The evolution of sex-biased gene expression in the *Drosophila* brain

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Genes with sex-biased expression in *Drosophila* are thought to underlie sexually dimorphic phenotypes and have been shown to possess unique evolutionary properties. However, the forces and constraints governing the evolution of sex-biased genes in the somatic tissues of *Drosophila* are largely unknown. By using population-scale RNA sequencing data, we show that sex-biased genes in the *Drosophila* brain are highly enriched on the X Chromosome and that most are biased in a species-specific manner. We show that X-linked male-biased genes, and to a lesser extent female-biased genes, are enriched for signatures of directional selection at the gene expression level. By examining the evolutionary properties of gene-flanking regions on the X Chromosome, we find evidence that adaptive *cis*-regulatory changes are more likely to drive the expression evolution of X-linked male-biased genes than other X-linked genes. Finally, we examine whether constraint owing to broad expression across multiple tissues and genetic constraint owing to the largely shared male and female genomes could be responsible for the observed patterns of gene expression evolution. We find that expression breadth does not constrain the directional evolution of gene expression in the brain. Additionally, we find that the shared genome between males and females imposes a substantial constraint on the expression evolution of sex-biased genes. Overall, these results significantly advance our understanding of the patterns and forces shaping the evolution of sexual dimorphism in the *Drosophila* brain.

Even though males and females share nearly identical genomes— with the exception of the Y Chromosome—they show clear phenotypic differences at physiological, morphological, and behavioral levels (Darwin 1871; Arnqvist and Rowe 2005). Thus, one of the most exciting challenges in evolutionary biology is understanding how males and females use shared genomes to produce sexually dimorphic phenotypes. A popular and tractable approach to understanding sexual dimorphism is to examine differences between dimorphic phenotypes. A popular and tractable approach to understanding sexual dimorphism is to examine differences between the sexes at the transcriptome level (Jin et al. 2001; Ellegren and Parsch 2007; Grath and Parsch 2016). Historically, most work in *Drosophila* has focused on genes with sex-biased expression in either gonad or whole-body samples, where the differential expression signal is dominated by gonad expression (Parisí et al. 2003; Ranz et al. 2003; Whittle and Extaour 2019). Studies examining the forces influencing the evolution of these genes found that sex-biased genes—and male-biased genes in particular—are highly enriched for genes showing signatures of positive selection at the protein level, suggesting that the evolution of sex-biased genes is central to adaptive evolution in *Drosophila* (Pröscl et al. 2006; Grath and Parsch 2016). Relative to female-biased and unbiased genes, the expression levels of male-biased genes tend to diverge rapidly between species (Ranz et al. 2003; Zhang et al. 2007). However, genes frequently gain and lose sex bias over evolutionary time (Ranz et al. 2003; Zhang et al. 2007). It has also been shown that sex-biased genes within gonads and whole-body samples are distributed among chromosomes in a highly nonrandom manner: Male-biased genes are significantly depleted on the X Chromosome, whereas female-biased genes are significantly enriched on the X Chromosome, which may be attributable to the lack of X Chromosome dosage compensation in the male germline (Parisí et al. 2003; Meiklejohn et al. 2011; Meiklejohn and Preregans 2012). Finally, several studies have examined sex-biased gene expression in the *Drosophila melanogaster* brain (Catalán et al. 2012; Huylmans and Parsch 2015; Pacifico et al. 2018). In contrast to findings in the gonads, male- and female-biased genes were both enriched on the X Chromosome (Catalán et al. 2012; Huylmans and Parsch 2015), suggesting that the chromosomal distributions of genes with sex-biased expression are highly tissue dependent. Despite these advances, the evolutionary properties of genes with sex-biased expression in sexually dimorphic somatic tissues such as the *Drosophila* brain remain largely unknown (Chang et al. 2011; Catalán et al. 2012).

Some possible forces, constraints, and phenomena associated with expression evolution include the faster-X effect, expression breadth across tissues, and genetic constraint. The faster-X effect is a phenomenon in which genes and regulatory regions tend to evolve more rapidly on the X Chromosome than on the autosomes (Charlesworth et al. 1987; Meisel et al. 2012; Meisel and Connallon 2013). This can be owing to the relatively rapid fixation of beneficial recessive mutations on the X Chromosome, and this effect has been observed in both sex-biased and unbiased genes but is usually most prominent in male-biased genes (Charlesworth et al. 1987; Orr and Betancourt 2001; Vicoso and Charlesworth 2006; Begun et al. 2007; Baines et al. 2008; Meisel et al. 2012; Meisel and Connallon 2013). Broad expression across multiple tissues may also act to constrain the expression evolution of genes in the brain. For instance, genes that are expressed in many tissues are likely to have an important role in at least several
tissues, and hence, their expression levels may be subjected to greater selective constraint (Orr 2000; Chen and Dokholyan 2006; Meisel et al. 2012; Papakostas et al. 2014). Additionally, the genetic constraints imposed by the largely shared male and female genomes have been shown to constrain the evolution of sexually dimorphic phenotypes in various species, including Drosophila (Poisant et al. 2010; Griffin et al. 2013). By understanding the forces and constraints governing the evolution of sex-biased gene expression in the Drosophila brain, we may begin to uncover the molecular rules underpinning the evolution of sexually dimorphic behaviors.

