. Pleiotropic constraints on Obp56h expression (because of correlated changes in CHCs) could therefore reduce the response to selection on male copulation latency.

Additional pleiotropic constraints are possible because Obps have myriad functions beyond chemical detection (Findlay et al. 2008; Arya et al. 2010; Benoit et al. 2017, Sun et al. 2018a). Obp56h expression specifically has been shown to affect mating behavior, avoidance of bitter tastants, and the expression of genes related to immune response and heat stress (Swarup et al. 2014; Shorter et al. 2016). Moreover, single base pair changes at the Obp56h locus in D. melanogaster cause sex-specific effects on a wide range of fitness-related traits, including heat resistance (Mokashi et al. 2021). Future studies should aim at determining whether Obp56h expression in III M and Y M male house flies is also associated with other phenotypes, such as CHC profiles, desiccation resistance, or tolerance to other environmental stressors.

In general, our results provide evidence that pleiotropy can reduce the response to sexual selection by creating genetic covariation amongst unrelated traits (Fitzpatrick 2004; Chenoweth and McGuigan 2010). Specifically, selection on Obp56h expression and male courtship behavior could be weakened by trade-offs between courtship behavior and stress response. In addition, the regulatory architecture underlying Obp56h expression likely creates additional pleiotropic constraints that could impede selection on copulation latency. One expected consequence of these pleiotropic constraints is that there will be genetic variance for sexually selected traits in natural populations (Kirkpatrick and Ryan 1991; Turelli and Barton 2004; Johnston et al. 2013; Heinen-Kay et al. 2020), which we observe in the form of variation in copulation latency between III M and Y M males.

**Sexually antagonistic alleles on Y chromosomes**

Our results suggest that III M males may have overcome the pleiotropic constraints associated with Obp56h expression via the Y-linkage of at least one trans regulator of Obp56h. We identified three candidate trans regulators of Obp56h on the house fly III M chromosome (Md-spdt, Md-gd, and Md-Gr35), all of which could be inhibitors of Obp56h expression (Fig. 8). This
is consistent with previous work suggesting that male-beneficial *trans* regulatory alleles could be important for the fitness effects of the house fly III\(^M\) chromosome (Adhikari et al. 2021). More generally, *trans* regulation of autosomal and X chromosome expression appears to be an important feature of the phenotypic and fitness effects of Y chromosomes (Chippindale and Rice 2001; Lemos et al. 2008, 2010; Brown et al. 2020; Morris et al. 2020; Kawamoto et al. 2021; Sandkam et al. 2021),

We hypothesize that the pleiotropic constraints associated with the regulation of *Obp56h* expression are a source of sexual antagonism. As described above, deleterious fitness effects could result from both pleiotropic effects of upregulation of the negative regulators of *Obp56h* or downstream effects of the downregulation of *Obp56h*. Regardless of the cause, these fitness costs are likely to affect both males and females. However, downregulation of *Obp56h* confers a fitness benefit to males that may offset any fitness costs. III\(^M\) males can realize this fitness benefit because the III\(^M\) chromosome carries at least one negative *trans* regulator of *Obp56h*. Therefore, our results provide evidence that sexual antagonism can arise via conflicts between sexual selection in males and opposing pleiotropic effects in females (Lande 1980; Fitzpatrick 2004; Mank et al. 2008). It also appears that the intersexual conflict over *Obp56h* regulation was (at least partially) resolved with the Y-linkage of the *trans* regulators in III\(^M\) males.

Upregulation of the III\(^M\) alleles of *Md-spdt*, *Md-gd*, and/or *Md-Gr35* alone cannot fully resolve the intersexual conflict over *Obp56h* regulation for at least two reasons. First, YM males do not realize the fitness benefit because they do not carry a III\(^M\) chromosome. Therefore, YM males remain evolutionarily constrained in their expression of *Obp56h*, suggesting that other fitness benefits are responsible for the maintenance of the YM chromosome. Previous work suggests that temperature-dependent fitness differences could be responsible—specifically YM males have higher fitness at colder temperatures (Feldmeyer et al. 2008; Delclos et al. 2021). This suggests that the III\(^M\) courtship advantage may be temperature-sensitive, which could create context-dependent fitness effects that favor YM males in specific environments. This is consistent with the prediction that genotype-by-environment interactions can maintain variation in sexually selected traits across heterogeneous environments (Kokko and Heubel 2008). In contrast to that prediction, we failed to detect effects of temperature on the III\(^M\) courtship advantage (Fig. 2), although we did not comprehensively test for environmental effects. Additional work could evaluate whether trade-offs across environments create opportunities for YM males to have courtship advantages in specific contexts.

A second reason that the intersexual conflict over *Obp56h* regulation is not fully resolved is that female house flies can carry the III\(^M\) chromosome. The male determining gene on the YM and III\(^M\) chromosome (*Mdmd*) is a negative regulator of *Md-tra* splicing, and there is a dominant allele (*Md-tra\(^D\)*) that is desensitized to the effects of *Mdmd* (Hediger et al. 2010; Sharma et al. 2017). This allows female house flies to carry YM or III\(^M\) if they also carry *Md-tra\(^D\)*. Females
with III<sup>M</sup> and Md-<i>tra</i><sup>D</sup> in their genotype may suffer a fitness cost from upregulation of Md-<i>spdt</i>, Md-<i>gd</i>, and Md-<i>Gr35</i> and/or downregulation of <i>Obp56h</i>. Therefore, even though the putatively sexually antagonistic alleles are Y-linked, they can still be carried by females, leaving the conflict unresolved. Consistent with this interpretation, previous work has determined that sexually antagonistic fitness effects of the house fly proto-Y chromosomes could contribute to the maintenance of the Y<sup>M</sup>-III<sup>M</sup> polymorphism within natural populations (Meisel et al. 2016; Meisel 2021). Our results provide a mechanism by which the III<sup>M</sup> chromosome could confer sexually antagonistic effects.

