## The evolution of ovary-specific gene expression in Hawaiian Drosophilidae

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## 13 1 Abstract

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As detailed data on gene expression become accessible from more species, we have an opportunity to test the 14 extent to which our understanding of developmental genetics from model organisms helps predict expression 15 patterns across species. Central to this is the question: how much variation in gene expression do we expect 16 to observe between species? Here we provide an answer by comparing RNAseq data between twelve species 17 of Hawaiian Drosophilidae flies, focusing on gene expression differences between the ovary and other tissues. 18 We show that there exists a cohort of ovary-specific genes that is stable across species, and that largely 19 corresponds to described expression patterns from laboratory model *Drosophila* species. However, our results 20 also show that, as phylogenetic distance increases, variation between species overwhelms variation between 21 tissues. Using ancestral state reconstruction of expression, we describe the distribution of evolutionary 22 changes in tissue-biased expression profiles, and use this to identify gains and losses of ovarian expression 23 across these twelve species. We then use this distribution to calculate the correlation in expression evolution 24 between genes, and demonstrate that genes with known interactions in *D. melanoqaster* are significantly more 25 correlated in their evolution than genes with no or unknown interactions. Finally, we use this correlation 26 matrix to infer new networks of genes that have similar evolutionary trajectories, and we provide these as a 27

28 dataset of novel testable hypotheses about genetic roles and interactions.

## $_{29}$ 2 Introduction

Data on when and where genes are expressed are now fundamental to the study of development and disease<sup>1</sup>. 30 With continually advancing RNA sequencing technologies, these data have been collected using RNA sequenc-31 ing from a wide variety of cells, treatments and species<sup>2,3</sup>. Statistical analysis of gene expression across these 32 differentials generates insights into how gene expression is connected to phenotypic differences in morphology 33 and behavior<sup>4</sup>. However, when comparing gene expression across species, most studies have been restricted 34 to pairwise comparisons, often between one model laboratory species and one other species of interest<sup>5</sup>. One 35 challenge with such pairwise comparisons is that they lack robust information about how much evolutionary 36 variation in expression we expect to observe, making it difficult to evaluate the significance of any inter-37 specific difference in variation<sup>5,6</sup>. Instead, we need phylogenetic comparisons of expression that take into 38 account the shared history between species<sup>7,8</sup>, and that describe significant changes in expression in relation 39 to other phenotypic traits of interest.<sup>9</sup> In this study we perform a phylogenetic comparison of gene expres-40 sion across the organs of twelve species of Hawaiian Drosophilidae flies with highly divergent ovary and egg 41 morphologies. From our results we identify individual genes that have undergone significant evolutionary 42 shifts in organ-specific expression, and describe global patterns in transcriptome variation across species that 43 can serve as a benchmark for future interspecific comparisons of gene expression. 44

Phylogenetic comparisons of developmental traits are particularly valuable for building context around com-45 parisons between well-studied model organisms and their non-model relatives<sup>10</sup>. Much more has been learned 46 about the genetics and development of laboratory model species like *D. melanogaster* than may ever be pos-47 sible for the vast majority of life<sup>11</sup>. But the usefulness of model species to understand general principles 48 depends in part on the extent to which biology in these species reflects the biology of other taxa, rather 49 than species-specific phenomena<sup>12</sup>. In the case of gene expression, there has been substantial debate about 50 the degree to which patterns observed in model organisms may be representative across species 13-16. Where 51 several studies showed that the expression profiles of organs within a species are more different the pro-files of homologous organs across species<sup>17-20</sup>, other work has questioned this finding<sup>13,14</sup>. More recently, 52 53 Breschi and colleagues  $(2016)^{21}$  demonstrated that, consistent with an evolutionary model of trait evolu-54 tion, species-level variation in gene expression increases with the time since divergence from the most recent 55 common ancestor. In addition, previous work by authors on this manuscript<sup>8</sup> showed that, while expression 56 patterns across tissues tend to be consistent between species, lineage-specific shifts in expression enrichment 57 can be identified by applying phylogenetic comparative methods. With the exception of the work by Munro 58 and colleagues  $(2021)^8$ , these studies have been, to our knowledge, performed almost exclusively in vertebrate 59 species<sup>17,18,20</sup>, and for the most part placental mammals<sup>13,14,16</sup>, meaning that far less is known about organ 60 and species-level expression differences when comparing across the tree of life. 61

The detailed atlases of expression data across  $\operatorname{organs}^{22}$  and developmental timepoints<sup>23</sup> is one of the strengths 62 of model systems like D. melanogaster. These public resources make it possible to explore global patterns of 63 expression to gain insight into potential gene regulation, interaction, and function  $2^{3-25}$ . As atlases such as 64 these have become increasingly detailed and available from more taxa, a new goal has been to compare these 65 expression profiles across species<sup>7,26,27</sup>. One objective of these cross-species comparisons is to shed light on 66 potential regulatory associations between genes<sup>7,9</sup>. This is especially advantageous for complex processes 67 such as ovarian function for which we have a fragmented understanding of gene regulation despite genetic 68 and transcriptome studies within single model organisms. Another objective of phylogenetic comparisons 69 of expression at lases is to estimate the evolutionary distance between species at which we might expect a 70 given gene to demonstrate a divergent pattern of  $expression^6$ . If this distance is relatively small, then we 71 predict atlases to contain large amounts of species-specific patterns. Alternatively, if as described above, 72 variation across tissues outweighs variation across the species being compared, we predict atlases to contain 73 large cohorts of tissue-specific genes that have been evolutionarily conserved. In this study we test for the 74 existence of a core suite of ovary-specific genes across species of Hawaiian Drosophilidae and describe its size 75 and composition in relation to the described atlas of expression in *D. melanogaster*. 76

The *Drosophila* ovary has several features<sup>28</sup> that make it a compelling organ in which to test hypotheses about expression evolution. Analyses of the FlyAtlas2 dataset<sup>29</sup> show that in *D. melanogaster*, more genes demonstrate highest expression enrichment in the ovary than any other adult organ (Fig. S1). Additionally,

all described signaling pathways are known to have a role in regulating ovarian development<sup>30</sup>. The ovary 80 performs several critical functions, including maintaining the germ line and manufacturing specialized egg 81 cells, volk, and egg-shell materials<sup>31</sup>. Genetic screens<sup>30,32</sup> and experimental manipulation in D. melanoque 82 have revealed functions of many genes involved in these processes, including yolk-protein genes required 83 for oogenesis<sup>33</sup> and embryonic patterning genes with localized mRNA like  $nanos^{34}$  and  $swallow^{35}$ . Here 84 we compare whole-ovary RNA profiles to assess the extent to which these genes and others demonstrate 85 consistent patterns of ovary-enrichment over evolutionary timescales in a clade with highly divergent ovary 86 and egg morphologies. 87

