Genomics analysis of hexanoic acid exposure in Drosophila species

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Abstract

Drosophila sechellia is a dietary specialist endemic to the Seychelles islands that has evolved to consume the fruit of Morinda citrifolia. When ripe, the fruit of M. citrifolia contains octanoic acid and hexanoic acid, two medium-chain fatty acid volatiles that deter and are toxic to generalist insects. Drosophila sechellia has evolved resistance to these volatiles allowing it to feed almost exclusively on this host plant. The genetic basis of octanoic acid resistance has been the focus of multiple recent studies, but the mechanisms that govern hexanoic acid resistance in D. sechellia remain unknown. To understand how D. sechellia has evolved to specialize on M. citrifolia fruit and avoid the toxic effects of hexanoic acid, we exposed adult D. sechellia, D. melanogaster and D. simulans to hexanoic acid and performed RNA sequencing comparing their transcriptional responses to identify D. sechellia specific responses. Our analysis identified many more genes responding transcriptionally to hexanoic acid in the susceptible generalist species than in the specialist D. sechellia. Interrogation of the sets of differentially expressed genes showed that generalists regulated the expression of many genes involved in metabolism and detoxification whereas the specialist primarily downregulated genes involved in the innate immunity. Using these data, we have identified interesting candidate genes that may be critically important in aspects of adaptation to their food source that contains high concentrations of HA. Understanding how gene expression evolves during dietary specialization is crucial for our understanding of how ecological communities are built and how evolution shapes trophic interactions.

Keywords: toxin resistance; host specialization; RNA-seq

Introduction

Insects have long been recognized as one of the most abundant and diverse groups of organisms on the planet, with a large fraction of them feeding on plants (Jaeinike 1990; Stork 2018). Many of these phytophagous insects have evolved to be highly host plant specific. The evolution of such specialized interactions is often guided by specific plant chemistry, with most plants responding to increased insect herbivory by the production of toxic secondary metabolites (Jaeinike 1990; Petschenka and Agrawal 2016). This leads to an evolutionary arms race as insects evolve resistance to these toxins (Heidel-Fischer and Vogel 2015). While host plant-insect adoptions are well studied in the literature, less is known about the underlying genetic mechanisms that contribute to the evolution of these complex ecological interactions (Ungerer et al. 2008).

Drosophila sechellia feeds and oviposits primarily on Morinda citrifolia, a fruit highly toxic to other Drosophila species (Legal et al. 1992). The plant produces ripe fruit year-round in the Seychelles island archipelago, the sole location where D. sechellia are found (Legal et al. 1992) providing abundant and consistent resources. Upon exposure to the ripe fruit, other Drosophila species display frantic behavior and wing movements, reduction in locomotor activity, and death (Legal et al. 1994). Drosophila sechellia eggs are able to hatch and develop on M. citrifolia fruit but the embryos of other Drosophila species die (Amlou et al. 1998). Unlike its generalist sister species, D. sechellia prefer M. citrifolia to other hosts and are drawn to the fruit from a long distance (R’Khā et al. 1991). Drosophila sechellia evolved resistance to the toxins in M. citrifolia from an ancestral sensitive state (R’Khā et al., 1997) and this tolerance provides D. sechellia a temporal advantage over other fruit fly species that can only lay eggs in the fruit once it has rotten and the toxic volatiles are reduced.