**Commonalities across sex chromosome evolution**

Our results have two additional implications for our understanding of the evolution of sex chromosomes. First, our results provide evidence that gene duplication is important for the acquisition of genes with male-specific functions during the evolution of Y chromosomes (Koerich et al. 2008). The most central gene in the co-expression module, LOC101887703, has a paralog in the house fly genome (LOC101890114) that is predicted to be on chromosome III. These two genes are homologous to <i>D. melanogaster</i> CG8745, which we hypothesize is negatively regulated by <i>Obp56h</i> based on conserved correlated expression in <i>D. melanogaster</i> and house fly (Fig. 6). The two transcripts encoded by LOC101887703 and LOC101890114 are <1% diverged in their nucleotide sequences, suggesting a recent duplication event. CG8745 is broadly expressed in many <i>D. melanogaster</i> tissues (Chintapalli et al. 2007), and broadly expressed genes often give rise to paralogs with sex-specific expression (Meisel et al. 2009).

We hypothesize that one of the two paralogs (LOC101887703 or LOC101890114) has evolved a male-specific function, which resolved an intersexual conflict associated with the coregulation of CG8745 and <i>Obp56h</i>. It is not clear which gene is ancestral and which is derived. However, a derived paralog with sex-specific expression is consistent with duplication of a broadly expressed gene to resolve an inter-sexual conflict (Connallon and Clark 2011; Gallach and Betrán 2011; Van Kuren and Long 2018). Those types of duplications often involve genes on sex chromosomes—either when a broadly expressed X-linked gene gives rise to a an autosomal gene with sex-specific expression, or when a gene is duplicated onto the Y chromosome and evolves male-specific expression (Betrán et al. 2002; Emerson et al. 2004; Koerich et al. 2008; Meisel et al. 2009; Hall et al. 2013; Mahajan and Bachtrog 2017; Ricchio et al. 2021). Consistent with this evolutionary trajectory, the only identified differences between the house fly Y<sup>M</sup> and X chromosomes thus far have been autosome-to-Y duplications (Meisel et al. 2017). Future work could address a potential sexually dimorphic subfunctionalization of the two CG8745 paralogs in the house fly genome.

Second, our results suggest that intersexual conflict may have been an important factor in the convergent evolution of Muller element A into a sex chromosome in both <i>Drosophila</i> and house fly. Muller element F was the X chromosome of the most recent common ancestor of <i>Drosophila</i>...
and house fly, and element A became the \textit{Drosophila} X chromosome after the divergence with most other flies (Vicoso and Bachtrog 2013). Element A also recently became a sex chromosome in house fly when it acquired an \textit{Mdmd} gene, creating the III\textsuperscript{M} chromosome (Meisel et al. 2017; Sharma et al. 2017; Son and Meisel 2021). Element A has become a sex chromosome in at least two other dipteran lineages: \textit{Glossina} (tsetse flies) and \textit{Anopheles} mosquitoes (Pease and Hahn 2012; Vicoso and Bachtrog 2015), raising the possibility that element A is primed to be recruited as a sex chromosome. Similar convergent sex-linkage of the same chromosomal region has been observed in vertebrates (O’Meally et al. 2012; Furman and Evans 2016; Ezaz et al. 2017), which could be explained by the same gene independently acquiring a sex determining allele in multiple independent lineages (Takehana et al. 2014).

We hypothesize that element A may be convergently recruited to be a sex chromosome because the \textit{trans} regulatory connections with \textit{Obp56h} create sexually antagonistic effects related to male mating behavior. This differs from the hypothesized cause of convergent evolution of sex chromosomes in vertebrates, which is based on the expectation that some chromosomes contain an excess of genes that are predisposed to become sex determiners. Sexually antagonistic alleles are expected to be an important selective force in the formation of new sex chromosomes because sex-limited inheritance can resolve the intersexual conflict (van Doorn and Kirkpatrick 2007, 2010; Roberts et al. 2009). Our results suggest that an enrichment of genes involved in a regulatory network with sexually antagonistic effects could promote the sex-linkage of the same chromosome in distantly related species without convergent evolution of a master sex determine.

\textbf{Conclusions}

We have identified \textit{Obp56h} expression as the likely cause of differences in courtship performance between Y\textsuperscript{M} and III\textsuperscript{M} house flies. We further identified multiple candidate \textit{trans} regulators of \textit{Obp56h} on the III\textsuperscript{M} chromosome, one of which (\textit{spdt}) we experimentally verified in \textit{D. melanogaster}. Our results demonstrate how the \textit{trans} regulators of gene expression could have sexually antagonistic effects, which are resolved via the Y-linkage of those \textit{trans} factors. This is the first time, to our knowledge, that a mechanism has been ascribed to the observation that the fitness effects of Y chromosomes can manifest via \textit{trans} effects on autosomal genes, which affect male courtship and other sexually selected traits (Chippindale and Rice 2001; Morris et al. 2020; Kawamoto et al. 2021; Sandkam et al. 2021).

