

RESEARCH ARTICLE

Divergence of the diapause transcriptome in apple maggot flies: winter regulation and post-winter transcriptional repression

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ABSTRACT

The duration of dormancy regulates seasonal timing in many organisms and may be modulated by day length and temperature. Though photoperiodic modulation has been well studied, temperature modulation of dormancy has received less attention. Here, we leverage genetic variation in diapause in the apple maggot fly, *Rhagoletis pomonella*, to test whether gene expression during winter or following spring warming regulates diapause duration. We used RNAseq to compare transcript abundance during and after simulated winter between an apple-infesting population and a hawthorn-infesting population where the apple population ends pupal diapause earlier than the hawthorn-infesting population. Marked differences in transcription between the two populations during winter suggests that the 'early' apple population is developmentally advanced compared with the 'late' hawthorn population prior to spring warming, with transcripts participating in growth and developmental processes relatively up-regulated in apple pupae during the winter cold period. Thus, regulatory differences during winter ultimately drive phenological differences that manifest themselves in the following summer. Expression and polymorphism analysis identify candidate genes in the Wnt and insulin signaling pathways that contribute to population differences in seasonality. Both populations remained in diapause and displayed a pattern of up- and then down-regulation (or vice versa) of growth-related transcripts following warming, consistent with transcriptional repression. The ability to repress growth stimulated by permissive temperatures is likely critical to avoid mismatched phenology and excessive metabolic demand. Compared with diapause studies in other insects, our results suggest some overlap in candidate genes/pathways, though the timing and direction of changes in transcription are likely species specific.

KEY WORDS: Diapause, Phenology, Overwintering, Gene expression, *Rhagoletis*

INTRODUCTION

The waxing and waning of climatic factors and resources over predictable time periods dictates nearly every aspect of an organism's life history in seasonal environments. Accordingly,

understanding the synchronization of growth and reproduction with permissive conditions and the physiological mechanisms that determine this synchrony has been an important focus of research on adaptive evolution. Through such studies, a general understanding has emerged of how life histories are shaped by environmental uncertainty (Childs et al., 2010; Cohen, 1970; Kingsolver, 1979), of environmental effects on growth and development (Amano et al., 2014; Powell et al., 2000; Taylor, 1981), and of the genetic architecture underlying seasonal timing (Bradshaw et al., 2012; Lair et al., 1997; Li et al., 2010; Schmidt et al., 2008). Similarly, the physiological mechanisms that regulate seasonal timing in animals and plants have been investigated, particularly endocrine mechanisms regulating dormant, overwintering life stages (diapause) in insects (Denlinger, 2002; Hahn and Denlinger, 2011).

Diapause in insects is typically induced and ended by environmental cues such as temperature and photoperiod. These cues are important even for univoltine species that enter an obligate diapause stage, because termination of obligate diapause is often sensitive to temperature and photoperiod (Tauber et al., 1986). Different physiological mechanisms likely transduce temperature, photoperiod and other important cues such as diet quality or quantity. However, the same major neuroendocrine systems regulate entrance into and exit from developmental arrest and metabolic suppression broadly across taxa, although regulatory mechanisms upstream of hormonal signals appear to be taxon specific (Denlinger et al., 2005). These neuroendocrine signals control the physiologically dynamic progression of insects through initiation, maintenance and termination phases collectively termed 'diapause development' (Kostal, 2006).

Because diapause involves a developmental progression, processes affecting the rate of diapause development will affect diapause duration. Diapause duration, in turn, determines the seasonal timing of exit from diapause and resumption of active growth and reproduction. Thus, regulatory mechanisms acting relatively early in diapause development may influence the timing of the end of diapause occurring weeks, months or even years later. These mechanisms may be less important for species that complete diapause in response to a specific photoperiodic cue. For example, diapausing pitcher plant mosquito larvae rapidly end diapause and resume active growth when switched from relatively short to long day lengths, constituting a reliable cue for permissive environmental conditions in the field (Bradshaw and Lounibos, 1977). However, temperature appears to be the primary factor determining the duration of diapause in many species, even for some insects that initiate diapause based on photoperiodic cues (Tauber et al., 1986). Thermal responses over the course of diapause in such species may thus regulate diapause development rate, and consequently the total duration of diapause.