The toxic properties of Morinda are attributed to the carboxylic acids present in the pulp of the ripe fruit (Legal et al. 1994). This pulp is largely characterized by carboxylic acids, primarily the fatty acids octanoic (OA) and hexanoic acid (HA) which comprise 58% and 19% of the volatile compounds found the ripe fruit respectively (Farine et al. 1996). While the genetic basis of D. sechellia resistance to the most abundant and toxic compound in M. citrifolia fruit, OA, has been characterized in previous studies (Dworkin and Jones 2009; Andrade Lopez et al. 2017; Lanno et al. 2017; Peyser et al. 2017; Lanno and Coolon 2019; Lanno et al. 2019a,
2019b), much less is known about the genes involved in HA resistance. In a study using fatty acid concentrations equivalent to 1.5 g of ripe M. citrifolia fruit, OA treatment alone killed all fruit fly species assayed except D. sechellia, while HA alone caused reversible knock-down in other Drosophilids (Farine et al. 1996). In a more recent study using higher concentrations of HA, mortality was observed in response to exposure to HA in D. melanogaster and D. simulans and D. sechellia is significantly more resistant to HA induced mortality than sister species (Peyser et al. 2017; Lanno and Coolon 2019). Surprisingly, tests of the three major detoxification gene families, cytochrome P450s (cyps), glutathione-S-transferases (GSTs) and esterases (Ests) found that none was involved in derived HA resistance in D. sechellia suggesting an alternative genetic mechanism must be involved in resistance. Studies have suggested that HA may be a more efficient D. sechellia attractant than OA, while OA is a more potent repellant of generalist species (Amlou et al. 1998). In a test using laboratory food medium supplemented with 0.5% of either OA or HA, D. sechellia exhibited oviposition preference for media supplemented with HA over OA (Amlou et al. 1998).

From a study analyzing the transcriptomic response of D. sechellia on OA, 104 genes were found to be differentially expressed in response to OA (Lanno et al. 2017). This study showed that several Osiris genes, including Os6 are upregulated in D. sechellia in response to OA. Another study showed that RNAmediated knockdown of Os6 expression drastically decreased survival in response to OA (Andrade Lopez et al. 2017). Given that HA makes up about of fifth of the volatile compounds found in M. citrifolia and produces unique effects on behavior (Farine et al. 1996) and has an unknown and less common basis for toxin resistance (Peyser et al. 2017; Lanno and Coolon 2019), identifying the genes responding to HA is necessary to understand how D. sechellia has specialized on M. citrifolia fruit, and may help pinpoint genes that are involved in resistance to HA. By also analyzing the genes responding to HA exposure in generalist Drosophila species D. melanogaster and D. simulans we can identify derived gene expression responses specific to D. sechellia that may be critical for HA associated traits (Coolon et al. 2009). In this study, adult female D. sechellia, D. melanogaster, and D. simulans flies were fed either control food or food supplemented with 0.23% HA and significantly differentially expressed genes (DEGs) were identified using RNA-seq. Comparison of the identified genes with those found to respond to OA (Lanno et al. 2017) and L-DOPA (Lanno et al. 2019), another highly abundant compound found in M. citrifolia fruit, identified several genes common in response to OA, HA, and L-DOPA as well as genes unique to HA suggesting these genes may play an important role in the evolved resistance and specialization of D. sechellia to M. citrifolia.

Methods

Fly strains and culture

Drosophila sechellia (14021-0428.25), D. simulans (14021-0251.195), and D. melanogaster (14021-0231.36) flies were reared on standard cornmeal medium under a 16:8 light: dark cycle maintained at 20°C.

RNA extraction, library preparation, and sequencing

Zero to 3-day posteclosion adult female flies were fed control food (0.75 g Drosophila instant medium Formula 4-24, Carolina Biological Supply Company) or identical food containing 0.23% hexanoic acid (HA). After 24 h, three replicates of ten whole flies per species and per treatment were homogenized and total RNA was extracted with a modified protocol of the Promega 5V extraction system (Coolon et al. 2013; Figure 1). RNA quality and quantity was assessed using agarose gel electrophoresis and Nanodrop spectrophotometry. RNA was sent to the University of Michigan Sequencing Core Facility for poly-A selection, cDNA synthesis, bar-coded library preparation with TruSeq library preparation kits and sequencing on an Illumina platform, generating 405,166,795 single-end 65 nt sequencing reads for D. sechellia and 51 nt sequencing reads for D. melanogaster and D. simulans (Table 1).