The Hawaiian Drosophilidae clade contains an estimated 1,000 extant species<sup>36</sup> that diverged from a common 88 ancestor with D. melanoque petween 25 and 40 million years  $ago^{37}$ . Extant species have been studied 89 in particular for the variation in ovary and egg morphology<sup>38,39</sup>. Species of Hawaiian Drosophilidae show 90 the largest range within the family of egg size, shape, and the number of egg-producing units in the ovary, 91 known as  $ovarioles^{40-42}$ . Previous studies by our research group and others have shown that these traits are 92 likely associated with evolutionary changes in the egg-laving substrate (e.g. rotting bark, flowers, leaves)<sup>38,40</sup>. 93 Furthermore, our previous work demonstrated that at least one developmental process, governing how the 94 number of ovarioles is specified in the adult *D. melanogaster* ovary, is conserved in Hawaiian *Drosophila*<sup>40</sup>. 95 The diversity of Hawaiian species and their relationship to model species make them a strong candidate 96 model clade for evo-devo research<sup>36,43</sup>. However, their relatively long generation times and species-specific 97 breeding requirements make laboratory culture more challenging than classic Drosophila models<sup>36</sup>. In this 98 study we leverage technologies that can be deployed on wild-caught individuals to gather rich developmental ٩q data to compare across species. 100

We compared the expression profiles of twelve species of wild-caught Hawaiian Drosophilidae species across 101 three tissues: the adult ovary, head, and the remaining carcass (Fig. 1). First, we characterized the 102 differentially expressed genes in the ovary of each species individually. By comparing these to each other, 103 and to records of ovary-enriched genes from D. melanogaster, we identified a core suite of ovary genes shared 104 across species. We then repeated this analysis for head-enriched genes, and compared the results across 105 these parallel analyses to test the extent to which global patterns of expression difference are influenced 106 by the identity of the tissues in question. We applied linear modeling to this dataset to test the overall 107 contribution of species- and tissue-level differences to expression variation across genes, and describe the 108 circumstances under which one is likely to dominate over the other. Finally, we used a phylogenetic analysis 109 of expression changes over evolutionary time to identify genes likely to have gained and lost tissue-enriched 110 expression. This evolutionary screen of expression changes allowed us to identify networks of genes that 111 demonstrate correlated changes in expression evolution. We provide these networks as a searchable dataset 112 of novel, testable hypotheses for gene regulation with respect to ovarian function. The results of this study 113 demonstrate both the power of Hawaiian *Drosophila* as a model clade for evo-devo, and the potential of using 114 phylogenetic methods to identify evolutionary variation in gene expression underlying phenotypic differences. 115

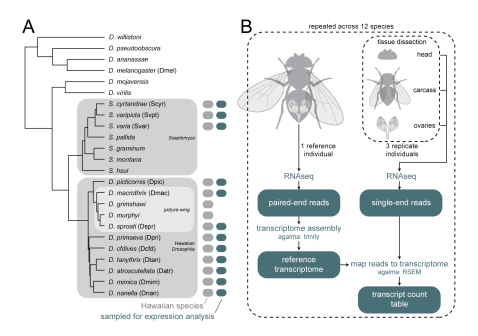


Figure 1: **Phylogeny of species and RNA sampling strategy.** A, Twelve species of Hawaiian Drosophilidae flies were collected in the wild and processed for RNA sequencing. The twelve reference transcriptomes assembled from these species were combined with twelve published genomes to generate the phylogeny shown here (originally published in Church and Extavour, 2021<sup>44</sup>). Three clades within the group are highlighted: the genus *Scaptomyza*, nested within the paraphyletic genus *Drosophila*; the Hawaiian *Drosophila*, which, along with *Scaptomyza*, make up the Hawaiian Drosophilidae; and the well-known *picture-wing* clade. Adjacent to tip labels are four letter species codes used throughout the manuscript. B, The experimental design used to generate the data in this manuscript. When sufficient specimens were available per species, one individual was used as a reference and three individuals were dissected into three separate tissues: the head, ovaries, and all remaining material (carcass). Reference individuals were sequenced to generate paired-end RNA reads and tissues were sequenced to generate single-end RNA reads. Tissue libraries were then mapped to the assembled reference to quantify transcript expression. Teal boxes indicate data files. Dashed-line boxes indicate a repeated step.

## 116 3 Methods

### 117 3.1 Field collection

Specimens used for transcriptome sampling were caught on the Hawaiian islands between May of 2016 and 118 May of 2017. Specimens were caught using a combination of net sweeping and fermented banana-mushroom 119 baits in various field sites on the Hawaiian islands of Kaua'i and Hawai'i (see Table S1 for locality data). 120 Field collections were performed under permits issued by the following: Hawai'i Department of Land and 121 Natural Resources, Hawai'i Island Forest Reserves, Kaua'i Island Forest Reserves, Koke'e State Park, and 122 Hawai'i Volcanoes National Park. Adult flies were maintained in the field on vials with sugar media and kept 123 at cool temperatures. They were transported alive back to Cambridge, MA where they were maintained on 124 standard Drosophila media at 18°C. Samples were processed for RNA extraction between 5 and 31 days after 125 collecting them live in the field (average 10.8 days, see Table S1). One species, *Scaptomyza varia*, was caught 126 in the field before the adult stage by sampling rotting *Clermontia sp.* flowers (the oviposition substrate). 127 For this species, male and female adult flies emerged in the lab, and were kept together until sampled for 128

### <sup>130</sup> 3.2 Species identification

<sup>131</sup> Species were identified using dichotomous keys<sup>45–49</sup>, when possible. Many keys for Hawaiian Drosophili-<sup>132</sup> dae are written focusing on male specific characters (e.g. sexually dimorphic features or male genitalia)<sup>47</sup>. <sup>133</sup> Therefore, for species where females could not be unambiguously identified by morphology, we verified their <sup>134</sup> identity using DNA barcoding. When males were caught from the same location, we identified males to <sup>135</sup> species using dichotomous keys and matched their barcode sequences to females included in our study. We <sup>136</sup> also matched barcodes from collected females to sequences previously uploaded to NCBI<sup>50–52</sup>.

<sup>137</sup> The following dichotomous keys were used to identify species: for *picture-wing* males and females, Magnacca

and Price  $(2012)^{45}$ ; for *antopocerus* males, Hardy  $(1977)^{46}$ ; for *Scaptomyza*, Hackman  $(1959)^{47}$ ; for species in the *mimica* subgroup of MM, O'Grady and colleagues  $(2003)^{48}$ ; for other miscellaneous species, Hardy

140  $(1965)^{49}$ .

<sup>141</sup> For DNA barcoding, DNA was extracted from one or two legs from male specimens using the Qiagen DNeasy

<sup>142</sup> blood and tissue extraction kit, or from the DNA of females isolated during RNA extraction (see below). We <sup>143</sup> amplified and sequenced the cytochrome oxidase I (COI), II (COII) and 16S rRNA genes using the primers

<sup>143</sup> amplified and sequenced the cytochrome oxidase I (COI), II (C <sup>144</sup> and protocols described in Sarikaya and colleagues  $(2019)^{40}$ .

For barcode matching, we aligned sequences using MAFFT, version v7.475<sup>53</sup>, and assembled gene trees using RAxML, version 8.2.9<sup>54</sup>. Definitive matches were considered when sequences for females formed a monophyletic clade with reference males or reference sequences from NCBI; see Table S2.

Female *D. primaeva*, *D. macrothrix*, *D. sproati*, and *D. picticornis* could be identified unambiguously using dichotomous keys. Female *D. atroscutellata*, *D. nanella*, *D. mimica*, *D. tanythrix*, *S. cyrtandrae*, *S. varipicta*, and *S. varia* were identified by matching barcodes to reference sequences from NCBI, reference males, or both. For the female *haleakalae* fly used in this study, no male flies were caught in the same location as these individuals, and no other sequences for *haleakalae* males on NCBI were an exact match with this species. Given its similar appearance to *Drosophila dives*, we are referring to it here as *Drosophila* cf *dives*, and we

await further molecular and taxonomic studies of this group that will resolve its identity.