To understand the evolutionary properties of genes showing sex-biased expression in the brain, we generated and analyzed gene expression data from female and male brains in multiple strains of D. melanogaster, Drosophila simulans, and Drosophila yakuba. To determine the types of selection affecting expression levels, we examined interspecific and intraspecific variability in gene expression using an analysis of variance (ANOVA) framework. We used this framework to study the selective forces affecting different classes of sex-biased and unbiased genes at the expression level. Finally, we examined how expression breadth across tissues and the shared genome between sexes constrains expression evolution.

Results

Large enrichment of sex-biased genes on the X Chromosome

By using strand-specific poly(A)° RNA-seq data collected from the brains of D. melanogaster, D. simulans, and D. yakuba (Fig. 1A), we analyzed the expression patterns of each species, and identified 1246 female- and 989 male-biased genes in D. melanogaster, 767 female- and 725 male-biased genes in D. simulans, and 417 female- and 551 male-biased genes in D. yakuba at a false-discovery rate (FDR) of 0.05 (Fig. 1B; Supplemental Table S1). By looking at the chromosomal distribution of female- and male-biased genes, we found that both sets of genes were highly enriched on the X Chromosome in all three species (P < 1 × 10−5, by a factor of about two- to threefold in all species) and depleted on the autosomes, suggesting that the enrichment of X-linked genes with sex-biased expression in the brain is conserved phenomenon (Fig. 2).

To determine if the significant enrichment of sex-biased genes on the X Chromosome was unique to the brain, we analyzed previously published expression data from D. melanogaster and D. yakuba in four other tissues: gonads, thorax, abdomen, and viscera (Yang et al. 2018). We found that the large enrichment of sex-biased genes on the X Chromosome was unique to the brain, we analyzed the expression patterns of each species, and identified 1246 female- and 989 male-biased genes in all three species at an FDR of 0.05. In all three species, male-biased autosomal genes were enriched for Gene Ontology terms relating to metabolic processes (Supplemental Table S2). Other classes of genes were not consistently enriched for similar terms across the three species. In fact, with the exception of male-biased genes in D. yakuba, X-linked sex-biased genes were not enriched for any terms in any of the species.

We compared the expression levels of sex-biased and unbiased genes. The expression patterns were largely consistent between the three species, and male-biased autosomal genes had relatively high expression (Supplemental Fig. S2). We then examined the expression fold-changes and significance-levels of sex-biased genes and found that in all three species X Chromosome male-biased genes had a significantly higher fold-change than autosomal male-biased genes (Supplemental Fig. S3). Moreover, in all three species and in both sexes, sex-biased genes on the X Chromosome were more significantly sex-biased (smaller in q-value) than genes on the autosomes (Supplemental Fig. S3). Given that female-biased X Chromosome genes did not show higher fold-changes than autosomal female-biased genes (with the exception of D. simulans) but were more significantly sex-biased, our results suggest that female-biased autosomal genes may show a high degree of expression variability.

Rates of sex-bias turnover are high and chromosome specific

These observations prompted us to look at the evolution of sex-biased genes at the expression level. First, we looked at the turnover of sex-biased genes between the three Drosophila species. By only considering annotated 1:1 orthologs (Thurmond et al. 2019), we found that 57% of male-biased genes and 65% of female-biased genes within the three species are sex-biased in a species-specific manner (Fig. 3A), supporting that there is rapid sex-biased gene expression turnover in the brain. This result is consistent with findings in Heliconius butterflies (Catalán et al. 2018), suggesting that fast evolution and turnover of sex-biased genes in the brain may be a pervasive and conserved phenomenon in animals. In addition, we found the ratio of multispecies to single-species sex-biased genes to be substantially higher on the X Chromosome compared with the autosomes in both sexes (male-biased: 1.42 vs. 0.46; female-biased: 1.01 vs. 0.34; P < 2.2 × 10−16 for both; Fisher’s exact test), suggesting that chromosome-dependent patterns of
levels of both between- and within-species variability is indicative of relaxed selective constraint, whereas genes with low interspecific and intraspecific variability are likely to be evolving under stabilizing selection (Nuzhdin et al. 2004).