BIOL310 genomics analysis

The genomics analysis of RNA-seq data presented in this manuscript was performed by 20 undergraduate and 6 graduate students as part of a semester-long course at Wesleyan University called Genomics Analysis (BIOL310). This is the third such manuscript (see Lanno et al. 2017, 2019a) made from this course where the aim is to provide undergraduate students an early opportunity with a course-based research experience with active participation in scientific discovery. Students in the course learn through engaging with never-before analyzed data using cutting edge genomics analysis techniques and bioinformatics tools through a discovery-based independent study. Every student in the course contributed to the quality control, analyses, write-up, and interpretation of the findings, providing their own unique perspective of the results and text written by each and every student was combined into this manuscript with very little modification.

After sequencing output files were obtained from the University of Michigan Sequencing Core (Table 1), fastq files containing raw sequencing reads were uploaded to the Galaxy platform (Afgan et al. 2016) and an RNA-seq analysis pipeline was performed (Figure 1) as previously described (Lanno et al. 2017, 2019a). Briefly, reads were assessed for quality using FASTQC (Andrews 2010), and any overrepresented sequences were analyzed using NCBI Blast (Altschul et al. 1990). Bowtie2 was used for mapping reads to the appropriate reference genome for each species with default parameters (Langmead and Salzberg 2012), with the most recent genomes for each species available at the time of analysis acquired from Ensembl (www.ensembl.org, Yates et al., 2020) (D. sechellia: Drosophila_sechellia.dsec_caf1.dna.toplevel.fa, D. simulans: Drosophila_simulans.ASM75419v3.dna.toplevel.fa and D. melanogaster: Drosophila_melanogaster.BDGP6.dna.toplevel.fa). The Bowtie2 output files were analyzed using Cuffdiff (Trapnell et al. 2010), which performs gene expression quantification and differential gene expression analysis using the aforementioned genome file along with the most recent annotated .gff3 file for each genome available at the time of analysis acquired from Ensembl (D. sechellia: Drosophila_sechellia.dsec_caf1.42.gff3, D. simulans: Drosophila_simulans.ASM75419v3.42.gff3 and D. melanogaster: Drosophila_melanogaster.BDGP6.95.gff3). In Cuffdiff, geometric normalization and library size correction was performed, along with bias correction using the reference genome, giving an output of DEGs for each species following false discovery rate multiple testing correction (Benjamini and Hochberg 1995, q < 0.05). Data were visualized using R (R Core Development Team 2010). In order to compare gene expression results across species, we obtained all 1:1:1 orthologs from D. sechellia, D. simulans, and D. melanogaster from Flybase (Thurmond...
et al. 2019). DEGs following D. sechellia exposure to OA or L-DOPA were downloaded from online databases (Lanno et al. 2017, 2019a). GO term enrichment was performed on D. melanogaster orthologs for each species using GeneOntology.org (www.geneontology.org, Ashburner et al. 2000; Mi et al. 2019; Carbon et al. 2021). KEGG pathway analysis was performed using the D. melanogaster ortholog for each DEG from each species (https://www.kegg.jp/kegg/tool/map_pathway1.html, Kanehisa and Sato 2020).