### 155 3.3 Sampling strategy

The target number of mature, healthy female flies per species was four, with three intended for dissection and species specific expression libraries and one intended as a whole-body reference library (Fig. 1). When four such individuals were not available, a reference library was assembled by combining the tissue-specific libraries from one of the other individuals. This was the case for the following species: *D. sproati*, which was dissected and had RNA extracted separately from the head, ovaries, and carcass, with RNA combined prior to library preparation; and *S. varia*, *S. cyrtandrae* and *D.* cf *dives*, for which RNA was extracted and libraries prepared for separate tissues, and raw reads were combined after sequencing.

For the other eight species, sufficient individual females were available such that reads for transcriptome assembly were sequenced from a separate individual. In these cases one entire female fly was dissected and photographed to assess whether vitellogenic eggs were present in the ovary, and all tissues were combined in the same tube and used for RNA extraction. Library preparation failed for one individual *D. atroscutellata* fly, as well as two tissue-specific libraries: one head sample from *D. mimica*, and one head sample from *D. sproati*.

### <sup>169</sup> 3.4 Dissection and RNA sequencing

<sup>170</sup> Female flies were anesthetized in 100% ethanol and were dissected in a phosphate-buffered saline solution.

<sup>171</sup> The ovary was separated from the abdomen, and the head was separated from the carcass. Photographs

<sup>172</sup> of each tissue were taken, and tissues were moved to pre-frozen eppendorf tubes, kept in dry ice, and

 $_{173}$  immediately transported to a -80°C freezer. Dissections were performed as quickly as possible to prevent

RNA degradation. Samples were stored at -80°C for between 90 and 336 days before RNA extraction (average
281.9 days, see Table S1).

RNA was extracted from frozen samples using the standard TRIzol protocol (http://tools.thermofisher. com/content/sfs/manuals/trizol\_reagent.pdf). One mL of TRIzol was added to each frozen sample, which were then homogenized using a sterile motorized mortar. The recommended protocol was followed without modifications, using 10 µg of glycogen, and resuspending in 20µL RNAse-free water-EDTA-SDS solution. DNA for subsequent barcoding was also extracted using the phenol-chloroform phase saved from the RNA extraction.

182 RNA concentration was checked using a Qubit fluorometer, and integrity was assessed with a Agilent TapeS-

tation 4200. RNA libraries were prepared following the PrepX polyA mRNA Isolation kit and the PrepX

<sup>184</sup> RNA-Seq for Illumina Library kit, using the 48 sample protocol on an Apollo 324 liquid handling robot in

the Harvard University Bauer Core Facilities. Final library concentration and integrity were again assessed

<sup>186</sup> using the QUbit and TapeStation protocols.

Samples intended for transcriptome assembly were sequenced on an Illumina HiSeq 2500, using the standard 187 version 4 protocol, at 125 base pairs of paired-end reads. Samples intended for tissue-specific expression 188 analyses were sequenced on an Illumina NextSeq 500, using a high output flow cell, at 75 base pairs of 189 single-end reads. A table of total read counts for each library can be found in Tables S3-S4. To account 190 for any possible batch effects across separate rounds of sequencing, each sequencing run was performed with 191 one or several overlapping samples. Principle component analysis of these libraries showed variation between 192 sequencing runs to be negligible relative to variation between tissue and individual (see Results and Fig. 193 S7). 194

### <sup>195</sup> 3.5 Transcriptome assembling and expression mapping

<sup>196</sup> Transcriptome assembly and expression mapping was performed using the agalma pipeline, version 2.0.0<sup>55</sup>. <sup>197</sup> For the twelve reference transcriptomes, reads from separate rounds of sequencing were concatenated and <sup>198</sup> inserted into the agalma catalog. Further details of transcriptome assembly and homology assessment are <sup>199</sup> included in our previous manuscript<sup>44</sup>.

Each tissue-specific expression library was mapped to the corresponding reference transcriptome using the 'expression' pipeline in agalma, which uses the software RSEM to estimate gene and isoform count levels from RNAseq data<sup>56</sup>. The agalma pipeline also includes steps to catalog the species, tissue type, and run information, which were exported as a single JavaScript object notation (JSON) file. This file is available in the GitHub repository in the directory analysis/data.

### 205 3.6 Phylogenetic analysis

The phylogenetic methods for inferring homology, orthology, and estimating gene and species trees are the same as those described in our previous manuscript<sup>44</sup>. Genetrees were additionally annotated with the software Phylog<sup>57</sup>.

### <sup>209</sup> 3.7 Annotating transcripts by sequence similarity

We leveraged the close relationship of these species to species of *Drosophila* with well-annotated genomes to annotate the transcripts considered here. For each transcript in the reference transcriptome, we performed four comparisons of sequence similarity using local BLAST: [1] comparing nucleotide transcript sequences to nucleotide sequences from *D. melanogaster* (blastn), [2] comparing translated nucleotide sequences to protein sequences of *D. melanogaster* (blastx), [3] comparing nucleotide sequences to a database of nucleotide sequences from *D. melanogaster*, *D. virilis*, and *D. grimshawi* (blastx), and [4] comparing translated nucleotide sequences to a database of protein sequences from the same three species (blastn). For downstream analyses,

we prioritized annotations from the second comparison, but we provide all sequence similarity reports in the GitHub repository under the directory analysis/BLAST.

To annotate homolog groups as defined by the homology inference step of agalma, we extracted the name and sequence ID from all *D. melanogaster* sequences in the group.

### 221 3.8 Normalization and differential gene expression

Transcript count tables were imported into R using the agalmar package, version 0.0.0.9000. Differential gene expression analysis was performed using the package DESeq2, version 1.32.0. For these analyses we used only one sequencing run per library, thereby excluding duplicate sequencing runs. Analyses of differential gene expression were calculated using the default approaches in DESeq2 for estimating size factors, dispersions, and calculating log<sub>2</sub> fold-change and p-values (Fig. S2A). Both individual and tissue were considered in the design formula. Transcripts were considered differentially expressed at a significance threshold of 0.01.