\textbf{ACKNOWLEDGEMENTS}

Rebecca Presley assisted with house fly courtship assays, and Poala Najera assisted with \textit{Drosophila} courtship assays. This work was completed in part through the use of the Maxwell Cluster provided by the Research Computing Data Core at the University of Houston. This material is based upon work supported by the National Science Foundation under Grant No.
REFERENCES

Abbott, J. K., A. K. Nordén, and B. Hansson. 2017. Sex chromosome evolution: historical insights and future perspectives. *Proc. Biol. Sci.* 284.

Adhikari, K., J. H. Son, A. H. Rensink, J. Jaweria, D. Bopp, L. W. Beukeboom, and R. P. Meisel. 2021. Temperature-dependent effects of house fly proto-Y chromosomes on gene expression could be responsible for fitness differences that maintain polygenic sex determination. *Mol. Ecol.* 30:5704-5720.

Arya, G. H., A. L. Weber, P. Wang, M. M. Magwire, Y. L. S. Negron, T. F. C. Mackay, and R. R. H. Anholt. 2010. Natural variation, functional pleiotropy and transcriptional contexts of odorant binding protein genes in *Drosophila melanogaster*. *Genetics* 186:1475–1485.

Atkinson, D. 1996. Ectotherm life-history responses to developmental temperature. Pp. 183–204 in A. F. B. Ian A. Johnston, ed. Animals and Temperature: Phenotypic and Evolutionary Adaptation.

Bachtrog, D., J. E. Mank, C. L. Peichel, M. Kirkpatrick, S. P. Otto, T.-L. Ashman, M. W. Hahn, J. Kitano, I. Mayrose, R. Ming, N. Perrin, L. Ross, N. Valenzuela, J. C. Vamosi, and Tree of Sex Consortium. 2014. Sex determination: why so many ways of doing it? *PLoS Biol.* 12:e1001899.

Bates, D., M. Mächler, B. Bolker, and S. Walker. 2015. Fitting linear mixed-effects models using lme4. *J. Stat. Software* 67:1–48.

Belvin, M. P., and K. V. Anderson. 1996. A conserved signaling pathway: the *Drosophila* toll-dorsal pathway. *Annu. Rev. Cell Dev. Biol.* 12:393–416.

Benjamini, Y., and Y. Hochberg. 1995. Controlling the false discovery rate: a practical and powerful approach to multiple testing. *J. R. Stat. Soc. Series B Stat. Methodol.* 57:289–300.

Benoit, J. B., A. Vigneron, N. A. Broderick, Y. Wu, J. S. Sun, J. R. Carlson, S. Aksoy, and B. L. Weiss. 2017. Symbiont-induced odorant binding proteins mediate insect host hematopoiesis. *eLife* 6.
Berson, J. D., F. Garcia-Gonzalez, and L. W. Simmons. 2019a. Experimental evidence for the role of sexual selection in the evolution of cuticular hydrocarbons in the dung beetle, *Onthophagus taurus*. *J. Evol. Biol.* 32:1186–1193.

Berson, J. D., M. Zuk, and L. W. Simmons. 2019b. Natural and sexual selection on cuticular hydrocarbons: a quantitative genetic analysis. *Proc. Biol. Sci.* 286:20190677.

Betrán, E., K. Thornton, and M. Long. 2002. Retroposed new genes out of the X in *Drosophila. Genome Res.* 12:1854–1859.

Beukeboom, L. W., and N. Perrin. 2014. The Evolution of Sex Determination. Oxford University Press.

Bliss, C. I. 1926. Temperature characteristics for prepupal development in *Drosophila melanogaster*. *J. Gen. Physiol.* 9:467–495.

Blomquist, G. J., and A.-G. Bagnères. 2010. Insect Hydrocarbons: Biology, Biochemistry, and Chemical Ecology. Cambridge University Press.

Brankatschk, M., and S. Eaton. 2010. Lipoprotein particles cross the blood-brain barrier in *Drosophila*. *J. Neurosci.* 30:10441–10447.

Bray, N. L., H. Pimentel, P. Melsted, and L. Pachter. 2016. Near-optimal probabilistic RNA-seq quantification. *Nat. Biotechnol.* 34:525–527.

Brown, E. J., A. H. Nguyen, and D. Bachtrog. 2020. The *Drosophila* Y chromosome affects heterochromatin integrity genome-wide. *Mol. Biol. Evol.* 37:2808–2824.

Brown, J. B., N. Boley, R. Eisman, G. E. May, M. H. Stoiber, M. O. Duff, B. W. Booth, J. Wen, S. Park, A. M. Suzuki, K. H. Wan, C. Yu, D. Zhang, J. W. Carlson, L. Cherbas, B. D. Eads, D. Miller, K. Mockaitis, J. Roberts, C. A. Davis, E. Frise, A. S. Hammonds, S. Olson, S. Shenker, D. Sturgill, A. Samsonova, R. Weiszmann, G. Robinson, J. Hernandez, J. Andrews, P. J. Bickel, P. Carninci, P. Cherbas, T. R. Gingeras, R. A. Hoskins, T. C. Kaufman, E. C. Lai, B. Oliver, N. Perrimon, B. R. Graveley, and S. E. Celniker. 2014. Diversity and dynamics of the *Drosophila* transcriptome. *Nature* 512:393–399.
Bryant, E. H. 1980. Geographic variation in components of mating success of the housefly, *Musca domestica* L., in the United States. *Am. Nat.* 116:655–669.