Table 1  Sequencing results and mapping percentage

| Sample | ID       | No. of reads | No. of mapped reads | % Mapped | Read length (nt) |
|--------|----------|--------------|---------------------|----------|-----------------|
| sim-C-1| 105545   | 28,056,123   | 26,210,691          | 93.42    | 51              |
| sim-C-2| 105546   | 26,058,213   | 24,449,785          | 93.82    | 51              |
| sim-C-3| 105547   | 24,095,284   | 22,589,715          | 93.75    | 51              |
| sim-HA-1| 105558  | 27,002,538   | 25,521,057          | 94.51    | 51              |
| sim-HA-2| 105559  | 23,222,911   | 21,786,539          | 93.81    | 51              |
| sim-HA-3| 105560  | 21,999,530   | 20,633,866          | 93.79    | 51              |
| mel-C-1| 105542   | 20,950,464   | 19,779,953          | 94.41    | 51              |
| mel-C-2| 105544   | 21,157,160   | 20,919,514          | 94.41    | 51              |
| mel-HA-1| 105554  | 19,811,200   | 18,745,553          | 94.62    | 51              |
| mel-HA-2| 105555  | 23,577,339   | 22,322,344          | 94.68    | 51              |
| mel-HA-3| 105556  | 23,222,911   | 21,786,539          | 94.51    | 51              |
| sec-C-1| 76332    | 19,122,060   | 18,496,450          | 96.23    | 65              |
| sec-C-2| 76333    | 20,740,811   | 19,440,620          | 93.89    | 65              |
| sec-C-3| 76334    | 17,966,868   | 17,123,579          | 96.76    | 65              |
| sec-HA-1| 76338   | 30,612,710   | 29,079,271          | 94.99    | 65              |
| sec-HA-2| 76339   | 25,873,099   | 25,202,396          | 97.41    | 65              |
| sec-HA-3| 76340   | 19,074,054   | 18,031,453          | 94.53    | 65              |
Results

Differential gene expression in response to HA treatment

In order to identify candidate genes that are important in *D. sechellia* host specialization and evolved resistance to HA, we sought genes that have altered expression levels in response to HA exposure. Previous studies have shown that such environmentally plastic gene regulation can indicate the importance of that gene’s function in that environment making identified genes good candidates for *D. sechellia* HA associated traits (Coolon et al. 2009; Lanno et al. 2017, 2019). To quantify gene expression response to HA we performed RNA-sequencing (RNA-seq) on adult female flies after exposure to control food and compared this to flies fed food containing 0.23% HA. Because many of the transcriptional responses to HA might be nonspecific, we measured gene expression responses in *D. sechellia*, *D. melanogaster*, and *D. simulans* to identify those responses (or loss of response) that are restricted to *D. sechellia* representing changes that might contribute to its unique phenotypes.

Using this approach, we identified 841 genes differentially expressed by *D. melanogaster* (Figure 2, A and D; Supplementary Table S1), 743 genes were differentially expressed by *D. simulans* (Figure 2, B and E; Supplementary Table S2) and only 93 genes were differentially expressed in *D. sechellia* (Figure 2, C and F; Supplementary Table S3) in response to HA. No significant difference in the number of upregulated genes (50/93) vs downregulated genes (43/93) was observed in *D. sechellia* in response to HA (Binomial Exact Test, *P* = 0.1066). In *D. simulans*, there was a significant difference in the number of upregulated genes (69/743) compared to the number of downregulated genes (674/743) in response to HA (Binomial Exact Test, *P* = 2.2e-16, Figure 2, B and E). In *D. melanogaster*, there was also a significant difference in the number of upregulated genes (171/841) compared to the number of downregulated genes (670/841) in response to HA (Binomial Exact Test, *P* = 2.2e-16, Figure 2, A and D). In *D. sechellia*, there were 39 DEGs identified that responded to HA that do not have annotated *D. melanogaster* orthologs. 27 of these genes were 5.8S rRNAs, two snoRNAs, and 7 genes of unknown function. Of the 27 5.8S rRNAs, all 27 were upregulated (Binomial Exact Test, *P* = 7.451e-09). In *D. simulans*, of the 743 DEGs there were annotated *D. melanogaster* orthologs for 673 genes. For the remainder of the analysis, only genes with known *D. melanogaster* orthologs are considered to allow functional interpretations of DEGs, and for all subsequent analyses the *D. melanogaster* ortholog name was used. This filtering resulted in 673 differentially genes in *D. simulans*, 841 in *D. melanogaster* and 54 in *D. sechellia* used in subsequent analyses.