We identified a cohort of core ovary-specific genes by first identifying a parent gene for each transcript using 228 a sequence similarity search against D. melanogaster (Fig. S2A). We then identified parent genes that had 229 at least one transcript significantly differentially upregulated in the ovary of more than ten of the twelve 230 species. Because multiple transcripts may match to a single parent-gene, core ovary-specific parent genes 231 may include transcripts that are also not differentially upregulated in the ovary, as long as at least one 232 transcript is for more than ten out of twelve species. This may be the case when transcripts are artificially 233 fragmented during reference transcriptome assembly, or when sequence-similar transcripts have biologically 234 distinct expression levels. 235

### <sup>236</sup> 3.9 Comparison of expression to *D. melanogaster*

We compared our differential gene expression results to a reference database of tissue expression from D. 237 melanogaster, known as the FlyAtlas2<sup>29</sup>. We downloaded this reference in July of 2021, from http://motif. 238 gla.ac.uk/downloads/FlyAtlas2 21.04.18.sql. This dataset provides data on transcript abundance and tissue 239 enrichment, including for female ovaries. Tissue enrichment is calculated using the same methods as in the 240 FlyAtlas2 web browser, defined as the fragments per kilobase of transcript per million mapped reads (FPKM) 241 for a given tissue divided by that value for the reference tissue (here, female whole body), with a pseudocount 242 of two counts added to empty values to avoid division by zero. We considered a FlyAtlas gene to be enriched 243 244 in the ovary, comparable to our data, if the ovary was the maximum enrichment value across all tissues excluding the head, brain, and eye tissues, as these were separated in our RNASeq procedure (Fig. S2A). 245 We considered a FlyAtlas gene to be head enriched if either the head, brain, or eye were the maximum 246 enrichment value, excluding the ovary. 247

# 3.10 Transforming data into comparable measurements of expression across species

Transcript counts are reported in transcripts per million (TPM), but this measurement is known to not be directly comparable across species due to differences in reference transcriptome size<sup>7,8</sup>. Therefore, we normalized TPM by species using the procedure described by Munro and colleagues (2021)<sup>8</sup>, where TPM values are multiplied by the number of genes in the reference, and this value is divided by 10<sup>4</sup> (Fig. S2B). TPM10k values were natural-log transformed.

An additional challenge when working with reference transcriptomes is the presence of fragmented transcripts created during the assembly process<sup>58</sup>. This fragmentation can result in noise in estimating the amount of transcript as reads are differentially mapped to these fragments. To reduce the impact of this noise on our analysis, we undertook a novel approach where transcripts were grouped according to inferred homology as estimated by the agalma pipeline using an all-by-all BLAST approach (Fig. S2B). For each sequenced library, we then found the average count value across all transcripts from the same homology group (see

Table S5 for statistics on homology group composition). For each species-tissue pair, we then averaged this value across all biological replicates, here replicate individuals.

### <sup>263</sup> 3.11 Linear modeling

We performed linear modeling to calculate the relative contribution of tissue- and species-level differences 264 to variation in gene expression (Fig. S2B), following the approach of Breschi and colleagues  $(2016)^{21}$ . These 265 analyses were performed separately on datasets of ovary vs. carcass and head vs. carcass expression. Using 266 the ANOVA script provided at https://github.com/abreschi/Rscripts/blob/master/anova.R, we built 267 a linear model for each gene that accounts for the contribution of the organ, species, and any residual error. 268 We then calculated the relative proportion of each factor divided by the total sum of squares for all factors. 269 We identified groups of highly variable genes, using the same metrics defined by Breschi and colleagues 270  $(2016)^{21}$ , as any gene for which either tissues or species explains at least 75% of the variance. Species 271 variable genes (SVGs) were defined as highly variable genes whose relative variation was two-fold greater 272 across species than tissues (vice-versa for tissue variable genes, TVGs). 273

We performed these linear model analyses over four nested clades: a clade containing two *picture-wing* species (*D. sproati* and *D. macrothrix*); a clade containing the four *picture-wing-Nudidrosophila-Ateledrosophila* species in this study; a clade containing the nine Hawaiian *Drosophila* species in this study; and a clade of all 12 Hawaiian *Drosophila* and *Scaptomyza* species in this study. We repeated these analyses excluding the species *S. varia*, which showed the lowest similarity in expression to the other eleven species. To compare our analysis to the more typical approach undertaken, we also performed these analyses on all pairwise combinations of these twelve species.

#### <sup>281</sup> 3.12 Reconstructing evolutionary history of differential expression

We calculated tissue bias as the ratio of counts in TPM10k for each tissue (ovary and head) to the reference tissue<sup>7</sup>, here the carcass (Fig. S3A). We subsequently performed the same transformation steps described above, averaging over ratios from the same homology group and across biological replicates, to calculate average expression bias per homology group per library. To avoid division by zero, we added a pseudocount of 0.01 to each TPM10k value. Ratio values were natural-log transformed so that positive values indicate enrichment in the tissue of interest relative to the reference tissue, negative values indicate the opposite, and values of zero indicate equivalent expression.

We reconstructed the evolutionary history of tissue bias for each homology group using the species tree published in Church and Extavour, 2021<sup>44</sup>, based on the same reference transcriptome data (Fig. S2C). First, we calibrated the tree estimated using IQtree (Fig 1A of that publication) to be ultrametric using the R function chronos in the package ape, version 5.5 (using a correlated model and a lambda value of 1). We then subset this tree to only include tips for which expression data was available, and annotated this tree to be able to identify specific branches and nodes in ancestral state reconstruction analyses.

Ancestral expression bias values were estimated with the R package Rphylopars, version 0.3.2, using the fast 295 ancestral state reconstruction algorithm based on Ho and Ané, 2014<sup>59</sup> (Fig. S3A). Tips for which expression 296 data were not available were dropped from each reconstruction, and ancestral state reconstruction was only 297 performed when more than 3 tips had data. Following ancestral state reconstruction, we calculated the 298 scaled change as the difference between the value at the ancestral and descendant nodes, divided by the 299 length of the branch. Scaled changes were compared between homology groups by identifying equivalent 300 branches as those that share the same parent and child node, following the procedure described in Munro 301 and colleagues  $(2021)^8$ . We identified qualitative changes in expression bias as changes that resulted in a 302 ratio changing from negative to positive values or vice versa. 303

### <sup>304</sup> 3.13 Estimating correlated evolution of expression across genes

<sup>305</sup> For each homology group that had representation across all twelve species, we calculated pairwise Pear-

<sup>306</sup> son's correlation coefficients by comparing scaled changes in expression bias across equivalent branches (Fig.

<sup>307</sup> S3B). For the twelve-species phylogeny, this meant each correlation coefficient was calculated using 22 in-

dividual data points (branches). This resulted in a correlation matrix of 1,306,449 pairwise comparisons of

<sup>309</sup> evolutionary correlation.

We compared this correlation network to data on protein interactions and genetic interactions downloaded from http://flybase.org in July, 2021. These data include pairwise observations of genetic enhancement and suppression interactions between parent genes in *D. melanogaster*. These interactions were matched to pairwise correlation coefficients by identifying the corresponding homology group for each *D. melanogaster* 

<sup>314</sup> parent gene ID (more than one parent gene may fall into the same homology group).

We tested whether correlation coefficients for known genetic interaction partners were higher than in genes with unknown interactions using two-sample t-tests. In each test we compared the coefficients for either enhancement or suppression interactions to a random sample of 5000 coefficients for which interactions are unknown. We repeated these t-tests 100 times using different random samples, and report the maximum p-value observed. We also compared the distribution of enhancement and suppression interaction coefficients to each other using a single t-test.

Strong correlations for the visualization of co-evolutionary networks were selected using a threshold correlation coefficient of 0.825.