Burke, N. W., and G. I. Holwell. 2021. Increased male mating success in the presence of prey and rivals in a sexually cannibalistic mantis. *Behav. Ecol.* 32:574–579.

Carroll, S. B. 2005. Evolution at two levels: on genes and form. *PLoS Biol.* 3:e245.

Castel, S. E., A. Levy-Moonshine, P. Mohammadi, E. Banks, and T. Lappalainen. 2015. Tools and best practices for data processing in allelic expression analysis. *Genome Biol.* 16:195.

Charlesworth, D., B. Charlesworth, and G. Marais. 2005. Steps in the evolution of heteromorphic sex chromosomes. *Heredity* 95:118–128.

Chenoweth, S. F., and K. McGuigan. 2010. The genetic basis of sexually selected variation. Annu. Rev. Ecol. Evol. Syst. 41:81–101. *Annual Reviews.*

Chintapalli, V. R., J. Wang, and J. A. T. Dow. 2007. Using FlyAtlas to identify better *Drosophila melanogaster* models of human disease. *Nat. Genet.* 39:715–720.

Chippindale, A. K., and W. R. Rice. 2001. Y chromosome polymorphism is a strong determinant of male fitness in *Drosophila melanogaster*. *Proc. Natl. Acad. Sci. USA.* 98:5677–5682.

Chung, H., and S. B. Carroll. 2015. Wax, sex and the origin of species: Dual roles of insect cuticular hydrocarbons in adaptation and mating. *Bioessays* 37:822–830.

Connallon, T., and A. G. Clark. 2011. The resolution of sexual antagonism by gene duplication. *Genetics* 187:919–937.

Delclos, P. J., K. Adhikari, J. E. Cambric, A. G. Matuk, R. I. Presley, J. Tran, and R. P. Meisel. 2021. Thermal tolerance and preference are both consistent with the clinal distribution of house fly proto-Y chromosomes. *Evol. Lett.* evl3.248.

Dobin, A., C. A. Davis, F. Schlesinger, J. Drenkow, C. Zaleski, S. Jha, P. Batut, M. Chaisson, and T. R. Gingeras. 2013. STAR: ultrafast universal RNA-seq aligner. *Bioinformatics* 29:15–21.

Edgar, R. C. 2004. MUSCLE: a multiple sequence alignment method with reduced time and space
complexity. *BMC Bioinformatics* 5:113.

Emerson, J. J., H. Kaessmann, E. Betrán, and M. Long. 2004. Extensive gene traffic on the mammalian X chromosome. *Science* 303:537–540.

Ewen-Campen, B., D. Yang-Zhou, V. R. Fernandes, D. P. González, L.-P. Liu, R. Tao, X. Ren, J. Sun, Y. Hu, J. Zirin, S. E. Mohr, J.-Q. Ni, and N. Perrimon. 2017. Optimized strategy for in vivo Cas9-activation in *Drosophila*. *Proc. Natl. Acad. Sci. USA*. 114:9409–9414.

Ezaz, T., K. Srikulnath, and J. A. M. Graves. 2017. Origin of amniote sex chromosomes: an ancestral super-sex chromosome, or common requirements? *J. Hered.* 108:94–105.

Feldmeyer, B., M. Kozielska, and B. Kuiper. 2008. Climatic variation and the geographical distribution of sex-determining mechanisms in the housefly. *Evol. Ecol. Res.* 10:797-809.

Findlay, G. D., X. Yi, M. J. MacCoss, and W. J. Swanson. 2008. Proteomics reveals novel *Drosophila* seminal fluid proteins transferred at mating. *PloS Biol.* 6:e178.

Fitzpatrick, M. J. 2004. Pleiotropy and the genomic location of sexually selected genes. *Am. Nat.* 163:800–808.

Foster, G. G., M. J. Whitten, C. Konovalov, J. T. A. Arnold, and G. Maffi. 1981. Autosomal genetic maps of the Australian Sheep Blowfly, *Lucilia cuprina dorsalis* R.-D. (Diptera: Calliphoridae), and possible correlations with the linkage maps of *Musca domestica* L. and *Drosophila melanogaster* (Mg.). *Genet. Res.* 37:55–69.

Fox, J., and M. S. Carvalho. 2012. TheRcmdrPlugin.survivalPackage: Extending theRCommander Interface to Survival Analysis. *J. Stat. Software* 49:1-32.

Fox, J., M. Friendly, and S. Weisberg. 2013. Hypothesis tests for multivariate linear models using the car package. *The R Journal* 5:39.

Franco, M. G., P. G. Rubini, and M. Vecchi. 1982. Sex-determinants and their distribution in various populations of *Musca domestica* L. of Western Europe. *Genet. Res.* 40:279–293.

Furman, B. L. S., and B. J. Evans. 2016. Sequential turnovers of sex chromosomes in African clawed...
frogs (*Xenopus*) suggest some genomic regions are good at sex determination. *G3* 6:3625–3633.

Gallach, M., and E. Betrán. 2011. Intralocus sexual conflict resolved through gene duplication. *Trends Ecol. Evol.* 26:222–228.

Gelbart, W. M., and D. B. Emmert. 2013. FlyBase High Throughput Expression Pattern Data.