Identifying functional enrichment in DEGs

To identify the biological pathways that are involved in responses to HA in *D. sechellia*, *D. melanogaster*, and *D. simulans* KEGG analyses of the upregulated and downregulated genes in each species were performed. These analyses show that many different metabolic and detoxification pathways along with proteins that localize to the lysosome are changing in response to HA exposure in *D. melanogaster* and *D. simulans*, whereas very few genes in these pathways are responding in *D. sechellia* (Figure 3, A and B). In *D. sechellia*, many of the genes downregulated in response to HA are involved in immune processes whereas this was much less prominent in *D. melanogaster* and *D. simulans* DEGs (Figure 3C).

To understand which biological and cellular processes are being altered in *D. sechellia*, *D. melanogaster*, and *D. simulans* in response to HA, Gene Ontology (GO) term enrichment analysis was performed (Supplementary Tables S16–S18). In *D. sechellia*, GO term enrichment analysis of DEGs for cellular component showed a significant enrichment for extracellular region genes (*P* = 4.83e-05), suggesting that several of the genes responding

![Figure 2](image-url)
to HA exposure have proteins that are secreted. For biological process GO term enrichment analysis, processes involved in the antibacterial humoral response were significantly enriched \((P = 1.75e-08)\). In D. sechellia, no molecular function processes were significantly enriched. In D. melanogaster, there was also significant enrichment for GO terms for extracellular region genes \((P = 4.42e-04)\), along with genes found inside the nucleus \((P = 8.40e-04)\). For biological process, GO terms associated with genes involved in the antibacterial humoral response were significantly enriched as in D. sechellia \((P = 3.02e-02)\), as were other terms involved in the Drosophila immunity. The D. melanogaster HA response also was enriched for genes involved in ribosome biogenesis \((P = 3.66e-02)\). In D. simulans, genes found inside the nucleus were significantly enriched in the set of HA responsive genes \((P = 2.65e-02)\). In an analysis of DEGs in D. simulans that are upregulated, genes involved in Notch signaling were significantly enriched \((P = 4.11e-04)\) alongside genes involved in vitellogenic membrane and chorion formation \((P = 2.56e-02)\). In D. sechellia, upregulated genes were significantly enriched for the larval serum protein complex \((P = 1.96e-02)\). Downregulated genes were significantly enriched with antibacterial humoral response GO terms \((P = 3.24e-10)\) along with the response to hyperoxia \((P = 1.52e-03)\). These downregulated genes were also enriched for the extracellular region \((P = 4.17e-05)\). In D. melanogaster, upregulated genes were significantly enriched to be involved in the larval serum protein complex \((P = 1.23e-03)\) and were enriched intracellularly \((P = 4.02e-16)\) and within intracellular organelles \((1.32e-13)\). These upregulated genes were significantly enriched in many biological processes, including chromatin silencing \((P = 3.25e-03)\), ecdysone receptor-mediated signaling \((P = 1.89e-02)\), and chorion assembly \((P = 2.25e-02)\). In D. melanogaster, downregulated genes were significantly enriched for the antibacterial humoral response \((P = 4.52e-03)\) and for the defense response to Gram-positive bacteria \((P = 5.07e-04)\). These downregulated genes were significantly enriched to be localized to the extracellular region \((P = 1.25e-04)\).