While physiological responses to photoperiod have been well established in a number of diapausing species, little is known

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about when and how temperature modifies diapause development. Photoperiodic responses can be readily manipulated in the lab, and numerous physiological and genetic studies have identified candidate genes and mechanisms that may transduce photoperiodic signals upstream and downstream of major hormonal cues (Emerson et al., 2010; Poelchau et al., 2013; Poupardin et al., 2015; Schmidt et al., 2008; Sim and Denlinger, 2008; Tauber et al., 2007; Wadsworth and Dopman, 2015; Williams et al., 2006). In contrast, it has been more difficult to experimentally manipulate temperature effects because the sensitivity of development to thermal conditions often changes over the course of diapause (Hodek and Hodkova, 1988). For example, many, though not all, diapause-overwintering insects have relatively low thermal thresholds above which diapause development does not proceed or progresses very slowly, similar to vernalization in plants. In the literature, such a threshold is often referred to as a ‘chilling requirement’, which is something of a misnomer because typically exposure to low temperature is not

necessarily required, but rather affects the rate at which insects progress through phases of diapause development (Hodek and Hodkova, 1988). In addition, thermal thresholds can change over the course of diapause. As a result, thermal conditions during the autumn, winter and spring may have varied effects on the timing of diapause termination.

Here, we tested for transcriptomic signatures of regulatory events that may determine the duration of diapause in the apple maggot fly [*Rhagoletis pomonella* (Walsh 1867); Diptera: Tephritidae]. Specifically, we tested for transcriptomic differences between apple- and hawthorn-infesting host races of *R. pomonella* that recently evolved seasonal differences in the timing of their diapause termination. Pupal diapause in *R. pomonella* is functionally obligate in the field (populations are univoltine) and is mainly influenced by overwinter and post-winter temperature (Neilson, 1962). The termination of diapause synchronizes adult flies with the availability of host fruits where they court, mate and oviposit their eggs, and their larvae develop. Recently derived populations of *R. pomonella* attacking apple (*Malus domestica*) have evolved to terminate their diapause earlier than flies attacking the ancestral hawthorn (*Crataegus* spp.) host to track the earlier fruiting time of apples compared with hawthorns (Bush, 1969; Feder and Filchak, 1999; Feder et al., 1993; Smith, 1988). Though hawthorn- and apple-infesting populations eat different fruits as larvae, apples and hawthorns are nutritionally similar, with the larger apples generally yielding larger-bodied adults with greater lipid reserves because of greater resource quantity (Ragland et al., 2012). However, common garden experiments rearing both populations through apple clearly demonstrate that differences in diet do not contribute to the pronounced population difference in eclosion time observed in the field or the laboratory (Dambroski and Feder, 2007; Feder and Filchak, 1999; Smith, 1988).

We applied an RNA sequencing (RNAseq) approach to test two hypotheses concerning the eclosion time difference between apple and hawthorn flies. We examined whether earlier diapause termination and adult eclosion in apple flies is associated with: (1) differences in gene expression during winter (‘during-winter’ hypothesis); or (2) changes in gene expression in response to warming temperatures following winter (‘after-winter’ hypothesis). If the during-winter hypothesis is true, then by the end of winter many genes associated with development will be differentially expressed in apple compared with hawthorn flies (Fig. 1A). In contrast, if the after-winter hypothesis is true, then developmentally related genes will be differentially expressed between the host races only following post-winter warming (Fig. 1B). Both hypotheses predict that apple flies up- or down-regulate development-related genes earlier than hawthorn flies, in accordance with apple flies terminating diapause and eclosing earlier as adults than hawthorn flies. The distinction concerns when the predicted change in gene expression occurs between the two host races: during winter or during spring/summer. We note that dietary differences between populations could contribute to observed gene expression differences, but are unlikely to account for population differences in the non-feeding, diapausing pupal stage given that diet does not account for population differences in *R. pomonella* seasonality (Dambroski and Feder, 2007; Smith, 1988). Finally, we tested whether single nucleotide polymorphisms (SNPs) in differentially expressed genes between apple and hawthorn flies display significant allele frequency differences between the host races in nature. Such a finding would imply that the differentially expressed loci could be the actual targets of divergent selection on diapause timing.