Graveley, B. R., A. N. Brooks, J. W. Carlson, M. O. Duff, J. M. Landolin, L. Yang, C. G. Artieri, M. J. van Baren, N. Boley, B. W. Booth, J. B. Brown, L. Cherbas, C. A. Davis, A. Dobin, R. Li, W. Lin, J. H. Malone, N. R. Mattiuzzo, D. Miller, D. Sturgill, B. B. Tuch, C. Zaleski, D. Zhang, M. Blanchette, S. Dudoit, B. Eads, R. E. Green, A. Hammonds, L. Jiang, P. Kapranov, L. Langton, N. Perrimon, J. E. Sandler, K. H. Wan, A. Willingham, Y. Zhang, Y. Zou, J. Andrews, P. J. Bickel, S. E. Brenner, M. R. Brent, P. Cherbas, T. R. Gingeras, R. A. Hoskins, T. C. Kaufman, B. Oliver, and S. E. Celniker. 2011. The developmental transcriptome of *Drosophila melanogaster*. *Nature* 471:473–479.

Hall, A. B., Y. Qi, V. Timoshevskiy, M. V. Sharakhova, I. V. Sharakhov, and Z. Tu. 2013. Six novel Y chromosome genes in Anopheles mosquitoes discovered by independently sequencing males and females. *BMC Genomics* 14:273.

Hamm, R. L., J.-R. Gao, G. G.-H. Lin, and J. G. Scott. 2009. Selective advantage for III$^M$ males over Y$^M$ males in cage competition, mating competition, and pupal emergence in *Musca domestica* L. (Diptera: Muscidae). *Environ. Entomol.* 38:499–504.

Hamm, R. L., R. P. Meisel, and J. G. Scott. 2015. The evolving puzzle of autosomal versus Y-linked male determination in *Musca domestica*. *G3* 5:371–384.

Hamm, R. L., T. Shono, and J. G. Scott. 2005. A cline in frequency of autosomal males is not associated with insecticide resistance in house fly (Diptera: Muscidae). *J. Econ. Entomol.* 98:171–176.

Hediger, M., C. Henggeler, N. Meier, R. Perez, G. Saccone, and D. Bopp. 2010. Molecular characterization of the key switch $F$ provides a basis for understanding the rapid divergence of the sex-determining pathway in the housefly. *Genetics* 184:155–170.

Heinen-Kay, J. L., R. E. Nichols, and M. Zuk. 2020. Sexual signal loss, pleiotropy, and maintenance of a
male reproductive polymorphism in crickets. *Evolution* 74:1002–1009.

Johnston, S. E., J. Gratten, C. Berenos, J. G. Pilkington, T. H. Clutton-Brock, J. M. Pemberton, and J. Slate. 2013. Life history trade-offs at a single locus maintain sexually selected genetic variation.

*Nature* 502:93–95.

Kawamoto, M., Y. Ishii, and M. Kawata. 2021. Genetic basis of orange spot formation in the guppy (*Poecilia reticulata*). *BMC Ecol. Evol.* 21:211.

Kirkpatrick, M., and M. J. Ryan. 1991. The evolution of mating preferences and the paradox of the lek.

*Nature* 350:33–38.

Koerich, L. B., X. Wang, A. G. Clark, and A. B. Carvalho. 2008. Low conservation of gene content in the *Drosophila* Y chromosome. *Nature* 456:949–951.

Kokko, H., and K. Heubel. 2008. Condition-dependence, genotype-by-environment interactions and the lek paradox. *Genetica* 134:55–62.

Konrad, K. D., T. J. Goralski, A. P. Mahowald, and J. L. Marsh. 1998. The gastrulation defective gene of *Drosophila melanogaster* is a member of the serine protease superfamily. *Proc. Natl. Acad. Sci. USA* 95:6819–6824.

Koštál, V., J. Korbelová, T. Štětina, R. Poupardin, H. Colinet, H. Zahradníčková, I. Opeakarová, M. Moos, and P. Šimek. 2016. Physiological basis for low-temperature survival and storage of quiescent larvae of the fruit fly *Drosophila melanogaster*. *Sci. Rep.* 6:32346.

Kozielska, M., B. Feldmeyer, I. Pen, F. J. Weissing, and L. W. Beukeboom. 2008. Are autosomal sex-determining factors of the housefly (*Musca domestica*) spreading north? *Genet. Res.* 90:157–165.

Kudron, M. M., A. Victorsen, L. Gevirtzman, L. W. Hillier, W. W. Fisher, D. Vafeados, M. Kirkey, A. S. Hammonds, J. Gersch, H. Ammouri, M. L. Wall, J. Moran, D. Steffen, M. Szynkarek, S. Seabrook-Sturgis, N. Jameel, M. Kadaba, J. Patton, R. Terrell, M. Corson, T. J. Durham, S. Park, S. Samanta, M. Han, J. Xu, K.-K. Yan, S. E. Celniker, K. P. White, L. Ma, M. Gerstein, V. Reinke, and R. H. Waterston. 2018. The ModERN resource: Genome-wide binding profiles for hundreds of *Drosophila*
and *Caenorhabditis elegans* transcription factors. *Genetics* 208:937–949.

Kumar, S., G. Stecher, M. Li, C. Knyaz, and K. Tamura. 2018. MEGA X: Molecular evolutionary genetics analysis across computing platforms. *Mol. Biol. Evol.* 35:1547–1549.

Lande, R. 1980. Sexual dimorphism, sexual selection, and adaptation in polygenic characters. *Evolution* 34:292–305.

Lande, R., and S. J. Arnold. 1983. The measurement of selection on correlated characters. *Evolution* 37:1210–1226.

Langfelder, P., and S. Horvath. 2008. WGCNA: an R package for weighted correlation network analysis. *BMC Bioinformatics* 9:559.