### Comparing DEGs identified in response to HA in D. melanogaster, D. simulans, and D. sechellia

In order to identify genes with D. sechellia specific responses to HA, we compared the DEGs from the three species when exposed to HA (Figure 4A). We identified 33 genes that were differentially expressed by D. sechellia flies exposed to HA that were not responsive to HA exposure in both D. melanogaster and D. simulans flies (Figure 4A, Supplementary Table S13). Gene ontology enrichment analysis was performed for the 33 DEGs that are unique to the response to HA in D. sechellia and the downregulated DEGs are significantly enriched for immune response GO terms and response to stress \((GO:0006950, \text{Supplementary Table S23})\). There is no significant enrichment of unique upregulated DEGs in D. sechellia. In order to identify those genes where D. sechellia specific loss of response to HA was observed we selected those genes with significant changes in response to HA in both D. melanogaster and D. simulans that were not significantly differentially expressed by D. sechellia in response to HA. This analysis yields a total of 213 genes with this expression pattern in our data (Supplementary Table S10). Of these 213 DEGs shared between D. melanogaster and D. simulans, 205 of these genes are downregulated in both species in response to HA (Supplementary Table S10). GO analysis of these DEGs shared between D. melanogaster and D. simulans gives no significant enrichment. There are only 4 DEGs unique to D. melanogaster and D. simulans that are upregulated in both species (CG11835, Cpr62Bc, cype, and Rp16). Interestingly, only 2 genes were identified, GG13114 and Fbp1, that had significant response to HA in all three species, with the expression of both increasing in D. simulans and decreasing in D. sechellia and D. melanogaster (Figure 4A, Supplementary Table S9).

### Common transcriptional responses of D. sechellia exposed to HA, OA, and L-DOPA suggests overlapping regulatory mechanisms

To assess the overlap in transcriptional responses of D. sechellia to both OA and HA, the predominant fatty acid volatiles in M. citrifolia, along with 3,4-dihydroxyphenylalanine (L-DOPA), which is found in M. citrifolia fruit and is important for the specialization
of D. sechellia to this fruit (Lavista-Llanos et al. 2014; Lanno et al. 2019a), we compared DEGs between our HA treatment, DEGs identified in a previous study using a 0.7% OA treatment (Lanno et al. 2017), and DEGs identified in a previous study of responses to 10 mg/mL of L-DOPA added to the fly food (Lanno et al. 2019a). Treatment with OA treatment yielded 103 D. sechellia genes with D. melanogaster orthologs that were significantly differentially expressed. Treatment with L-DOPA yielded 643 D. sechellia genes with D. melanogaster orthologs that were significantly differentially expressed. Comparison of genes responsive to HA, OA, and L-DOPA identified 12 DEGs that respond to HA, OA, and L-DOPA treatment in D. sechellia (Figure 4B, Supplementary Table S6). Interestingly, of the 19 shared DEGs with D. melanogaster orthologs between OA and HA treatments, all genes are downregulated except for P(spl)ngamma-HLH, which is upregulated upon both OA and HA treatment (Supplementary Table S4). To compare metabolic and cellular pathways involved in the conserved response between OA and HA treatment in D. sechellia, GO term enrichment analysis was performed on shared DEGs. Genes involved with biological processes related to the humoral immune response were overrepresented (P = 2.95e-08, Supplementary Table S22). No significantly enriched processes were found for molecular function or cellular component. A KEGG pathway analysis of DEGs in D. sechellia similarly found that upon OA, D. sechellia downregulated genes involved in both the Toll and Imd signaling pathways as well as genes involved in metabolic processes (Supplementary Table S21 and Figure S1). Interestingly, in response to OA and HA, D. sechellia downregulated genes involved in the humoral immune response (AttC, CecA2, Def, DptB, Dro, edin, and PGRP-SB1 Supplementary Table S4).

Discussion

Understanding the genetic basis of how organisms evolve to occupy different ecological niches and adapt to their environments is crucial to understanding the evolution of plant and animal interactions. Insect-host plant specialization is an excellent example of the evolution of such interactions and has been the subject of numerous ecological studies. While the phenomenon is well documented, the genetic basis of evolved host specialization is still not widely understood. Here, we focus on the specialization of D. sechellia to feed almost exclusively on one host plant, M. citrifolia because it is an excellent model to understand the genetic basis of dietary specialization. This is in part because it is has evolved recently and very closely related to the genetic model generalist species D. melanogaster. Fortuitously, we can take advantage of the wealth of genetic tools and information about D. melanogaster and sister species to understand the evolution of dietary specialization in this group (Groen and Whiteman 2016).