## 323 4 Data Availability

All data, results, and code for this manuscript are available at GitHub, under the repository 324 shchurch/hawaiian\_drosophilidae\_expression\_2021, commit 67d8e6f. The code to perform all 325 agalma commands was performed in clean anaconda environment, installed following the instructions 326 at https://bitbucket.org/caseywdunn/agalma. All R commands were performed with a fresh install 327 of R, and the session information including all package versions is available in the GitHub repository 328 under the file r\_session\_info.txt. The code to generate all plots as well as the text of this manuscript 329 is available in several R scripts and Rmarkdown files at the same location. The resulting correlation 330 matrix can be interactively visualized and queried at the accompanying data visualization for this paper 331 (https://github.com/shchurch/hawaiian fly dataviz 2021). 332

## 333 5 Results

## 5.1 Differential gene expression reveals a cohort of consistently ovary-specific genes

We observed several patterns in tissue-specific gene expression that are consistent across all twelve species. 336 First, in all species the main axis of variation separated ovary RNA libraries from head and carcass (Fig. 337 S4). In all species this axis accounted for at least 50% of variation, and in several species greater than 70% of 338 variation. To test for possible variation due to different runs on the sequencer, we resequenced several libraries 339 and compared them using principle component analysis. We found variation between sequencing runs to be 340 negligible compared to variation across tissues and individuals (Fig. S7). Second, in all species we observed 341 that there was a larger amount of significantly downregulated transcripts than upregulated in the ovary 342 relative to the carcass (Fig. 2A-B, S5). Across species, we observed an average of 27.7% to be significantly 343 downregulated and 15.5% of transcripts to be significantly upregulated. In contrast, when comparing the 344 head to the carcass, we observed an average of 10% of transcripts to be significantly upregulated and 10.5%345 to be significantly downregulated (Fig. S6). Therefore the ovary shows a larger number of both upregulated 346

 $_{347}$   $\,$  and downregulated genes relative to the carcass than the head, indicating the ovary has a particularly distinct

expression profile. These differences may also reflect variation in the complexity and diversity of functions

<sup>349</sup> of the tissues being compared.

We used the results of our differential gene expression analysis within species to test for the existence of a suite of genes that show consistent ovary-specific expression across species. We found a cohort of 131 genes, grouped according to BLAST sequence similarity to *D. melanogaster*, for which at least one transcript was significantly upregulated in the ovaries of more than ten species (Fig. 2C). Transcripts matching these genes made up on average 24.6% of the significantly ovary-upregulated transcripts across species, indicating that a substantial portion of ovary-specific genes have conserved expression patterns over evolutionary time (17.7%,

 $_{356}$  excluding the species *S. varia* that had the most distinct expression profile of all species).

We then tested the extent to which these core ovary genes correspond to observations in well-studied laboratory *Drosophila* models. To accomplish this, we compared expression across Hawaiian species to reported tissue-specific expression levels from *D. melanogaster*<sup>29</sup>. We found that Hawaiian core ovary-specific genes show nearly universal enrichment in the ovary of *D. melanogaster* as well, as reported in the FlyAtlas2 dataset<sup>29</sup> (Fig. 2D). We likewise observed that genes reported in *D. melanogaster* to have highest enrichment in the ovary largely correspond to genes that are significantly upregulated in the ovaries of Hawaiian species (Fig. S8).

<sup>364</sup> The 131 core ovary genes include several well-known members involved in oogenesis and germline stem cell

renewal such as  $nanos^{34}$ ,  $swallow^{35}$ , and  $oskar^{60}$  (Fig. 2E). We found only two genes that were identified as

Hawaiian core ovary genes that are not reported in the FlyAtlas2 dataset<sup>29</sup> to be enriched in the ovary of D.

 $_{367}$  melanogaster: the SET domain binding factor sbf, and Rfx, which are reported to be enriched in the heart,

 $_{368}$  brain, and other non-reproductive tissues<sup>29</sup>.

We used the same approach to identify a core suite of 52 head-specific genes (Fig. S9). There was no 369 overlap between the sets of core head genes and core ovary genes. To test whether the correspondence 370 between expression observations in Hawaiian flies and *D. melanogaster* might be due to factors beyond 371 tissue identity, we compared head expression values to ovary enrichment data from D. melanoqueter, as we 372 had done for ovary expression values above. We did not observe a correspondence in either direction between 373 expression in the head of Hawaiian species and enrichment in the ovary of D. melanogaster (Fig. S10A). 374 In contrast, we did find a correspondence between head-specific expression and genes enriched in the D. 375 melanogaster brain, eye, and head (Fig. S10B). Core head genes include Rhodopsin photoreceptor genes and 376 genes such as *hikaru genki* with involvement in synaptic centers<sup>61</sup>. 377

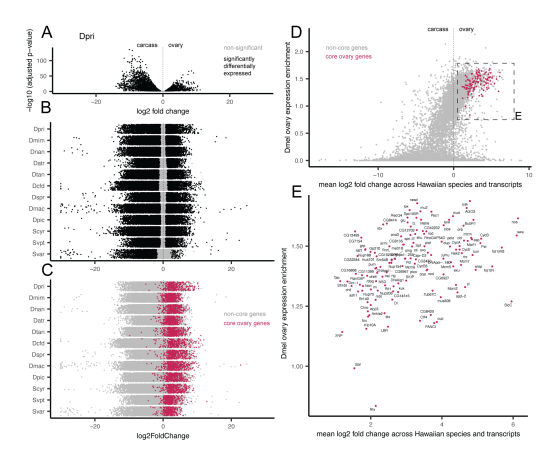


Figure 2: Identifying a cohort of ovary-specific genes across drosophilid species. A, Volcano plot for one example species, *D. primaeva* (Dpri), showing the results of a differential gene expression analysis comparing the ovary to the carcass. The x-axis shows the  $\log_2$  fold change of expression across transcripts, and the y-axis shows the adjusted p-value,  $\log_{10}$  transformed. Points that are significantly differentially expressed are shown in black. B, Jitter plots showing the results of the same analysis across the twelve species studied here. The x-axis shows the  $\log_2$  fold change of expression across transcripts, and points are arranged with random jitter within species on the y-axis. C, The same jitter plots, but now colored according to whether or not transcripts belong to a cohort of core ovary genes. These are defined as genes, grouped by BLAST similarity to *D. melanogaster* transcripts, for which at least one transcript is upregulated in the ovary of ten or more of the twelve species. D, A comparison of mean expression change across Hawaiian species to reported ovary-enrichment values from *D. melanogaster*, as reported in FlyAtlas2<sup>29</sup>. Core ovary genes are marked in magenta. E, The same plot, now showing only core ovary genes, annotated with the gene symbol from *D. melanogaster*.

### 5.2 Modeling reveals the phylogenetic decay of expression similarity between tissues

Many studies have investigated the question of whether we expect expression to be more similar across the same organ in different species, or across different organs within the same species<sup>13-20</sup>. Recent studies have suggested that the answer to this question will depend on the phylogenetic distance separating the species being compared<sup>21</sup>. Here we used a modeling approach to investigate this question with respect to the ovaries of Hawaiian drosophilids.