Langfelder, P., P. S. Mischel, and S. Horvath. 2013. When is hub gene selection better than standard meta-analysis? *PLoS One* 8:e61505.

Leek, J. T., W. E. Johnson, H. S. Parker, A. E. Jaffe, and J. D. Storey. 2012. The sva package for removing batch effects and other unwanted variation in high-throughput experiments. *Bioinformatics* 28:882–883.

Lemos, B., L. O. Araripe, and D. L. Hartl. 2008. Polymorphic Y chromosomes harbor cryptic variation with manifold functional consequences. *Science* 319:91–93.

Lemos, B., A. T. Branco, and D. L. Hartl. 2010. Epigenetic effects of polymorphic Y chromosomes modulate chromatin components, immune response, and sexual conflict. *Proc. Natl. Acad. Sci. USA.* 107:15826–15831.

LeMosy, E. K., Y. Q. Tan, and C. Hashimoto. 2001. Activation of a protease cascade involved in patterning the *Drosophila* embryo. *Proc. Natl. Acad. Sci. USA.* 98:5055–5060.

Li, A., and S. Horvath. 2007. Network neighborhood analysis with the multi-node topological overlap measure. *Bioinformatics* 23:222–231.

Lockey, K. H. 1988. Lipids of the insect cuticle: origin, composition and function. *Comp. Biochem.* *Physiol. B: Comp. Biochem.* 89:595–645.
Loeb, J., and J. H. Northrop. 1917. On the influence of food and temperature upon the duration of life. *J. Biol. Chem.* 32:103–121.

Love, M. I., W. Huber, and S. Anders. 2014. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. *Genome Biol.* 15:550.

Maere, S., K. Heymans, and M. Kuiper. 2005. BiNGO: a Cytoscape plugin to assess overrepresentation of Gene Ontology categories in Biological Networks. *Bioinformatics* 21:3448–3449.

Mahajan, S., and D. Bachtrog. 2017. Convergent evolution of Y chromosome gene content in flies. *Nat. Commun.* 8:785.

Mank, J. E., D. J. Hosken, and N. Wedell. 2014. Conflict on the sex chromosomes: cause, effect, and complexity. *Cold Spring Harb. Perspect. Biol.* 6:a017715.

Mank, J. E., L. Hultin-Rosenberg, M. Zwahlen, and H. Ellegren. 2008. Pleiotropic constraint hampers the resolution of sexual antagonism in vertebrate gene expression. *Am. Nat.* 171:35–43.

Marcillac, F., Y. Grosjean, and J.-F. Ferveur. 2005. A single mutation alters production and discrimination of *Drosophila* sex pheromones. *Proc. Biol. Sci.* 272:303–309.

May, C. E., A. Vaziri, Y. Q. Lin, O. Grushko, M. Khabiri, Q.-P. Wang, K. J. Holme, S. D. Pletcher, P. L. Freddolino, G. G. Neely, and M. Dus. 2019. High dietary sugar reshapes sweet taste to promote feeding behavior in *Drosophila melanogaster*. *Cell Rep.* 27:1675–1685.e7.

McGraw, E. A., Y. H. Ye, B. Foley, S. F. Chenoweth, M. Higbie, E. Hine, and M. W. Blows. 2011. High-dimensional variance partitioning reveals the modular genetic basis of adaptive divergence in gene expression during reproductive character displacement. *Evolution* 65:3126–3137.

McKenna, A., M. Hanna, E. Banks, A. Sivachenko, K. Cibulskis, A. Kernytsky, K. Garimella, D. Altshuler, S. Gabriel, M. Daly, and M. A. DePristo. 2010. The Genome Analysis Toolkit: a MapReduce framework for analyzing next-generation DNA sequencing data. *Genome Res.* 20:1297–1303.

Meisel, R. P. 2021. The maintenance of polygenic sex determination depends on the dominance of fitness...
effects which are predictive of the role of sexual antagonism. *G3* jkab149.

Meisel, R. P., T. Davey, J. H. Son, A. C. Gerry, T. Shono, and J. G. Scott. 2016. Is multifactorial sex
determination in the house fly, *Musca domestica* (L.), stable over time? *J. Hered.* 107:615–625.

Meisel, R. P., C. A. Gonzales, and H. Luu. 2017. The house fly Y Chromosome is young and minimally
differentiated from its ancient X Chromosome partner. *Genome Res.* 27:1417–1426.

Meisel, R. P., M. V. Han, and M. W. Hahn. 2009. A complex suite of forces drives gene traffic from
*Drosophila* X chromosomes. *Genome Biol. Evol.* 1:176–188.

Meisel, R. P., and J. G. Scott. 2018. Using genomic data to study insecticide resistance in the house fly,
*Musca domestica*. *Pestic. Biochem. Physiol.* 151:76–81.

Meisel, R. P., J. G. Scott, and A. G. Clark. 2015. Transcriptome differences between alternative sex
determining genotypes in the house fly, *Musca domestica*. *Genome Biol. Evol.* 7:2051–2061.

Menuz, K., N. K. Larter, J. Park, and J. R. Carlson. 2014. An RNA-seq screen of the *Drosophila* antenna
identifies a transporter necessary for ammonia detection. *PLoS Genet.* 10:e1004810.

Mohapatra, P., and K. Menuz. 2019. Molecular profiling of the *Drosophila* antenna reveals conserved
genes underlying olfaction in insects. *G3* 9:3753–3771.

