

Postmating transcriptional changes in reproductive tracts of con- and heterospecifically mated *Drosophila mojavensis* females

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In internally fertilizing organisms, mating involves a series of highly coordinated molecular interactions between the sexes that occur within the female reproductive tract. In species where females mate multiply, traits involved in postcopulatory interactions are expected to evolve rapidly, potentially leading to postmating-prezygotic (PMPZ) reproductive isolation between diverging populations. Here, we investigate the postmating transcriptional response of the lower reproductive tract of *Drosophila mojavensis* females following copulation with either conspecific or heterospecific (*Drosophila arizonae*) males at three time points postmating. Relatively few genes (15 total) were differentially regulated in the female lower reproductive tract in response to conspecific mating. Heterospecifically mated females exhibited significant perturbations in the expression of the majority of these genes, and also down-regulated transcription of a number of others, including several involved in mitochondrial function. These striking regulatory differences indicate failed postcopulatory molecular interactions between the sexes consistent with the strong PMPZ isolation observed for this cross. We also report the transfer of male accessory-gland protein (Acp) transcripts from males to females during copulation, a finding with potentially broad implications for understanding postcopulatory molecular interactions between the sexes.

gene expression | reproduction | sexual selection | sexual conflict | speciation

In internally fertilizing organisms, the female reproductive tract serves as the arena for a series of highly coevolved molecular interactions between the sexes that are critical for successful reproduction (1, 2). Postcopulatory interactions should further increase in complexity in species in which females mate with more than one male, as intense sexual selection propels the rapid evolution of traits mediating female choice, male competitive ability, and sexual conflict (3, 4). This, in turn, may facilitate divergence of such traits between populations following different coevolutionary trajectories, leading to postmating-prezygotic (PMPZ) reproductive isolation (5). Consistent with these expectations are the rapid evolution of morphological and molecular reproductive traits associated with postcopulatory processes (6) and the recognition that PMPZ barriers can serve as potent and rapidly evolving forms of reproductive isolation (5).

The availability of genomic resources for an increasing number of species provides a platform for elucidating the molecular basis of postcopulatory molecular interactions between males and females. For example, recent genomic studies on *Drosophila melanogaster* (7–14), *Anopheles gambiae* (15), and *Apis mellifera* (16, 17) have begun to characterize the female postmating response by identifying changes in the transcriptome and/or proteome of mated females. In *D. melanogaster*, sperm or other specific components of the seminal fluid are known to induce some of these changes, which ultimately trigger physiological responses in females (18). Male accessory-gland proteins (Acps), in particular, modulate a variety of physiological processes in *D. melanogaster* females including immune response, oogenesis, oviposition, sperm transfer and storage, and female receptivity

(18). Although considerable progress has been made in understanding the nature and scope of postcopulatory molecular interactions between males and females, comparable studies on additional species, especially those with different mating systems, are necessary to generalize these findings. Moreover, although accumulating evidence suggests that postcopulatory incompatibilities between the sexes often result in significant PMPZ reproductive isolation between species (5), the molecular and genetic bases of such incompatibilities have yet to be identified.

Drosophila mojavensis and *Drosophila arizonae* are recently diverged (<1 Mya) sister species (19) with partially overlapping distributions in the arid regions of southwestern United States and northwestern Mexico. The mating systems of these two species are characterized by frequent female remating relative to *D. melanogaster* (20), along with extensive intersexual coevolution of postcopulatory traits (21–23), including rapid evolution of both male and female reproductive proteins (24–28). Consistent with expectations, interspecific crosses also exhibit strong PMPZ isolation, particularly those involving *D. mojavensis* females. Heterospecifically mated *D. mojavensis* females exhibit perturbations in a number of processes occurring within the female reproductive tract that result in a high incidence of failed fertilizations, a reduced rate of oviposition, and ultimately the production of few hybrid offspring (25). These problems are associated with deficiencies in the heterospecifically mated female's storage and retention of sperm, and also in degrading the insemination reaction, a temporary mass that forms in the uterus immediately after conspecific copulation (25). In contrast to conspecific matings, where the mass is typically eliminated within several hours, following heterospecific matings the mass often persists for days, interfering with oviposition and in some cases even permanently sterilizing females (21, 25). Whereas premating and postzygotic isolating barriers between *D. mojavensis* and *D. arizonae* vary in strength depending on the source population of males and females (29–31), PMPZ isolation is strong in all crosses involving *D. mojavensis* females, suggesting that this barrier may have been among the earliest to evolve.