Considering only 1:1:1 orthologs between the three species, we found that male-biased genes on the X Chromosome (defined as genes with male bias in at least one species) were enriched for signatures of directional selection in both sexes, and female-biased genes (defined as genes with female bias in at least one species) were enriched for signatures of directional selection in females (ratios and P-values see Fig. 4A; Supplemental Fig. S4; Supplemental Table S4). Female-biased autosomal genes showed lower than expected median values for $M_{\text{species}}/M_{\text{strain}}$, which is indicative of balancing selection or adaptive variation. To determine if there is a faster-X effect for gene expression evolution in the brain, we compared $M_{\text{species}}/M_{\text{strain}}$ between genes on the X Chromosome and the autosomes. We found that sex-biased genes showed evidence of faster-X evolution in both males and females (Supplemental Table S5). No effect was found for unbiased X-linked genes.

We then examined the marginal distributions of $M_{\text{species}}$ and $M_{\text{strain}}$ and noticed that male-biased autosomal genes showed lower than expected values for both statistics in both sexes, which is suggestive of stabilizing selection (Supplemental Fig. S5A,B). To further investigate whether certain classes of genes were enriched for signatures of stabilizing selection, we examined the fold-enrichment of genes with lower than genome-wide median values for both $M_{\text{species}}$ and $M_{\text{strain}}$. We found an enrichment of signatures of stabilizing selection in two gene classes: (1) male-biased autosomal genes in both sexes and (2) unbiased genes on the X Chromosome in males (Supplemental Fig. S6). However, female-biased autosomal genes were depleted for signatures of stabilizing selection.

If the dosage compensation complex or other sex-specific trans-acting factors were the main drivers of adaptive gene expression evolution for X-linked sex-biased genes, one would expect correlations between variability ratios in males and females to be fairly low. However, we observed a high correlation (Spearman’s $r = 0.78–0.88$) between the variability ratios of sex-biased genes in males and in females (Fig. 4B). This suggests that sex-specific trans-acting factors are unlikely to be the main drivers of adaptive gene expression evolution for X-linked sex-biased genes.

To determine if the observed patterns of gene expression evolution were associated with changes in sex-biased state, we compared genes with sex bias in only one species with genes with conserved sex bias in all three species (Supplemental Fig. S7). With the exception of female-biased autosomal genes, most

**Figure 2.** Chromosomal distributions of genes with sex-biased expression in the Drosophila brain. Sex-biased genes are significantly enriched on the X Chromosome and depleted on the autosomes in the three Drosophila species. The red points indicate the observed number of sex-biased genes on a given chromosome arm (at an FDR of 0.05), and the box plots indicate the simulated numbers of sex-biased genes. (****) $P$-value $< 1 \times 10^{-15}$; (***) $P$-value $< 1 \times 10^{-12}$; (**) $P$-value $< 1 \times 10^{-4}$; (*) $P$-value $< 5 \times 10^{-2}$. The $P$-values shown are two times the fraction of simulations, which resulted in a more extreme number of sex-biased genes on a given chromosome arm.

**Selection forces.**

The selection forces acting on gene expression levels in the Drosophila brain can be analyzed using an ANOVA framework similar to the method described by Nuzhdin et al. (2004). In particular, we took the mean sum of squares for species ($M_{\text{species}}$) as a quantification of interspecific variability in expression and the mean sum of squares for strain ($M_{\text{strain}}$) as a quantification of intraspecific variability in gene expression. We then calculated the ratio of $M_{\text{species}}$ to $M_{\text{strain}}$. Genes with higher ratios are presumably evolving under the influence of directional selection, whereas genes with low ratios may be evolving under balancing selection. Genes with intermediate ratios require a closer examination of their levels of between- and within-species variability. Relatively high

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sex-biased genes showed similar patterns regardless of conservation.

Sex-biased genes with large $M_{\text{species}}/M_{\text{strain}}$ ratios (i.e., whose expression is likely evolving under directional selection) include the X-linked gene cinnamon (FBgn0000316; ratio$_{\text{male}}$ = 86.15, ratio$_{\text{female}}$ = 302.26). Cinnamon, which is involved in the biosynthesis of the molybdenum cofactor and inter-uterine aggression (Kamdar et al. 1994; Ramin et al. 2019), has changed sex bias in its recent evolutionary history: It has female-biased expression in both D. melanogaster and D. simulans and male-biased expression in D. yakuba.