First, we determined an appropriate unit of comparison across species, based on an assessment of homologous features between reference transcriptomes. The agalma pipeline provides a method for determining

homologous and orthologous sequences using an all-by-all BLAST approach to determine clusters of recipro-387 cally similar sequences (homology groups). These can then be divided into orthology groups by estimating 388 genetrees and identifying maximally inclusive subtrees with no more than one sequence per taxon<sup>55</sup>. We 389 compared the representation of species across homology and orthology groups, and observed that while the 390 representation of homology groups increases with the number of species compared, representation of or-391 thology groups decreases (Fig. S11). This is a known obstacle in comparative transcriptomics, attributed 392 to many transcripts being artifactually fragmented during reference transcriptome assembly<sup>58</sup>. To reduce 393 the impact of this on our downstream analyses, we averaged TPM values across all transcripts within a 394 homology group for each sequenced RNA library. Principle component analysis of this average expression 305 dataset showed that the first principle component divides ovary libraries from the rest, while the second com-396 ponent separates samples along an axis that largely corresponds to phylogenetic distance between species 397 (Fig. S12). While this averaging approach reduces noise due to variable mapping affinities of fragments of 398 the same transcript, it comes at the cost of averaging over potential variation between genuine transcripts 399 that fall into the same homology group. Future analyses using improved assemblies for transcriptomes or 400 genomes will likely be able to avoid this trade off and compare transcript counts directly. 401

With average expression counts for homologous transcripts across species, we tested the degree to which 402 variation across this dataset could be attributed to tissue-specific variation (here, ovary vs. carcass), species 403 specific variation, or neither (residual variation). Using the linear modeling approach adapted from Breschi 404 and colleagues  $(2016)^{21}$ , we found the proportion of variance across the dataset attributed to tissue differences 405 decreased with phylogenetic distance, while the proportion attributed to species difference increased (Fig. 406 3A-C). In addition, we found that, when comparing ovary and carcass tissues, the Hawaiian drosophilid clade 407 encompasses the crossover point where variation across species swamps variation across tissues (crossed lines, 408 Fig. 3A). When comparing across the two species from the *picture-wing* group included in this study, an 409 average of 45.6% of the variation can be attributed to tissue differences. For the same comparison, 960 genes 410 were identified as tissue variable genes, defined as residual variation accounting for <25% and a two-fold 411 increase in variation attributed to tissues than to species (Fig. 3B, S13). In contrast, when comparing across 412 all twelve Hawaiian drosophilid species studied here, 34.7% of the variation can be attributed to tissue, with 413 240 TVGs (Fig. 3B, S13). Across different clades of comparisons, the number of species-variable genes 414 (SVGs) remains relatively stable (from 304 to 260, Fig. 3B). 415

We then leveraged the results of this linear modeling approach across all twelve species to perform an additional screen for genes that are consistently upregulated in ovaries across species. We compared the proportion of variation explained by tissue for each homology group to the average log<sub>2</sub> fold change from the results of our differential gene expression analysis (Fig 3D). This comparison allowed us to identify genes that fall above our threshold for TVGs that are also upregulated in the ovary (Fig 3E). This group of genes includes many of the same members as the core ovary genes (e.g. *nanos* and *swallow*), as well as several new candidates (e.g. *singed*).

To test the importance of tissue identity, we repeated the same analysis comparing variation across species and tissues using the head in place of the ovary. Consistent with what we describe for the ovary and carcass, as phylogenetic distance increases the proportion of variation across tissues decreases while variation across species increases. In contrast to the above findings, however, for the head and carcass far less of the variation in gene expression can be attributed to tissue differences (Fig. S14). For these tissues, the crossover point between total proportion of variation occurs roughly at the distance separating the two *picture-wing* species.

To verify these results were not driven by the species *S. varia*, which had the most distinct expression patterns of all species, we repeated these analyses excluding this species and recovered largely equivalent results (Fig. S15). To compare our findings to those that would be recovered using a more typical pairwise approach, we repeated the linear modeling analysis on ovary and carcass data using every pairwise combination of the twelve species. We recovered the same trend of decreasing contribution of tissue-level variation with increasing phylogenetic distance, and observed that the variance in mean proportion attributed to either species- or tissue-level differences increased as well (Fig. S16).

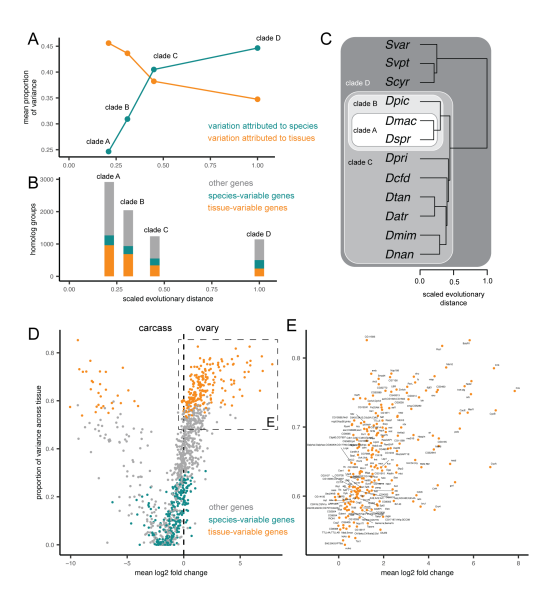


Figure 3: Linear modeling shows the proportion of variance explained by differences across tissues and species. A, The results of a linear modeling approach to calculate expression variation for each gene, attributed to variation across organs, species, or residual variation, as described in Breschi and colleagues  $(2016)^{21}$ . The average proportion of variation attributed to tissues is higher than that attributed to species for the two *picture-wing* species in clade A, while the opposite is true for all twelve species in clade D. B, The number of genes, defined by homology group, classified as tissue variable genes (TVGs), species variable genes (SVGs), or neither in each clade comparison. C, The phylogeny of the twelve species studied here, showing the four clades compared in A-B. Scaled evolutionary distance is calculated as the relative distance from the most recent common ancestor of Hawaiian drosophilids to extant species. D, Comparing approach on the y-axis (variation across tissues). Genes are colored according to TVGs and SVGs. The inset box highlights TVGs that are upregulated in the ovary relative to the carcass. E, The same plot, now showing only upregulated TVGs, annotated with the gene symbol from the *D. melanogaster* sequences in the same homology group.

### <sup>436</sup> 5.3 Identifying gains and losses of ovary bias across genes and the phylogeny

While a substantial fraction of ovary-specific transcripts belong to the cohort of core ovary genes, on average 437 75.4% of transcripts are upregulated in the ovaries of one or several species, but not consistently across ten or 438 more of the species studied here (Fig 2B-C). This is suggestive of many evolutionary gains and losses of ovary 439 biased expression of genes. We characterized the evolution of these gains and losses using an ancestral state 440 reconstruction approach. First we quantified expression bias between tissues as the ratio of read counts<sup>7</sup>, 441 then reconstructed the value of this continuous trait for each gene (defined using homology groups) at each 442 node of the estimated species tree (Fig. S3). We then calculated the scaled change of expression bias along 443 each branch, which allowed us to describe how relative expression values between tissues had changed the 444 course of evolutionary time (Fig 4A). Visualizing the distribution of scaled changes by genes shows that 445 most scaled changes are small and centered around zero, representing little change in gene expression bias 446 between tissues (Fig. 4B-C). 447

Using this dataset of scaled changes across genes and branches, we identified branches for which the direction 448 of tissue bias had changed (e.g. from higher expression in the ovary biased than the carcass to lower, or vice 449 versa). Visualizing this dataset according to branches reveals that the majority of these changes in bias are 450 located on the root and terminal branches, rather than internal branches (Fig 4D-E). This is likely because 451 internal branches for this rapid radiation tend to be very short; even when scaling evolutionary changes to 452 branch length, it is less probable to experience a shift to and from ovary biased on a short branch than a 453 long branch. Visualizing the distribution of genes by ancestral and descendant values allows us to identify 454 shifts in bias which represent the largest swings in expression values (Fig. 4F, points a-d). Highlighting the 455 top two such shifts in both directions, we identify four example genes which acquired or lost ovary-specific 456 expression in the phylogeny of Hawaiian Drosophilidae. In the case of FMRFaR and GABA, a few Hawaiian 457 species have gained ovary biased expression of these genes, while most species and the ancestral state indicate 458 non-ovary bias (Fig. 4Fa-b). In the case of *vilya* and the unnamed gene CG9109, each shows a pattern where 459 one species has lost ovary bias from a biased ancestral state (Fig. 4Fc-d). 460

Repeating the same analysis using the head in place of the ovary revealed a set of evolutionary gains and
 losses in head-specific expression (Fig. S17). Identifying the top four changes in head expression shows gains

and losses of head expression in the genes *hiro*, *stil*, *Jhe*, and, consistent with the ovary, *vilya*. In the case of

the latter, these results may be driven by substantial changes in expression of vilya in carcass tissues across

species, resulting in major differences in both ovary and head-biased expression.