In the present study we sought to identify and compare the transcriptional changes that occur in female *D. mojavensis* reproductive tracts following conspecific and heterospecific matings at three postcopulatory time points. We first compared virgin and conspecifically mated females to identify genes involved in the normal female postmating response. We then compared this

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transcript set with the transcript differences observed in con- vs. heterospecific crosses, as genes found in both sets are candidates for involvement in PMPZ isolation. Our design thus allows us to investigate the genetic basis of PMPZ isolation in an internally fertilizing organism at its very early stages.

Results

We used *D. mojavensis* whole genome microarrays to compare patterns of transcript abundance in the lower reproductive tracts of virgin and mated females at 15 min, 2 h, and 6 h postcopulation. In addition to finding differences in the expression of lower reproductive tract genes in the three female treatments, we also found that a significant number of male-derived transcripts had been transferred at mating.

Few Transcripts Differ in Abundance Between Conspecifically Mated and Virgin Females. Using the false discovery rate (FDR) to control for multiple testing ($Q = 0.05$), we identified 18 genes for which transcript levels differed between conspecifically mated and virgin females at one or more time points (Table 1). The number of genes in this set is small compared with the much larger set (160) that differed between heterospecifically and conspecifically mated females, reflecting a relative paucity of regulatory changes in the female reproductive tract following conspecific mating. The ratio of differentially regulated genes in the two comparisons is largely unaffected by the choice of FDR cutoff, as a more liberal cutoff ($Q = 0.1$) yields similar results (27 genes in the conspecific–virgin comparison; 246 genes in the conspecific–heterospecific comparison). We subsequently determined that 3 of the 18 transcripts in this set came from males (see below *Males Transfer ACP mRNA Transcripts to Females During Mating*). Twelve of the remaining 15 were definitively of female origin and thus represent genes differentially regulated in response to mating. We were unable to determine the sex of origin of the 3 remaining transcripts; in the absence of evidence to the contrary, we assume that they also represent genes that were differentially regulated in females.

At 15 min postmating, only six genes showed differences in expression (excluding male-derived genes) between mated and virgin females, with transcript levels being significantly higher in mated females for all but one of the genes (Table 1). Although most of these genes have predicted *D. melanogaster* orthologs (some we identified by BLAST to the *D. melanogaster* genome; Table S1), analysis of gene ontology terms revealed that the

molecular function is known for only two: a highly up-regulated gene that codes for an odorant binding protein (*Obp 93A*) and a down-regulated gene that is involved in translation initiation (*Adam*) (Table 1). At 2 h postmating, only two additional genes showed differences in expression between mated and virgin females, including one gene involved in protein binding (CG15515). The peak number of genes exhibiting differences in expression occurred at 6 h postmating and was associated with a shift from primarily up-regulation at earlier time points to more down-regulation at 6 h (Table 1). Most notably, three genes with predicted roles in immune/stress response (a pair of *Thor* homologs and CG6770) were down-regulated at this time.

Regulation of Mating Responsive Genes Is Perturbed in Heterospecific Crosses. Striking differences in patterns of relative transcript abundance between conspecifically and heterospecifically mated females occurred. The majority (9/15) of mating-responsive transcripts identified from the virgin–conspecific comparison also differed between conspecifically and heterospecifically mated females, suggesting that the normal postmating transcriptional response was highly perturbed in heterospecifically mated females (Fig. 1). Moreover, transcriptional variation between the crosses was not limited to mating responsive genes identified from the conspecifically mated–virgin comparison, as heterospecifically mated transcript levels also differed for an additional 148 genes (Table S2). Analysis of gene ontology terms revealed that a number of terms associated with mitochondrial function were overrepresented in this gene set, with almost all being down-regulated in the heterospecific cross (Table S3). The contrast between the two sets of transcript differences is striking and consistent with the previous physical evidence of severe mismatch between heterospecific reproductive tracts.