Patterns of gene expression evolution are associated with the evolution of flanking sequences

To determine if DNA sequence evolution in gene-flanking regions was associated with the observed patterns of selection on gene expression, we examined the levels of nucleotide diversity in regions 20 kb upstream of and downstream from sex-biased genes in D. melanogaster. We found, compared with unbiased genes and female-biased genes, a significant depression of nucleotide diversity in male-biased genes, a significant depression of nucleotide diversity in the flanking regions of genes on the X Chromosome (excluding the genes themselves), we found that male-biased genes had significantly lower minimum HKAI values than both unbiased and female-biased X Chromosome genes ($P=0.001$ and $P=0.004$, respectively; two-sided Mann–Whitney U test) (Supplemental Fig. S8A). Overall, these results suggest that the observed enrichment of directional selection on X-linked male-biased genes is at least partly owing to adaptive evolution in the flanking (putative regulatory) sequences of these genes.

We noticed that female-biased X Chromosome genes showed no clear adaptive signatures in their flanking regions despite showing signatures of adaptation in their expression (Figs. 4A, 5). To explain this, we hypothesized that the expression evolution of male-biased X-linked genes is influenced by cis-regulatory changes to a greater extent than other classes of genes on the X Chromosome. To test this hypothesis, we first compared the nucleotide diversities in the flanking regions of diverged and conserved genes (at the gene expression level, we defined diverged genes as genes that have ratios of $M_{\text{species}}$ to $M_{\text{strain}}$ in the top 5% genome wide; conserved genes are not diverged in either sex) and found significantly reduced nucleotide diversity in the flanking regions of diverged genes (Fig. 5B). We then compared the nucleotide diversities in the flanking regions of diverged sex-biased and unbiased genes. We observed that diverged male-biased genes on the X Chromosome had significantly reduced nucleotide diversity in their flanking regions compared with other X-linked genes (Fig. 5C). When comparing HKAI in the flanking regions of highly diverged sex-biased and unbiased genes, we noticed a similar trend; however, the differences were not significant (Supplemental Fig. S8C).

Broad expression across tissues does not play a major role in constraining the directional expression evolution of genes in the Drosophila brain

To examine whether broad expression played a role in constraining expression evolution in the brain, we calculated a tissue specificity statistic (Yanai et al. 2005) for genes expressed in males and females of both D. melanogaster and D. yakuba using published expression data from four tissues (abdomen, thorax, viscera, and gonads) (Yang et al. 2018), along with our brain data. A tissue specificity near zero indicates broad expression, whereas a tissue specificity near one indicates that expression is highly tissue specific. We found a very weak negative relationship between tissue specificity (in both sexes and species) and $M_{\text{species}}/M_{\text{strain}}$ in the brain, indicating that broad expression does not constrain directional gene expression evolution in the Drosophila brain (Fig. 6A–D).

To determine if tissue specificity was associated with the observed patterns of directional expression evolution for sex-biased genes, we controlled for the effects of tissue specificity on $M_{\text{species}}/M_{\text{strain}}$ (Supplemental Fig. S9). We found that all of the patterns we observed earlier (Fig. 4A; Supplemental Fig. S4) still held. Hence, it appears as though broad expression imposes little constraint on the directional evolution of gene expression in the brain and it cannot explain the signatures of directional selection observed for sex-biased genes on the X Chromosome.

Despite the observation that expression breadth does not constrain adaptive gene expression evolution, we wondered if it could be constraining gene expression variability. Therefore, we examined the association between tissue specificity and interspecific and intraspecific gene expression variability ($M_{\text{species}}$, and
We found that tissue specificity was positively correlated with both classes of gene expression variability in the brain. Broadly expressed genes tend to have low expression variability, whereas tissue-specific genes tend to have larger variability in expression.

To test whether expression breadth could explain the enrichment of signatures of stabilizing selection on the expression levels of autosomal male-biased and X-linked unbiased genes, we examined the tissue specificities of each gene class (Supplemental Fig. S11). We observed that male-biased genes are generally broadly expressed and male-biased autosomal genes have particularly low tissue specificities. This suggests that expression breadth may partially explain the low inter- and intra-specific variability observed in male-biased autosomal genes. To examine this in greater depth, we controlled for the effects of tissue specificity on MS\textsubscript{species} and MS\textsubscript{strain} and ran our test for stabilizing selection (Supplemental Fig. S12). The patterns we observed earlier (Supplemental Fig. S6) were similar, but significance for male-biased autosomal genes was lost. This suggests that broad expression may be responsible for our observation of stabilizing selection on the expression levels of male-biased autosomal genes.