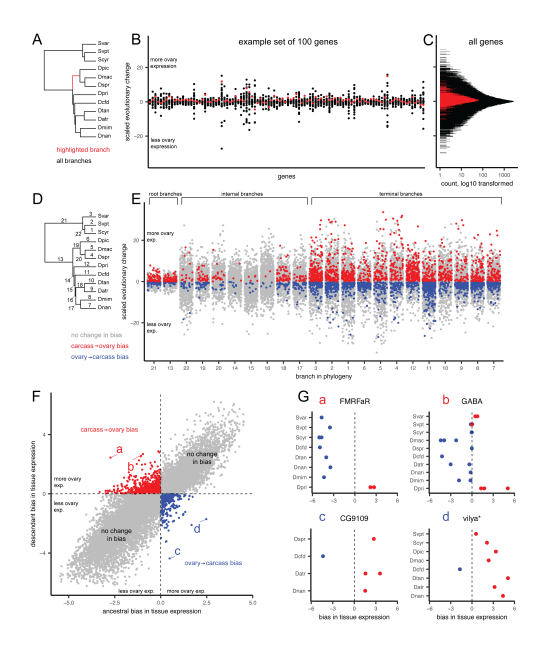


Figure 4: Identifying genes that have gained and lost ovary expression across the phylogeny. A, The phylogeny of the twelve species studied here, highlighting one example branch of the 22 for which we inferred the scaled evolutionary change in expression bias. B, The distribution of changes, grouped by gene, for 100 randomly selected genes, defined by homology group. Each point represents one of the 22 branches from A, with the red point corresponding to the highlighted branch from that panel. C, The distribution,  $log_{10}$  transformed, of scaled genes across all branches and all genes. Changes on the highlighted branch in red. D, The phylogeny with all 22 branches numbered. E, The distribution of changes, grouped by branch, with random jitter on the x-axis within each group. Points colored according to the qualitative change in bias, either from more expression in ovary than carcass to less (blue), the reverse (red), or no change in overall bias (gray). F, The distribution of ancestral and descendant values, showing the two quadrants that represent qualitative changes in bias. Points that represent large swings in expression within those quadrants are labeled a-d. G, The four genes with large swings from F, showing the expression bias for each transcript colored according to more expression in the ovary (red) or carcass (blue). Panels annotated with the gene symbol from the *D. melanogaster* sequences in the same homology group, with the exception of *vilya*\*, which was annotated using a direct BLAST search since no *D. melanogaster* sequence was present in that group.

### 466 5.4 Genes with a strong correlation of expression evolution

We tested the estimated evolutionary changes in expression bias for evidence of correlated expression evo-467 lution between genes. For every gene represented across all species, we performed a pairwise comparison of 468 changes in expression bias, using as data points the scaled change in ovary bias on the 22 branches in the 469 phylogenetic tree. This resulted in 1,306,449 pairwise measures of evolutionary correlation between genes. 470 Because the number of gene pairs being compared is much larger than the number of values used to esti-471 mate correlation, this method has the potential to produce many spurious correlations<sup>7</sup>. To test the degree 472 to which the correlations observed here reflect known biological interactions between genes, we compared 473 these measures to reported protein and genetic interactions between genes, using the database of published 474 genetic experiments in *D. melanogaster*, available at http://flybase.org. We found that the mean correlation 475 coefficient for genes that are known to physically interact as proteins was higher than for genes with no or 476 unknown interaction (p-value=<0.001, Fig. 5A). This indicates that even with a relatively small number 477 of observations, there is sufficient information in the matrix to detect biological signal between gene pairs. 478 These results were calculated based on the correlation in expression bias between the ovary and carcass. 479 However, following the same procedure using correlations in changes in head-biased expression showed no 480 significant difference between the two groups (p-value=0.256, Fig. S18), suggesting the strength of this signal 481 may be dependent on the tissues being compared. 482

We also found that genes known to interact via genetic enhancement or suppression have a significantly 483 higher mean correlation than genes with no or unknown genetic interactions (unknown vs. enhancement 484 p-value < 0.001, unknown vs. suppression = < 0.001, Fig. 5B). Comparing genes with known enhancement and 485 suppression interactions to each other showed no significant difference (p-value=0.497). However, for genetic 486 interactions, the range of correlation coefficients was higher in the group of no or unknown interactions (Fig. 487 5B). This indicates that, while the average correlation of expression evolution might be higher for interaction 488 partners, stronger positive and negative correlations exist between pairs of genes which do not interact, or 489 for which interactions have not yet been tested. 490

As evidence of this, we compared the interaction network for known ovary-specific genes, to test whether the 491 network inferred based on strong correlation of expression evolution was consistent with known interaction 492 partners from D. melanogaster. We selected as an example the gene volk-protein gene family, which are 493 known to be expressed in the reproductive system, among other tissues<sup>62</sup> (Fig. 5C). We found 8 distinct 494 homologous gene groups, comprising 14 unique D. melanogaster parent genes, that had a strong evolutionary 495 correlation with volk-protein genes (absolute coefficient greater than 0.825, Fig. 5D). None of these correlated 496 genes correspond to those listed on  $FlyBase^{63}$  as having known interactions with yolk-protein genes in D. 497 melanogaster (Fig. 5E). We consider the strong evolutionary correlations to be a set of new predictions about 498 evolutionary and genetic relationships between genes which can be tested in wild and laboratory model 499 species of *Drosophila*. The dataset of pairwise correlation coefficients can be visualized and interrogated 500 at the accompanying data visualization for this manuscript (https://github.com/shchurch/hawaiian fly 501 dataviz 2021). 502