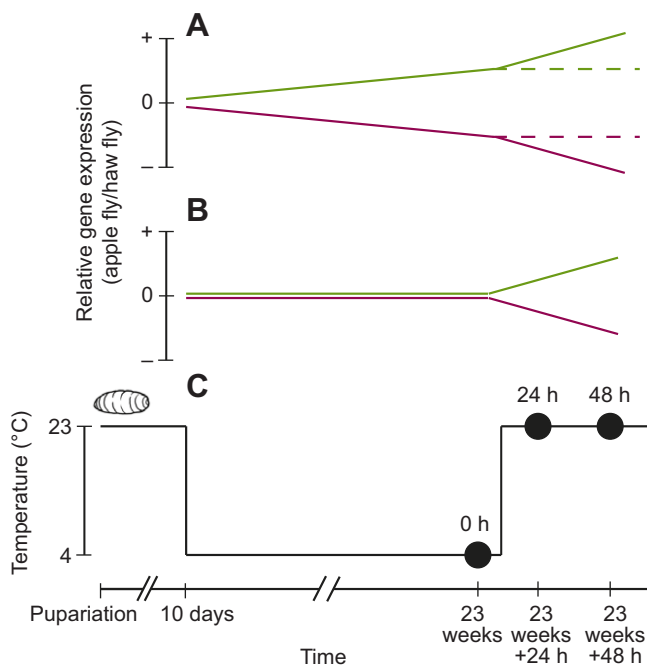


Fig. 1. Conceptual illustration of the ‘during-winter’ and ‘after-winter’ hypotheses for apple versus hawthorn fly race divergence in gene expression and the experimental design. (A,B) Transcripts regulating earlier diapause termination in apple relative to hawthorn flies could be differentially regulated between populations during winter (‘during-winter’ hypothesis; A), or only after exposure to warmer temperatures necessary to begin post-diapause development (‘after-winter’ hypothesis; B). Green and pink represent up- and down-regulation of transcripts in apple relative to hawthorn flies and not absolute values, so the key feature is when the shifts in transcription levels occur between the host races, i.e. the fork in up- and down-regulation can be seen prior to 0 h in the during-winter hypothesis and extends afterward, whereas it is initiated only after the 0 h time point in the after-winter hypothesis. Note that both positive and negative values of this relative measure may reflect up- or down-regulation of transcripts in one population compared with the other. Solid and dashed lines in the upper panel denote accelerating and constant differences between populations following the temperature shift. (C) Experimental design: upon pupariation, apple and haw fly pupae were exposed to common, simulated overwinter conditions, then sampled for RNAseq at three time points – one at the end of winter (0 h), one 24 h after the end of winter, and one 48 h after transfer to warmer conditions.

MATERIALS AND METHODS

Study system and fly rearing

Rhagoletis pomonella is a frugivorous fly native to North America that infests fruits of various hawthorn species in eastern North America (Berlocher and McPheron, 1996). After apples were introduced from Eurasia (~400 years ago), a population of these flies evolved to specialize on apple fruit during the mid-nineteenth century (Feder et al., 1988; Walsh, 1867). The derived apple- and ancestral hawthorn-infesting populations or ‘host races’ are hypothesized to be in the early stages of speciation-with-gene-flow, with natural selection maintaining genetic divergence despite ongoing gene flow (up to 4% migrants per year; Feder et al., 1994). Seasonal timing is a primary target of divergent natural selection, driven by differences in fruiting time between apples and hawthorns. At a typical site in the midwestern USA with sympatric apple and hawthorn fly host races, apple trees fruit on average 3 weeks earlier than hawthorn trees (Feder and Filchak, 1999). The fly has one generation per year, wherein adults rendezvous and mate on host fruit, females oviposit into fruit, larvae consume the fruit then exit, burrowing into the soil and entering a pupal diapause that lasts until the following growing season. Natural selection to synchronize adults with host fruit availability is very strong because the fruit is only available for a month or less, and adult flies typically only live a few weeks (Feder and Filchak, 1999). Individuals overwinter as diapausing pupae. Thus, there is strong selection for apple flies to terminate pupal diapause earlier compared with hawthorn flies, because the timing of the end of diapause determines the timing of adult emergence. Genetic association studies revealed that loci most strongly associated with eclosion timing are also the most genetically divergent between haw- and apple-infesting populations (Feder et al., 1993; Michel et al., 2010).