The shared genome between the sexes constrains the expression evolution of sex-biased genes in the \textit{Drosophila} brain

The shared genome has been shown to constrain the evolution of sexually dimorphic phenotypes in several species (Poissant et al. 2010; Griffin et al. 2013). An alternative to this is that sex-specific factors tend to govern the expression evolution of sex-biased genes. To distinguish between these two scenarios, we examined changes in the expression levels of sex-biased genes relative to their ancestral state in both sexes. The ancestral expression level for a given gene was estimated by maximizing the likelihood of a Brownian motion evolutionary model on the three-species phylogenetic tree. For both male- and female-biased genes in all three species, there was a strong correlation between the relative change in male expression and the relative change in female expression, indicating that male and female expression levels of sex-biased genes largely evolve in the same direction (Supplemental Fig. S13A, B).

We then sought to examine the role of genetic constraint in shaping the expression of sex-biased genes on a gene-by-gene basis. To do so, we looked at correlations between male and female expression levels for each sex-biased gene in \textit{D. melanogaster}. Because males and females within a given strain possess nearly identical genomes, genes with positive correlations between male and female expression levels are genetically constrained. Regardless of sex bias and chromosomal location, the distribution of correlations was heavily positively skewed.

Figure 4. Ratio of interspecific to intraspecific variability in expression for genes in the \textit{Drosophila} brain. (A) Male-biased genes on the X Chromosome show significantly elevated variability ratios, consistent with directional selection. Similarly, female-biased genes on the X Chromosome show signatures of directional selection in females. Female-biased autosomal genes show significantly reduced ratios, consistent with balancing selection on gene expression. Significance of medians relative to the genome-wide median is shown. P-values determined using 1 million simulations per group and Benjamini–Hochberg correction. (****) P-value <1 × 10^{-5}; (*** P-value <1 × 10^{-4}; (**) P-value <1 × 10^{-3}; (*) P-value <1 × 10^{-2}; (‘) P-value <5 × 10^{-2}. A boxplot version of the data showing dispersion around the median is presented in Supplemental Figure S4. (B) Ratios of interspecific to intraspecific variability in males (on x-axis) show large correlations (Spearman’s) with ratios in females (y-axis) irrespective of sex bias or chromosomal location. Only 1:1:1 orthologs were considered. Genes were considered male- or female-biased if they showed that bias in at least one species at an FDR of 0.05. For any given gene, the chromosomal location was taken to be the location of the \textit{D. melanogaster} ortholog.
indicating that the shared genome imposes a significant constraint on the expression evolution of sex-biased genes (Fig. 7). Given that we only have expression data from six *D. melanogaster* strains, it is likely that we consistently underestimated the correlations because it appears as though male–female correlations are generally strongly positive.

**Evolution of sex-biased genes at the protein level in *D. melanogaster***

We then examined the forces governing the evolution of sex-biased genes at the protein level in *D. melanogaster*. First, we looked for evidence of selection on sex-biased genes by examining \( d_\text{N}/d_\text{S} \) values generated from published data comparing *D. melanogaster* to *D. simulans* (Zhao and Begun 2017). \( d_\text{N}/d_\text{S} \) is the ratio of nonsynonymous substitutions per nonsynonymous site to synonymous substitutions per synonymous site, with high ratios (>1) being indicative of positive selection and low ratios being indicative of purifying selection. We found that autosomal male-biased genes are significantly enriched for signatures of purifying selection (Supplemental Fig. S14). This result, along with the relatively high expression of autosomal male-biased genes, relatively low sex-biased fold-change in all three species, low gene expression variability within and between species, broad expression, and enrichment of metabolism-related GO terms, suggests that autosomal male-biased genes have important, conserved functions in *Drosophila*.

To further understand whether positive selection has influenced the evolution of coding sequences of sex-biased genes, we examined published McDonald–Kreitman (MK) test data of annotated protein-coding genes in *D. melanogaster* (Langley et al. 2012). The MK test infers the presence of positive selection by comparing the numbers of fixed and polymorphic, synonymous and nonsynonymous substitutions (McDonald and Kreitman 1991). We found that sex-biased genes on the X Chromosome were significantly enriched, relative to the set of all genes, for genes with significant MK tests (Supplemental Table S6A). However, when comparing the numbers of genes with positive MK tests between sex-biased genes on the X Chromosome and all genes on the X Chromosome, we found that sex-biased genes showed a slight but nonsignificant enrichment (Supplemental Table S6B).

**Discussion**

We have performed the first systematic study on the evolution of sex-biased gene expression in the *Drosophila* brain. The outcomes of our analysis in combination with past studies suggest that the evolutionary properties of sex-biased genes may be tissue specific (Naqvi et al. 2019). We found a significant enrichment of genes showing male- and female-biased expression on the X Chromosome in three species of *Drosophila* (Fig. 2). The X Chromosome harbored a significantly higher relative number of genes with conserved sex-biased expression in multiple species. In a possible paradox, however, we observed an enrichment of sex-biased genes with female bias in one species and male bias in at least one other species on the X Chromosome (Fig. 3).