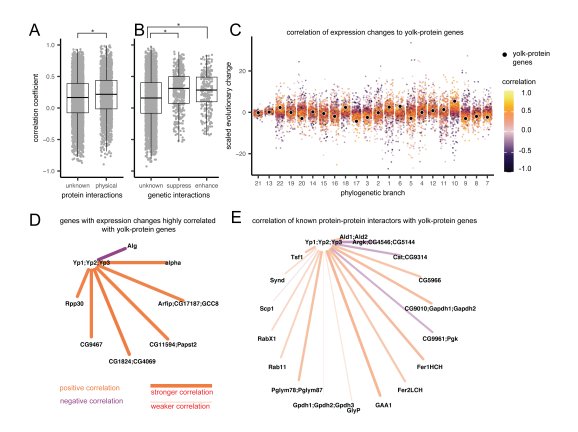


Figure 5: Estimating pairwise correlation coefficients across genes reveals new networks of correlated expression evolution. A-B Comparison of the distribution of Pearson's correlation coefficients based on ovary-biased expression evolution between genes. Asterisks indicate a significant t-test comparison. A, Genes with no or unknown protein-protein interactions compared to those with reported interactions in FlyBase<sup>63</sup> (p-value=<0.001). B, Correlation comparison between genes with no or unknown genetic interactions and those reported to have enhancement or suppression interactions in FlyBase (unknown vs. enhancement p-value=<0.001; unknown vs. suppression=<0.001; enhancement vs. suppression=0.497). C, Each point represents a scaled change in expression bias, colored by Pearson's correlation coefficients relative to one example gene-family, the yolk-protein genes (black points), arranged by phylogenetic branch (numbers shown in Fig. 4D). Yellow=strong positive correlation, purple=strong negative correlation. D, The network of strong correlation. Stronger correlations are shown by brighter colors, and thicker, shorter lines. Nodes are annotated with the gene symbols from the D. melanogaster sequences from that homology group. E, The correlation between known protein-protein interaction partners<sup>63</sup> with the yolk-protein genes.

## 503 6 Discussion

The results of this study show the importance of placing any comparison of gene expression across species 504 in an evolutionary context. When making comparisons that involve model organism for the study of 505 development and disease, this means identifying the crossover point at which variation between species 506 begins to swamp variation across the tissues or treatments in question. In such comparisons, the possibility 507 that any individual gene may show a divergent pattern of expression from the model organism increases 508 substantially. This study provides evidence that confirms we should expect variation in gene expression to 509 increase with the phylogenetic distance separating the species being compared. In addition, our results using 510 ovary and head expression data show that our expectation should also depend on the identity of the tissues 511

<sup>512</sup> being compared. Our dataset demonstrates that for some tissues, like the fly head, this crossover point may <sup>513</sup> be met even when comparing between two relatively closely related species.

Despite substantial variation across species, we describe here the composition of core suites of ovary- and 514 head-expressed genes that have maintained conservation of expression patterns over millions of years of 515 evolution. The core ovary genes include some of the most well-studied genes in relation to oogenesis, such 516 as nanos and oskar, as well as many genes that have yet to be studied in depth (e.g. unnamed genes such 517 as CG3430). We provide the full list of core ovary and head genes as a reference against which future 518 genetic studies may be informed and compared (Tables S6-S7). Furthermore, the existence of these suites of 519 genes suggests that equivalent groups are likely to exist within the many gene expression atlases currently 520 being published<sup>64,65</sup>. New technologies such as single-cell RNA sequencing that use global signatures of gene 521 expression to identify cells are ripe for interspecific comparisons that may reveal evolutionary conserved 522 gene modules<sup>66</sup>. Developing robust comparative methods for comparing these atlases across species has 523 the potential to reveal ancestral expression patterns in cells and organs, as well as pinpoint important 524 evolutionary shifts in expression regulation. 525

Our results indicate that genes known to interact, both physically as proteins and through genetic enhance-526 ment and suppression, likely experience more correlated changes in expression than would be expected for 527 genes chosen at random. However, we also find the difference in mean correlation between these groups to 528 be relatively small, and dependent on the context of the tissue in question. One possible explanation for 529 this finding is that interactions between genes with strong correlations of expression evolution have yet to be 530 described. We provide an interactive tool to explore highly correlated genes that can inform future genetic 531 studies in *D. melanogaster* and other related species (https://github.com/shchurch/hawaiian\_fly\_dataviz\_ 532 2021). Another possibility we consider likely is that interactions between genes represent one factor among 533 many that dictate the probability of correlated changes in expression. We hypothesize that other features, 534 such as shared regulatory architecture, will also influence evolutionary correlation of expression. 535

As more studies undertake phylogenetic comparisons of functional genomic data, new factors that influence 536 the evolutionary associations between genes are likely to be revealed<sup>7</sup>. The strength of these phylogenetic 537 comparisons will depend in part on comparing across a sufficient number of taxa such that there are multiple 538 branches on which to calculate and compare evolutionary changes. However, even as functional genomic 539 data become more accessible for more species, the number of features being compared (e.g. thousands 540 of genes) will likely continue to outnumber the number of evolutionary observations (e.g. changes along 541 branches)<sup>7</sup>. One encouraging result from this study is that, using our matrix of gene expression changes 542 along 22 branches, there is sufficient information to detect the biological signal associated with physical and 543 genetic interactions. While this is true, we assume that some fraction of the correlations that we report here 544 represent false positives, and that the strength of correlation of these genes would decrease with the addition 545 of more taxa to the comparison. For this reason we present the correlation matrix as a set of hypotheses to 546 be tested in future studies using additional lines of evidence. 547

One outstanding challenge in expression evolution is the quality of the references available against which 548 RNA reads can be mapped<sup>58</sup>. In this study we account for the statistical noise in our data by averaging 549 expression values over groups of homologous genes, as identified by sequence similarity to high quality refer-550 ence genomes. This approach has the advantage of accounting for problems associated with fragmentation of 551 genes in transcriptome assembly. However, it comes at the cost of averaging over possible biological variation 552 in expression between genes from the same gene family. The strong concordance of our results with published 553 records from D. melanogaster suggests that the approach we have used here is robust for our dataset. How-554 ever, as the quality and accessibility of genomes from diverse species continue to increase, future studies will 555 likely be able to compare directly between orthologous genes without needing to account for fragmentation. 556 For those future studies, a phylogenetic comparative approach like the one used here and  $elsewhere^{8}$  can 557 serve as an analytical framework to move expression comparisons beyond pairwise comparisons. 558

A goal of evolutionary-developmental biology is to identify changes in developmental mechanisms that underlie phenotypic differences<sup>12</sup>. Many evo-devo studies approach this by identifying phenotypic variation between species and then searching for differences in gene content or expression using one or several emerging model organisms in the lab<sup>12</sup>. To narrow down the field of search, this approach often requires outside knowledge of candidate genes, gained from developmental research in related models or other methods of

<sup>564</sup> filtering the genome. Furthermore, because these approaches usually lack global measurements of gene ex-

<sup>565</sup> pression variation across species, identifying an expression difference does not always constitute a smoking

<sup>566</sup> gun<sup>6</sup>. For example, observing a difference between candidate gene expression would not be unexpected if we <sup>567</sup> frequently observe differences of that magnitude between genes chosen at random. An alternative approach,

<sup>567</sup> frequently observe differences of that magnitude between genes chosen at random. An alternative approach, <sup>568</sup> as demonstrated here, is to characterize all the evolutionary changes in expression across the transcriptome,

and then identify the changes that are significantly associated with traits of interest<sup>9</sup>. As expression data

<sup>570</sup> become available from an ever wider array of species, this "evolutionary screen" approach becomes increas-

<sup>571</sup> ingly possible. One advantage of this approach is that it may reveal associations that would otherwise

escape detection when comparisons are centered on model organisms; for example, when genes, traits, or

<sup>573</sup> processes happen to not be present in our laboratory model species<sup>10</sup>. By leveraging phylogenetic compara-

<sup>574</sup> tive methods on high-dimensional functional genomic data, the objective of connecting genomic variation to

<sup>575</sup> developmental mechanisms and phenotypic differences will be accelerated.

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