Males Transfer Acp mRNA Transcripts to Females During Mating. Because Acp's are typically expressed only in male tissues, we were surprised that conspecifically and heterospecifically mated females showed large differences in relative transcript abundance for several previously identified Acp's and/or male reproductive transcripts. Given that most transcripts contain fixed sequence differences between *D. mojavensis* and *D. arizonae*, we used a combination of RT-PCR and sequencing to confirm that at least 12 transcripts in the female reproductive tract were of male origin (Table 2), including 10 that were in higher abun-

Table 1. Identity, predicted molecular function, and fold change in expression (relative to virgin females) of mating responsive transcripts in the lower reproductive tracts of *D. mojavensis* females at three time points postmating

<i>D. mojavensis</i> gene	<i>D. melanogaster</i> homolog	Molecular function	Fold change		
			15 min	2 h	6 h
GI10424	None	Unknown			2
GI10632	<i>Npc2h*</i>	Unknown	19.9		
GI11600	None	Unknown	10		
GI14543	<i>Adam*</i>	Translation initiation factor	–1.6		
GI14761	<i>ripped pocket*</i>	Sodium ion transport	82.3	64	25.8
GI14846	None	Unknown	10.2		
GI14885	CG6770	Stress response			–2.4
GI14996	<i>Thor*</i>	Immune/stress response			–2.7
GI15007	<i>Thor</i>	Immune/stress response			–3.2
GI16692	CG13936	Protein binding			–2.2
GI17134	CG14069	Unknown	10.7		
GI18586	CG34193*	Unknown		4.2	
GI20303	CG30273*	Unknown	78	66.8	33.7
GI22307	CG15515	Protein binding		5.9	
GI23227	CG6972	Protein binding			1.6
GI23324	CG7685	Hydrolase, glycosidase			1.8
GI23726	<i>Obp93A</i>	Odorant binding	11.3	6.3	
GI23890	<i>Scpr-C*</i>	Unknown			5.7

*Homology determined by BLAST to the *D. melanogaster* genome (results in Table S1).

Table 2. Transcripts that differed between con- and heterospecifically mated females that were (i) candidate *D. mojavensis*/*D. arizonae* Acps (ii), genes from *D. melanogaster* male reproductive tracts, or (iii) proteins that were transferred in the ejaculate of *D. melanogaster*

<i>D. moj</i> ID	Gene	Source of transcript	Confirm [§]	<i>D. moj</i> Acp	<i>D. mel</i> male RT	Protein transferred in <i>D. mel</i>	<i>D. mel</i> ortholog	Conspecific mating effect	Reference
GI17858		Male	No [†]	Yes	Yes		CG14034		45, 47
GI20219		Male	Yes	Yes	Yes	Yes	CG30488		45–47
GI19546		Male	No [†]	Yes		Yes	CG34002		45, 46
GI20999	<i>DmojlAcp11</i>	Male	Yes	Yes			—		27
GI20988	<i>DmojlAcp2</i>	Male	Yes	Yes			—		27, 45
GI11629	<i>Adk3</i>	Male	Yes		Yes		CG6612		47
GI23890	<i>ScprC</i>	Male	Yes		Yes		CG5106	Yes	47
GI14761	<i>rpk</i>	Male	No				CG1058	Yes	
GI11382		Male	Yes				—		
GI17134		Male	Yes				CG14069	Yes	
GI23381		Male		Yes	Yes	Yes	CG17097		45–47
GI23009		Male		Yes	Yes		CG10284		45, 47
GI16594		Female		Yes			CG1318		45
GI22128		Female		Yes			CG34215		45
GI13447	<i>Cam</i>	Female			Yes		CG8472		47
GI10528		Unknown [†]		Yes	Yes	Yes	CG14061		45–47
GI18622	<i>DmojlAcp2b*</i>	Unknown [†]		Yes			—		45
GI10529		Unknown [†]		Yes		Yes	CG9997		45, 46
GI10530		Unknown [†]		Yes			—		45
GI13594		Unknown [†]		Yes			CG18233		45
GI13596		Unknown [†]		Yes			CG18233		45
GI20607		Unknown [†]		Yes			CG4812		45
GI21941	<i>capt</i>	Unknown [†]			Yes		CG33979		47

Transcripts were screened by RT-PCR and sequencing to determine whether they were of male or female origin.

*Previously unannotated *D. mojavensis* gene.

[†]Did not amplify.

[‡]Amplified, but problems sequencing.

[§]Independent replicate experiment.

^{||}Female origin in replicate experiment.