When examining the evolutionary forces acting on gene expression, we noticed that sex-biased X-linked genes were enriched for signatures of directional selection (Fig. 4A). Moreover, when comparing X-linked genes with autosomal genes, we found evidence for a faster-X effect for the expression evolution of sex-biased genes (Supplemental Table S5). The assumptions underlying the most common explanation for faster-X evolution of gene
expression are as follows: The mutations largely influencing gene expression on the X Chromosome are X-linked, and the mutations are at least partly recessive with respect to expression or fitness (Meisel et al. 2012; Meisel and Connallon 2013). The signatures of adaptive evolution we observed in the flanking regions of X-linked genes—especially in male-biased genes—are consistent with the first assumption (Fig. 5). Our observation of clearer signatures of adaptive evolution in genomic regions proximal to X-linked male-biased genes relative to other X-linked genes suggests that adaptive evolution in expression may be more likely to occur owing to adaptation in cis-regulatory regions for male-biased genes than unbiased and especially female-biased genes (Fig. 5C). This is consistent with previous findings suggesting that adaptive evolution of cis-regulatory regions for male-biased X-linked genes, in particular, could be partly responsible for a faster-X effect in expression for those genes (Coolon et al. 2015). Additionally, it has been observed that trans-factors responsible for the expression of X-linked genes tend to disproportionately reside on the X Chromosome (Coolon et al. 2015). Therefore, it is possible that evolution in trans-regulatory regions is partly responsible for adaptive expression evolution in sex-biased genes. Given that we observed a faster-X effect for expression in female-biased genes but not in unbiased genes may indicate that trans-factors regulating the expression of X-linked female-biased genes may be more likely to reside on the X Chromosome than trans-factors regulating unbiased X-linked genes.

We examined the role that broad expression and shared genomic constraints impose on the expression evolution of sex-biased genes and found that broad expression across tissues generally does not constrain the directional evolution of gene expression in the Drosophila brain (Fig. 6). However, we found that expression breadth does appear to constrain expression variability, both between and within species (Supplemental Fig. S10). The fact that expression breadth does not seem to inhibit directional gene expression evolution suggests that expression evolution in the brain might in part be attributable to selection acting on expression in other tissues which in turn “drag along” expression levels in the brain and vice versa.

In accordance with findings made in other systems, we found that the shared genome between the sexes significantly constrains the expression evolution of sex-biased genes (Fig. 7; Supplemental Fig. S13). This indicates that selection acting upon the expression level of a certain sex-biased gene in one sex will likely “drag along” the expression of that gene in the other sex. It has previously been shown that the chromatin structures of the X Chromosome in male and female Drosophila are similar, despite the fact that dosage compensa-
Figure 7. The shared genome constrains expression evolution of sex-biased genes in the Drosophila brain. Correlations between male and female expression levels for sex-biased genes in D. melanogaster. For each category (on the y-axis), a Pearson correlation coefficient (x-axis) was calculated for each gene using six male–female data points (one for each strain). The median correlations for each class of sex-biased gene are nonzero. One-sample Wilcoxon signed-rank test P-values <2.2 × 10^{-16} for each class (female autosomal, female X, male autosomal, and male X). Sex-biased genes are defined in D. melanogaster at an FDR of 0.05.

of a study of sex-biased expression in unmated or young flies would compare to our results. Ultimately, the long-term goal is to reveal how sex-biased gene expression in somatic tissues contributes to reproduction and fitness in a species.

Methods

Fly lines and tissue collection

We collected data from inbred strains of three Drosophila species: The D. melanogaster strains come from the Drosophila melanogaster Genetic Reference Panel (DGRP) that are wild caught flies—RAL304, RAL307, RAL357, RAL360, RAL399, RAL517—from Raleigh, North Carolina, USA (MacKay et al. 2012); D. simulans strains were FAIR3 (Zhao et al. 2015) and w^{501} (Begun et al. 2007); and D. yakuba strains were Tai18E2 (Drosophila 12 Genomes Consortium 2007) and CY07, which is a Cameroonian isofemale line (Bachtrog et al. 2006). All lines were maintained on standard media under a 12:12 light/dark cycle at 24°C.

Before the dissections, all fly lines were expanded and maintained under an undisturbed 12:12 light/dark cycle at 24°C in low density vials with fly food. All individuals were mated and precisely 4 d old. Flies were very briefly anesthetized with CO2, and heads were collected with a clean razor blade and then were transferred to a glass dissection plate with PBS for brain collection. Dissections were performed within a 3-h window always at the same circadian zeitgeber time (ZT1–ZT4; in this experiment the light was turned on at 8:00 a.m., and the dissections were performed between 9:00 a.m. and 12:00 p.m. within a 3-h interval). Male and female brains of the same line were dissected in parallel the same day at the same time. Dissected brains were immediately transferred into a low retention Eppendorf tube containing 100 µL TRIzol (Thermo Fisher Scientific) and kept frozen at −20°C until RNA extraction. We collected three biological replicates for each genotype by sex combination, and each replicate comprised a pool of exactly 15 brains.