Mokashi, S. S., V. Shankar, J. A. Johnstun, W. Huang, T. F. C. Mackay, and R. R. H. Anholt. 2021.
Systems genetics of single nucleotide polymorphisms at the *Drosophila* Obp56h locus. *bioRxiv*
2021-06.

Morris, J., I. Darolti, W. van der Bijl, and J. E. Mank. 2020. High-resolution characterization of male
ornamentation and re-evaluation of sex linkage in guppies. *Proc. Biol. Sci.* 287:20201677.

Olafson, P. U., S. Aksoy, G. M. Attardo, G. Buckmeier, X. Chen, C. J. Coates, M. Davis, J. Dykema, S. J.
Emrich, M. Friedrich, C. J. Holmes, P. Ioannidis, E. N. Jansen, E. C. Jennings, D. Lawson, E. O.
Martinson, G. L. Maslen, R. P. Meisel, T. D. Murphy, D. Nayduch, D. R. Nelson, K. J. Oyen, T. J.
Raszick, J. M. C. Ribeiro, H. M. Robertson, A. J. Rosendale, T. B. Sackton, P. Saelao, S. L. Swiger,
S.-H. Sze, A. M. Tarone, D. B. Taylor, W. C. Warren, R. M. Waterhouse, M. T. Weirauch, J. H.
Werren, R. K. Wilson, E. M. Zdobnov, and J. B. Benoit. 2021. The genome of the stable fly, *Stomoxys calcitrans*, reveals potential mechanisms underlying reproduction, host interactions, and novel targets for pest control. *BMC Biol.* 19:1-31.

Olafson, P. U., and C. A. Saski. 2020. Chemosensory-related gene family members of the horn fly, (Diptera: Muscidae), identified by transcriptome analysis. *Insects* 11.

O’Meally, D., T. Ezaz, A. Georges, S. D. Sarre, and J. A. M. Graves. 2012. Are some chromosomes particularly good at sex? Insights from amniotes. *Chromosome Res.* 20:7–19.

Otte, T., M. Hilker, and S. Geiselhardt. 2018. Phenotypic plasticity of cuticular hydrocarbon profiles in insects. *J. Chem. Ecol.* 44:235–247.

Pan, J. W., J. McLaughlin, H. Yang, C. Leo, P. Rambarat, S. Okuwa, A. Monroy-Eklund, S. Clark, C. D. Jones, and P. C. Volkan. 2017. Comparative analysis of behavioral and transcriptional variation underlying CO2 sensory neuron function and development in *Drosophila Fly* 11:239–252.

Park, S. K., K. J. Mann, H. Lin, E. Starostina, A. Kolski-Andreaco, and C. W. Pikielny. 2006. A *Drosophila* protein specific to pheromone-sensing gustatory hairs delays males’ copulation attempts. *Curr. Biol.* 16:1154–1159.

Pease, J. B., and M. W. Hahn. 2012. Sex chromosomes evolved from independent ancestral linkage groups in winged insects. *Mol. Biol. Evol.* 29:1645–1653.

R Core Team. 2019. R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria.

Ricchio, J., F. Uno, and A. B. Carvalho. 2021. New genes in the *Drosophila* Y chromosome: lessons from *D. willistoni*. *Genes* 12.

Rice, W. R. 1996. Evolution of the Y sex chromosome in animals. *Bioscience* 46:331–343.

Ritchie, M. E., B. Phipson, D. Wu, Y. Hu, C. W. Law, W. Shi, and G. K. Smyth. 2015. limma powers differential expression analyses for RNA-sequencing and microarray studies. *Nucleic Acids Res.* 43:e47.
Roberts, R. B., J. R. Ser, and T. D. Kocher. 2009. Sexual conflict resolved by invasion of a novel sex
determiner in Lake Malawi cichlid fishes. Science 326:998–1001.

Rowe, L., and H. D. Rundle. 2021. The alignment of natural and sexual selection. Annu. Rev. Ecol. Evol.
Syst. 52:499–517. Annual Reviews.

Sandkam, B. A., P. Almeida, I. Darolti, B. L. S. Furman, W. van der Bijl, J. Morris, G. R. Bourne, F.
Breden, and J. E. Mank. 2021. Extreme Y chromosome polymorphism corresponds to five male
reproductive morphs of a freshwater fish. Nat. Ecol. Evol. 5:939–948.

Schaeffer, S. W. 2018. Muller “elements” in Drosophila: How the search for the genetic basis for
speciation led to the birth of comparative genomics. Genetics 210:3–13.

Scott, J. G., P. Sridhar, and N. Liu. 1996. Adult specific expression and induction of cytochrome P450lpr
in house flies. Arch. Insect Biochem. Physiol. 31:313–323.

Scott, J. G., W. C. Warren, L. W. Beukeboom, D. Bopp, A. G. Clark, S. D. Giers, M. Hediger, A. K.
Jones, S. Kasai, C. A. Leichter, M. Li, R. P. Meisel, P. Minx, T. D. Murphy, D. R. Nelson, W. R.
Reid, F. D. Rinkevich, H. M. Robertson, T. B. Sackton, D. B. Sattelle, F. Thibaud-Nissen, C.
Tomlinson, L. van de Zande, K. K. O. Walden, R. K. Wilson, and N. Liu. 2014. Genome of the
house fly, Musca domestica L., a global vector of diseases with adaptations to a septic environment.
Genome Biol. 15:466.

Shannon, P., A. Markiel, O. Ozier, N. S. Baliga, J. T. Wang, D. Ramage, N. Amin, B. Schwikowski, and
T. Ideker. 2003. Cytoscape: a software environment for integrated models of biomolecular
interaction networks. Genome Res. 13:2498–2504.