^{||}Orthologous call made by BLAST to *D. melanogaster* genome; see Table S1.

melanogaster. Although the production of AMPs in response to mating is clouded by these issues, three other genes involved in immunity or stress response (two *Thor* homologs and CG6770) were down-regulated in mated females at 6 h postmating (Table 1). These same genes were also down-regulated after mating in several *D. melanogaster* studies, with the only exception being that *Thor* was up-regulated in one study that used whole body flies (Table S5).

Response of the Female Reproductive Tract to Heterospecific Mating.

This study clearly demonstrates that transcriptional regulation of mating-responsive genes is highly disrupted in heterospecifically mated *D. mojavensis* female reproductive tracts. Transcript levels of many of the mating-responsive genes were considerably higher in the conspecific cross (most more than twofold) and in most cases these differences were detectable by 15 min postmating and persisted throughout the time course that we examined. The fact that most of the mating-responsive genes were up-regulated in conspecifically mated females relative to virgins indicates that heterospecifically mated females fail to ramp up transcription of key genes involved in the normal female postmating response. In some cases the pattern of regulation was similar between the crosses (i.e., the gene was up-regulated relative to virgins in both crosses), but transcript levels were higher in the conspecific cross, whereas in other cases heterospecifically mated females completely failed to up-regulate key mating genes (Fig. 1). An interesting exception to this general pattern is seen for the three genes predicted to function in immune and/or stress response (two homologs of *Thor* and CG6770), which were down-regulated in both crosses, but more so in heterospecifically mated females.

The overall pattern of misexpression of mating-responsive genes in heterospecifically mated females presumably reflects failed or suboptimal interactions between components of the ejaculate of heterospecific males and the female reproductive tract. In addition,

the fact that transcript abundance also differed for a relatively large number of nonmating-responsive genes (148) points to a complex transcriptional response to the ejaculate of heterospecific males that is not necessarily limited only to genes directly involved in mating. Transcript abundance for the majority of these genes was lower in the heterospecific cross, with genes involved in mitochondrial function, in particular, showing lower transcript levels. Although the reason for this is unclear, the difference was detectable at all time points, and thus could have important long-term metabolic consequences for heterospecifically mated females.

Transfer and Persistence of Male Acp Transcripts in the Female Reproductive Tract.

Our study clearly demonstrates that males transfer mRNA transcripts to females during mating. At least 7 of the 12 male-donated transcripts are previously identified candidate *D. mojavensis* Acps, and two others are orthologous to genes expressed in *D. melanogaster* male reproductive tracts (Table 2). Although the role of these transcripts within the female is unclear, the fact that they represent a small, repeatable subset of transcripts from the male reproductive tract suggests that their inclusion in the ejaculate is not random. Moreover, all were still present at elevated levels in females after 6 h (Fig. 2), indicating that they are not rapidly degraded, as might be expected if they served no functional purpose.

We do not know whether male transcripts that are passed to females are contained within cells or other vessels or whether they are extracellular, but in either case they could play an important functional role. For example, recent evidence in humans suggests that sperm carry mRNA transcripts, some of which are translated de novo by mitochondrial-like ribosomal proteins and appear to play an important role in sperm motility, capacitation, and fertilization (34, 35). Moreover, other transcripts are delivered to the oocyte where they may influence early embryonic development