RNA extractions and library preparation

All RNA extractions were performed according to TRIzol manufacturer protocol and immediately followed by a DNase treatment us-

ing the TURBO DNase from Thermo Fisher Scientific. RNA quality was assessed by a Bioanalyzer run of an Agilent eukaryote total RNA pico chip, whereas RNA quantity was measured with a NanoDrop One (ABI). About 50 ng total RNA was used for library preparation. Libraries were fragmented and enriched for mRNA using NEBNext poly(A) magnetic isolation module (NEB E7490) and prepared using NEBNext ultra II directional RNA library prep kit (NEB E7765) and dual indexing from the NEBNext multiplex oligos kit (NEB 7600) following manufacturer protocol including beads size selection for 200-bp inserts. Library quality was first assessed on Agilent D1000 ScreenTapes for TapeStation and then by Qubit and Agilent Bioanalyzer. Finally, 150-bp paired-ends libraries were sequenced on five lanes of an Illumina NextSeq 500 platform (Supplemental Table S7).

Read mapping and identification of sex-biased genes

Adaptors from RNA-seq reads were trimmed using Trimmmomatic (Bolger et al. 2014). In addition, low-quality bases were trimmed using Trimmomatic settings LEADING:1 TRAILING:1 SLIDING WINDOW:20:25 and MINLEN:36. HISAT2 (Sirén et al. 2014) was then used to align reads to their respective reference genomes: dmel_r6.15 for D. melanogaster, dsim_r2.02 for D. simulans, and dyak_r1.05 for D. yakuba. Gene and transcript expression levels (transcripts per million [TPM]) were then quantified using StringTie, and read counts were extracted using a script, prepDE.py, provided by StringTie (Pertea et al. 2015). We performed quality control on the data obtained from the 60 libraries: Three of them (one replicate of male FAIR3, male R517, and female R360) were excluded from further analysis because of low correlation with other replicates or deviation from other samples in the PCA plot.

The read counts were then analyzed using DESeq2 (Love et al. 2014) to identify sex-biased genes in each species. Genes with fewer than 10 read counts (summed across samples within a species) were excluded from the analysis. The DESeq2 model used to determine differential expression between the sexes was counts ~ strain + sex. Genes with statistically significant “sex” terms (FDR < 0.05) were considered to be sex-biased in the given species. In all future analyses in which we compared sex-biased genes to unbiased genes, we considered genes to be unbiased if DESeq2 assigned an insignificant P-value to the sex term for a gene. Genes for which a P-value could not be determined (“NA,” filtered out by DESeq2) were not considered for further analysis.

To directly compare gene expression levels between samples from different species we used trimmed mean M (TMM) values normalized TPM values. TMM normalization was implemented with the “r” package edgeR (Robinson et al. 2010).

Chromosomal distributions of sex-biased genes

To determine significance for the enrichment of sex-biased genes on the X Chromosome we used random sampling. For each simulation, the total number of male- and female-biased genes was taken to be the same as the number observed at an FDR of 0.05. That number of genes was then randomly sampled from a list of all genes with finite P-values (from DESeq2), and the number of sampled genes on each chromosome arm was recorded. Simulations were repeated 100,000 times for each bias-species combination. The P-values were calculated as two times the fraction of simulations, which resulted in a more extreme number of sex-biased genes on a given chromosome arm.
Expression turnover of sex-biased genes

To get a conservative estimate for the expression turnover of sex-biased genes, equal numbers of genes were taken to be sex-biased in each of the three species. This was performed by increasing the FDR in D. simulans and D. yakuba so that the numbers of sex-biased genes in each species was the same as the number found in D. melanogaster at an FDR of 0.05. Within these three sets, we subsequently only considered 1:1:1 orthologs identified using a list obtained from FlyBase (Thurmond et al. 2019). Genes that are sex-biased in one, two, or all three species were identified, as well as genes that “switched” sex bias in their recent evolutionary history. “Switched” genes are genes that are male-biased in at least one species and female-biased in at least one other species.