Drosophila sechellia upregulated a single cytochrome P450 (Cyp46I) in response to HA whereas D. simulans downregulates this same gene (Supplementary Table S2), but previous work has shown that cytochrome P450s are not the evolved mechanism by which D. sechellia is able to survive OA or HA exposure (Peyser et al. 2017). Previous work has shown that HA induces a “reversible coma” in generalist D. melanogaster, D. simulans, and D. mauritiana flies (Farine et al. 1996), and D. sechellia prefers to oviposit on HA compared to OA and control food sources (Amlou et al. 1998). Drosophila sechellia has a premature stop codon in Obp56e as well as Obp57d and Obp57e alleles that reduce their avoidance to noni volatiles (Matsuo 2007; Dworkin and Jones 2009). The mechanisms that drive this attraction may be through changes in gene expression and may be reflected in these predicted regulatory networks. Drosophila sechellia downregulates many genes involved in the immune response when exposed to both HA and OA, suggesting that somehow these two medium-chain fatty acids are interacting with negative effectors of Imd or Toll signaling. As these two immune pathways have similar and overlapping target genes, further analysis is needed to determine which immune pathways are involved in these interactions (Hanson and Lemaitre 2020). This interaction between D. sechellia and the volatiles from M. citrifolia reduces the immune response, which could cause alterations in gut microbiota composition that aids in the detoxification of relevant plant secondary metabolites. E(spl)ngamma-HLH is a Notch responsive Myc-like transcription factor that has been shown to interact with Relish, the main regulator of Imd immune signaling (Dushay et al. 1996) through recent yeast two-hybrid assays (Shokri et al. 2019) and is upregulated in D. sechellia in response to both OA and HA exposure (Figure 4B, Supplementary Table S4). The plastic response of insect immune systems allows them to fend off pathogens when needed, but also allow for the management of endosymbiots (Login et al. 2011; Vilcinskas 2013). Unlike its generalist sister species, Drosophila sechellia was previously shown to lack an immune response when confronted with parasitic wasps, hinting that immune system responses in this species may be unlike its generalist sister species (Salazar-Jaramillo et al. 2017). In addition, prior work showed that free fatty acids are involved in regulating

Figure 4 DEG overlap between species and treatments. (A) The number of differentially expressed genes in response to HA after RNA-seq are shown for each species, D. sechellia, D. simulans, and D. melanogaster. Overlap and species-specific number of DEGs are indicated. (B) The number of differentially expressed genes in D. sechellia when it is exposed to OA, HA, or L-DOPA treatment as well as the number of specific and overlapping genes are shown.
immune responses in mammals (Alvarez-Curto and Milligan 2016), and hexanoic acid priming of plants can activate the jasmonic acid pathway to increase plant resistance to fungal pathogens (García-Robles et al. 2013; Aranega-Bou et al. 2014). Using functional studies to alter the expression of known immune effectors and test if this alters resistance or preference for noni volatiles would be useful in understanding the interaction between immunity and these chemicals. Likewise, further study comparing the microbiomes between D. sechellia and D. simulans and how they may change from feeding on M. citrifolia fruit as well as how each responds to pathogen challenge in this context would help to understand why D. sechellia is downregulating its immune response when there could be a serious fitness cost of this action.