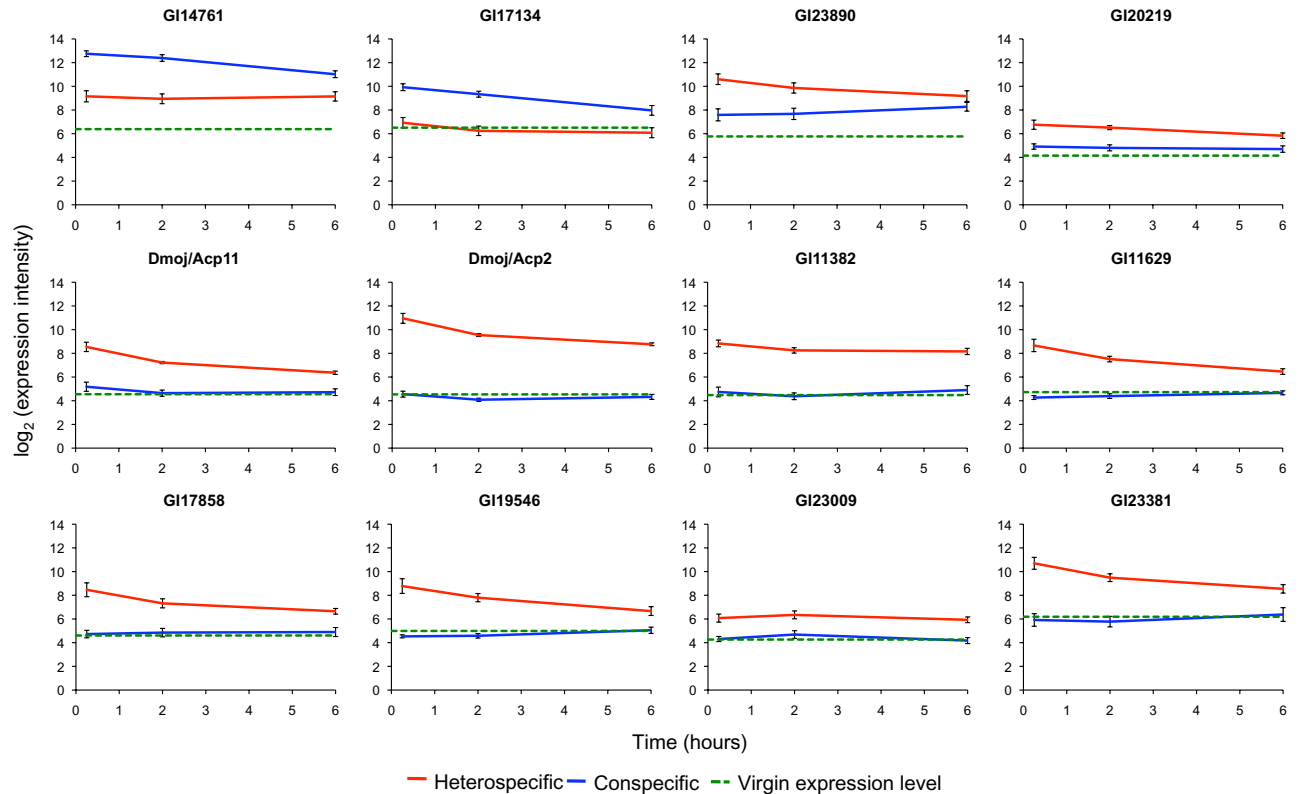


Fig. 2. Relative transcript abundance of male-donated transcripts in conspecifically mated females, heterospecifically mated females, and virgins over the time course of the experiment. The line for virgins corresponds to the average of the two replicates.

(36). Given that at least some of the male-donated transcripts are orthologous to genes in the *D. melanogaster* testes and seminal vesicle, the possibility that sperm carry these transcripts is plausible. In addition to this possibility, we also note that *D. mojavensis* females are unusual among *Drosophila* in that they incorporate significant levels of peptide components of the male ejaculate into their somatic tissues (37). The mechanism for this is unknown, but could conceivably provide a route for the uptake of male-donated mRNA transcripts into female cells where they could be subsequently translated. Given that Acp's play a critical role in many postcopulatory processes (2, 18), the translation of these transcripts into functional proteins within the female, if it occurs, would have important implications for understanding fertilization success, postcopulatory sexual selection, and the evolution of PMPZ isolating barriers.

Even if transferred male transcripts are extracellular and not translated, they still may be functionally important. An increasing number of studies have demonstrated that extracellular nucleic acids play pivotal roles in processes such as blood clotting and immune response (38, 39). For example, recent evidence from a study on the mechanisms of mammalian blood clotting demonstrated that extracellular RNA serves as a procoagulant, providing a template for adhesion of blood coagulation factors in response to vascular injury (38). Similarly, following bacterial infection in the insect, *Galleria mellonella*, extracellular RNA released from damaged cells forms a net-like coagulation structure that efficiently entraps bacteria and triggers important immune defenses (39). In light of this, it seems plausible that male-derived mRNA transcripts could be involved in the formation of the insemination reaction, which at least superficially resembles a coagulatory response in the uterus of *D. mojavensis* females. Although speculative, this intriguing possibility warrants further investigation.