Sharma, A., S. D. Heinze, Y. Wu, T. Kohlbrenner, I. Morilla, C. Brunner, E. A. Wimmer, L. van de
Zande, M. D. Robinson, L. W. Beukeboom, and D. Bopp. 2017. Male sex in houseflies is
determined by Mdmd, a paralog of the generic splice factor gene CWC22. Science 356:642–645.

Shiao, M.-S., W.-L. Fan, S. Fang, M.-Y. J. Lu, R. Kondo, and W.-H. Li. 2013. Transcriptional profiling
of adult Drosophila antennae by high-throughput sequencing. Zool. Stud. 52:1–10.
Shono, T., and J. G. Scott. 2003. Spinosad resistance in the housefly, *Musca domestica*, is due to a recessive factor on autosome 1. *Pestic. Biochem. Physiol.* 75:1–7.

Shorter, J. R., L. M. Dembeck, L. J. Everett, T. V. Morozova, G. H. Arya, L. Turlapati, G. E. St Armour, C. Schal, T. F. C. Mackay, and R. R. H. Anholt. 2016. *Obp56h* modulates mating behavior in *Drosophila melanogaster*. G3 6:3335–3342.

Sink, H., E. J. Rehm, L. Richstone, Y. M. Bulls, and C. S. Goodman. 2001. *sidestep* encodes a target-derived attractant essential for motor axon guidance in *Drosophila*. *Cell* 105:57–67.

Son, J. H., T. Kohlbrenner, S. Heinze, L. W. Beukeboom, D. Bopp, and R. P. Meisel. 2019. Minimal effects of proto-Y chromosomes on house fly gene expression in spite of evidence that selection maintains stable polygenic sex determination. *Genetics* 213:313–327.

Son, J. H., and R. P. Meisel. 2021. Gene-level, but not chromosome-wide, divergence between a very young house fly proto-Y chromosome and its homologous proto-X chromosome. *Mol. Biol. Evol.* 38:606–618.

Sun, J. S., N. K. Larter, J. S. Chahda, D. Rioux, A. Gumaste, and J. R. Carlson. 2018a. Humidity response depends on the small soluble protein *Obp59a* in *Drosophila*. *eLife* 7:e39249.

Sun, J. S., S. Xiao, and J. R. Carlson. 2018b. The diverse small proteins called odorant-binding proteins. *Open Biol.* 8:180208.

Swarup, S., T. V. Morozova, S. Sridhar, M. Nokes, and R. R. H. Anholt. 2014. Modulation of feeding behavior by odorant-binding proteins in *Drosophila melanogaster*. *Chemical Senses* 39:125-132.

Takehana, Y., M. Matsuda, T. Myosho, M. L. Suster, K. Kawakami, T. Shin-I, Y. Kohara, Y. Kuroki, A. Toyoda, A. Fujiyama, S. Hamaguchi, M. Sakaizumi, and K. Naruse. 2014. Co-option of *Sox3* as the male-determining factor on the Y chromosome in the fish *Oryzias dancena*. *Nat. Commun.* 5:4157.

Thomas, M. L., and L. W. Simmons. 2009. Sexual selection on cuticular hydrocarbons in the Australian field cricket, *Teleogryllus oceanicus*. *BMC Evol. Biol.* 9:1-12.

Tomita, T., and Y. Wada. 1989. Multifactorial sex determination in natural populations of the housefly
(Musca domestica) in Japan. *Japanese J. Genetics* 64:373–382.

Turelli, M., and N. H. Barton. 2004. Polygenic variation maintained by balancing selection: pleiotropy, sex-dependent allelic effects and G x E interactions. *Genetics* 166:1053–1079.

Valanne, S., J.-H. Wang, and M. Rämet. 2011. The *Drosophila Toll* signaling pathway. *J. Immunol.* 186:649–656.

Van De Bor, V., G. Zimniak, L. Papone, D. Cerezo, M. Malbouyres, T. Juan, F. Ruggiero, and S. Noselli. 2015. Companion blood cells control ovarian stem cell niche microenvironment and homeostasis. *Cell Rep.* 13:546–560.

van Doorn, G. S., and M. Kirkpatrick. 2010. Transitions between male and female heterogamety caused by sex-antagonistic selection. *Genetics* 186:629–645.

van Doorn, G. S., and M. Kirkpatrick. 2007. Turnover of sex chromosomes induced by sexual conflict. *Nature* 449:909–912.

Van Kuren, N. W., and M. Long. 2018. Gene duplicates resolving sexual conflict rapidly evolved essential gametogenesis functions. *Nat. Ecol. Evol.* 2:705–712.

Vicoso, B., and D. Bachtrog. 2015. Numerous transitions of sex chromosomes in Diptera. *PLoS Biol.* 13:e1002078.

Vicoso, B., and D. Bachtrog. 2013. Reversal of an ancient sex chromosome to an autosome in *Drosophila*. *Nature* 499:332–335.

Weller, G. L., and G. G. Foster. 1993. Genetic maps of the sheep blowfly *Lucilia cuprina*: linkage-group correlations with other dipteran genera. *Genome* 36:495–506.

Zhang, Y. Q., C. K. Rodesch, and K. Broadie. 2002. Living synaptic vesicle marker: synaptotagmin-GFP. *Genesis* 34:142–145.

Zhou, J. J. 2010. Odorant-binding proteins in insects. *Vitamins & Hormones* 83:241–272.