Gene expression evolution

One-to-one-to-one orthologs between the three species were identified using a list obtained from FlyBase (Thurmond et al. 2019). TMM normalization of TPM values was then applied across all samples and replicates simultaneously for orthologous genes. Only genes with expression higher than 1 TPM in at least one of the three species were considered for further analysis. The resulting expression values were then log2 transformed and Pareto scaled. For every single gene, we fit a model expression ~ species + strain. Strain was taken to be a random effect and was nested within species. The model was fit using the MIXED procedure in SAS. The model was fit using the type III sum of squares. We then took the ratio of mean sum of squares (MS) for the species term as a quantification of interspecific variability and MS for the strain term as a quantification of intraspecific variability. A high value for the ratio MSspecies/MSstrain was taken as indicative of directional selection on expression, whereas low values of the ratio were taken as indicative of balancing selection. To compare the ratios, interspecific variabilities, and intraspecific variabilities of various gene classes to genome-wide medians, we used simulations. We randomly sampled gene sets of identical size to the gene set of interest and computed medians for the statistic of interest. To do so, we used the function “gam,” from the “R” library mgcv, with formula y ~ s(x, bs = “cs”) (R Core Team 2019). The residuals were then calculated and plotted.

Controlling for tissue specificity

To control for the effects of tissue specificity on MSspecies/MSstrain, MSspecies, and MSstrain, we used a generalized additive model fit to the relationship between log(“statistic”) and tissue specificity (τ), where “statistic” is the statistic of interest. To do so, we used the function “gam,” from the “R” library mgcv, with formula y ~ s(x, bs = “cs”) (R Core Team 2019). The residuals were then calculated and plotted.

Direction of gene expression evolution and the shared genome constraint

To infer ancestral expression levels for a given gene, we maximized the likelihood of a Brownian motion model on the three species phylogenetic tree. Because our results provide evidence that sex-biased genes are enriched for signatures of selection at the expression level, we only used a Brownian model as an approximation. In particular, we used TMM normalized and log2 transformed expression levels (across all samples in all species) in combination with the anc.ML function from the “R” library phytools to infer the ancestral expression levels (R Core Team 2019). This was performed for sex-biased genes with 1:1:1 orthologs in all three species. Additionally, we required that all genes considered had a median expression TPM > 1 in at least one species in each sex. The ancestral state was estimated independently for males and females. For a given gene, the change in expression relative to its ancestral state was defined as (current–ancestral)/abs(ancestral), where “current” is the log2(TPM+1) of the extant expression and “ancestral” is the log2(TPM + 1) of the inferred ancestral expression. Pearson correlations between male and female changes relative to ancestral state were then calculated. To investigate the role of genetic constraint on a gene-by-gene basis, we first TMM-normalized the D. melanogaster TPM data. Then for each sex, strain, and gene, we calculated median expression, leading to six male–female pairs for each gene. By using these six data points (per gene), we calculated Pearson correlations for each sex-biased gene in D. melanogaster.

Analysis of nucleotide diversity in gene-flanking regions

SNP data for D. melanogaster were obtained from DGRP freeze2 (MacKay et al. 2012). Nucleotide diversity (π) within all DGRP strains was then calculated genome-wide in 500-bp windows using VCFTools (Danecek et al. 2011). Genes were considered “diverged” at the expression level if they had a ratio of MSspecies/MSstrain in the highest 5% genome wide in either sex. For each gene in a given class (i.e., male-biased X Chromosome, unbiased autosomes, etc.), π windows overlapping ±20-kb flanking regions were considered. The π of the gene is represented as a single point at zero. All of the data (distance from gene for each π window and the value of π within each window) from each gene in a given class were aggregated, and generalized additive model smoothing was performed. To smooth the data, we used the function “gam” from the “R” library mgcv, with formula y ~ s(x, bs = “cs”), distribution family “Gamma,” and link function “log” (R Core Team 2019). The resulting curves were plotted with the function “geom_smooth” from the “R” library “ggplot2” (Wickham 2016).

Protein evolution analysis

Protein coding gene sequences and orthologous information were downloaded from FlyBase (Thurmond et al. 2019). One-to-one orthologous genes in D. melanogaster and D. simulans were aligned using GeneWise to remove insertions and deletions and then re-aligned using PRANK with the –codon option (Zhao and Begun 2005).

Calculation of tissue specificity

RNA-seq data from the thorax, abdomen, gonads, and viscera of D. melanogaster and D. yakuba were obtained from Yang et al. (2018). Reads were aligned and expression was quantified in the same manner as for our brain data (except adaptors were not trimmed before alignment). Within each species, sex, and tissue (including our brain data), we computed median TPM values, which were then log2 transformed. Tissue specificity (τ) was then calculated in each sex and species using the formula

\[ \tau = \frac{\sum (1 - f_i)}{n - 1}; f_i = \frac{x_i}{\max(x)} \]

where \( x_i \) is the expression of gene \( x \) in tissue \( i \) (Yanai et al. 2005).

The sum and maximum are taken over all tissues, with \( n \) being the total number of tissues.
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