Our study represents a step forward in understanding the genetic basis of sexual conflict and reproductive compensation (40), postcopulatory molecular interactions between the sexes, and

how disruptions in these interactions contribute to the evolution of reproductive isolation between diverging populations. In addition, our discovery of the transfer of male ACP transcripts to females during copulation raises new questions about the nature and complexity of reproductive tract interactions, paving the way for exciting avenues of future research.

Materials and Methods

Mating and Sample Preparation. We used one *D. mojavensis* isofemale line collected from Anza Borrego Desert State Park, Borrego Springs, CA, in April of 2002, and one *D. arizonae* isofemale line collected from Guaymas, Sonora, Mexico, in March of 2007 for our experiments. We collected virgin males and females within 24 h of emergence (flies take at least 5 d to reach reproductive maturity) and aged them separately for 10–12 d. For mating experiments, one *D. mojavensis* female was introduced to a vial containing two *D. mojavensis* or *D. arizonae* males. Following copulation males were removed from vials and females were kept in isolation until lower reproductive tracts were removed at either 15 min, 2 h, or 6 h postmating; we also removed lower reproductive tracts from virgin females at this time. Tissues were placed immediately in TRIzol and frozen at -80°C until total RNA was extracted using a standard TRIzol protocol. We performed two biological replicates for each time point by pooling 20–25 lower reproductive tracts for each sample (14 samples total). Following RNA extraction, mRNA was amplified using the MessageAmp II aRNA amplification kit (Applied Biosystems/Ambion).

Microarray Assay and Data Analysis. The *D. mojavensis* microarray, which included 14,591 genes, was designed for the NimbleGen 4-Plex platform. Before analysis, hybridization intensities were normalized using the robust multichip average (RMA) method, which includes quantile normalization of intensities (41, 42). We analyzed the $\log_2$ of the RMA normalized intensities using a two-step mixed model ANOVA (43). More details on the microarray design and analysis can be found in (*SI Materials and Methods*). All expression data have been deposited in the Gene Expression Omnibus under series entry GSE27454.

We verified microarray results for three genes (*DmoglAcp2*, G117858, and G123890) using quantitative PCR. Working from the same mRNA pools used in the microarray, we synthesized cDNA using ABI's high-capacity RNA-to-cDNA kit. Quantitative PCR reactions were performed on an ABI 7000 Sequence Detection system machine using ABI's Power SYBR Green PCR kit. We ran each gene (including a control: Ribosomal subunit 18S) in triplicate using gene specific primers (Table S4). Statistical significance was calculated by performing 10,000 bootstraps using the REST 2008 software (44).

Analysis of the representation of gene ontology terms was performed using the online Database for Annotation, Visualization and Integrated Discovery (DAVID) at <http://david.abcc.ncifcrf.gov/>. The analysis was based on the level of gene ontology term representation of *D. melanogaster* orthologs of the differentially expressed *D. mojavensis* genes.

Verification of Male-Derived Transcripts. To determine whether the increase in transcript abundance of previously identified male Acp's following mating was due to female up-regulation of these genes or whether they were included in the male ejaculate, we created cDNA libraries from the original aRNA of two samples used in the microarray experiment (conspicuously mated/15 min and heterospecifically mated/15 min). Libraries were constructed using the Bio-Rad Iscript select cDNA synthesis kit with random primers. We used PCR to amplify 23 transcripts that included previously identified *D. mojavensis/D. arizonae* Acp's (27, 45) and/or previously identified transcripts from *D. melanogaster* male reproductive tract (46, 47), in

addition to the 18 genes that differed in transcript abundance between the conspecifically mated females and virgins (Table 1). We then sequenced the amplified products and used fixed differences between the species to determine whether transcripts from the heterospecific cross were from *D. arizonae* (i.e., of male origin) or *D. mojavensis* (i.e., of female origin). Sequences from this analysis that are longer than 200 bp are deposited under GenBank accession nos. JF512479–JF512494; all sequences, including those under 200 bp, are included in Dataset S1.

To independently verify the transfer of male transcripts, we repeated the heterospecific matings using the same *D. mojavensis* line and randomly chosen males from 12 *D. arizonae* lines from Guaymas, Sonora, Mexico (not including the original line used in the microarray). All procedures (matings, RNA extraction, cDNA synthesis) were identical to the original experiment except that we did not perform the mRNA amplification step.

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