Recent studies have shown that D. melanogaster uses sweet-tasting gustatory receptor neurons to sense OA and HA (Masek and Keene 2013; Chen and Amrein 2017; Tauber et al. 2017). Our data show that upon exposure to HA, only D. sechellia significantly upregulates another odorant-binding protein, Obp56a. In contrast to HA treatment, in response to OA D. sechellia adults do not change expression of any odorant-binding proteins (Lanno et al. 2017). Both D. simulans and D. melanogaster downregulate the expression of Obp57a in response to HA (Supplementary Table S10), and D. sechellia upregulates the expression of Obp56a in response to HA (Supplementary Table S13). Recently, studies examining the gustatory and behavioral basis of the attraction of D. sechellia to noni have shown that Or22a neurons in the fly brain are involved in the attraction to noni (Auer et al. 2020), as are Or85b/c and Ir75b neurons (Prieto-Godino et al., 2017). Oro mutant D. sechellia flies lose olfactory responses to both 2-heptanone and 1-hexanol, two compounds found in ripe noni fruit (Auer et al. 2020). Further work on the role of these genes in D. sechellia and the regulatory mechanisms responsible for the change of odorant-binding protein expression in response to HA may help to elucidate how D. sechellia has evolved to specialize on M. citrifolia.

Drosophila sechellia is altering the expression of far fewer genes in response to HA than in the generalist species D. melanogaster and D. simulans (Supplementary Tables S1–S3). Drosophila sechellia appears to be downregulating many genes involved in the immune response whereas D. melanogaster and D. simulans alter the expression of genes involved in many different metabolic pathways (Figure 4, A–C). As generalist insects feed on many different plants which produce different secondary metabolites to defend themselves from predators, perhaps generalist species have a more plastic regulatory response to subvert toxicity whereas the specialist D. sechellia is resistant to the toxicity of its host through a specific constitutive mechanism. A recent study comparing the fitness of D. melanogaster to D. sechellia larvae fed different food sources showed a loss of carbohydrate metabolic responses in D. sechellia, as they have specialized on a fruit with a relatively low sugar content, M. citrifolia (Watanabe et al. 2019). Of the many significant genes differentially expressed in D. melanogaster and D. simulans, most DEGs are not drastically differentially expressed in HA treatment compared to controls (Figure 2, D and E). Drosophila sechellia conversely alters the expression of relatively far fewer genes, but many of the DEGs in response to HA are drastically differentially expressed (Figure 2F). Similarly, another study found that when adapted to a grass diet, Spodoptera littoralis had a smaller transcriptional response when fed maize compared to more generalist S. littoralis (Roy et al. 2016). Determining the scale of these responsive regulatory effects and their role in toxic resistance will help elucidate how D. sechellia has evolved to avoid the toxicity of M. citrifolia.

In response to OA, D. sechellia increases its expression of several Osiris genes (Lanno et al. 2017). Previous work using RNAi in D. melanogaster to knock-down the expression of individual genes and examine survival in these flies when exposed to OA showed that the reduction of Osi6, Osi7, and Osi8 expression decreased survival (Andrade Lopez et al. 2017). In response to HA, neither D. sechellia nor D. melanogaster significantly alter the expression of any Osiris genes, but D. simulans downregulates the expression of Osi6, Osi7, and Osi15 (Supplementary Table S2). The cellular and physiological function of these genes is unknown, so understanding what these genes are doing and how they may be helping to shape the interactions between these insects and their toxic hosts will be useful to better understand how these interactions evolve (Coolon et al. 2019).

Examining and comparing the changes in transcriptional output of insects when exposed to these different plant chemicals in specialized vs generalist species provides a framework to understand how these interactions have evolved. Pathway analyses of these genes are useful in determining the physiological function of altered expression in response to these chemicals, but functional analyses of transcription factors that alter expression of these genes are necessary to better understand the regulatory mechanisms involved in the dietary specialization. Comparing transcription factors responding to plant chemicals may help elucidate regulatory mechanisms involved in these responses and shed light on how insects use changes in the transcription of target genes in order to compete against plants in this evolutionary arms race to adapt to toxic food sources.

Data availability

All RNA-seq data generated in this manuscript have been submitted to the NCBI Gene Expression Omnibus under accession number GSE185176. Supplementary tables for this manuscript have been uploaded to GSA figshare: https://doi.org/10.25387/g3.14745684.

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Conflicts of interest

The authors declare that there is no conflict of interest.

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