The goal of our experiment was to identify transcriptional differences between the apple and hawthorn host races during diapause development that may underlie their observed difference in seasonality (Fig. 1). We collected larval-infested apple fruit from East Lansing, MI, USA, on 15 September 2013, and hawthorn fruits from the University of Notre Dame campus (Notre Dame, IN, USA) on 15 October 2013. Although apple and hawthorn flies from these two collecting sites do not represent a co-occurring sympatric population pair, they are located in the same ecogeographic region at similar latitudes and the ~1 month fruiting time difference between East Lansing and Notre Dame mirrors differences observed at sympatric sites. Temperatures prior to diapause can affect diapause incidence and duration, but thermal exposure after larval wandering is of primary importance (Feder et al., 1997a; Neilson, 1962), and this exposure was controlled and equivalent for the two populations in our experimental design.

Infested fruit were placed on wire mesh trays held over plastic collecting bins in a greenhouse maintained at 23°C (natural light conditions). Wandering third instar larvae emerging from fruit were collected daily, placed in Petri dishes maintained at 85% relative humidity, and allowed to complete pupal development at 23°C for 10 days. The difference in the collection times of apple and hawthorn flies from East Lansing and Notre Dame resulted in a difference in photoperiod exposure between the host races in the greenhouse reflecting natural conditions. However, photoperiod during pre-winter development affects diapause development of *R. pomonella* minimally and does not affect pupae or account for host race differences in diapause development (Filchak et al., 2001). After 10 days, pupae were moved to a refrigerator and held at 4°C in constant darkness for 23 weeks (±3 days) to simulate winter.

Previous studies have shown that when pupae are subsequently removed from the simulated winter and exposed to spring-like temperatures (20–25°C), apple flies terminate pupal diapause earlier than hawthorn flies (Feder and Filchak, 1999; Feder et al., 1997a; Smith, 1988). We note that while a shift from cold to warm temperatures is necessary to terminate diapause, diapause does not end immediately. Pupae remain developmentally arrested and metabolically depressed for days to weeks before initiating pupal-to-adult apolysis (molting) and beginning adult morphogenesis (Ragland et al., 2011, 2009).

Experimental design

The experiment was designed to test whether the apple and hawthorn host races diverge in growth-related transcription levels during winter (during-winter hypothesis) or only after a shift from cold to warm temperatures simulating spring (after-winter hypothesis; Fig. 1). To answer this question, we compared apple versus hawthorn fly transcription profiles at three sampling time points: (1) directly out of the 4°C, 23 week winter treatment, (2) 24 h after removal from the cold and transfer to 23°C, and (3) 48 h after moving from 4°C to 23°C, hereafter ‘0 h’, ‘24 h’ and ‘48 h’ time points (Fig. 1C). Thus, transcriptional differences between the host races at the 0 h time point reflect regulatory differences induced during overwintering, while differences after the thermal shift reflect effects of post-winter warming.

Diapause developmental progression is under neuroendocrine control, so we sampled the transcriptome of only fly pupal heads containing the brain, ring gland and subesophageal ganglion. For the 0 h sample, pupae were flash-frozen in liquid nitrogen and rapidly decapitated with a sterilized razor blade, then transferred to Ambion TRI Reagent (Life Technologies, NY, USA). To obtain sufficient amounts of RNA and to account for intra-population variation in developmental progression, we pooled the heads of 10 flies from the same race in a single extraction. Heads were pestle-homogenized and stored at –80°C for no more than 4 weeks before RNA was extracted using Ambion RiboPure kits following the manufacturer’s recommendations. To confirm that pupae sampled after the shift to 23°C were still in diapause, we conducted stop-flow respirometry on each individual following Ragland et al. (2009), using a LI-COR 6252 CO₂ analyzer (LI-COR Biosciences, Lincoln, NB, USA) coupled to Sable Systems pumping and metering components (Sable Systems International, Las Vegas, NV, USA) to measure metabolic rates prior to sampling at 24 and 48 h after the temperature shift. All pupae measured at 24 and 48 h exhibited metabolic rates indicative of diapause (supplement S1.1, deposited in Dryad; see ‘Data availability’, below). After respirometry, pupae sampled at 24 and 48 h were flash frozen, decapitated, pooled and extracted as described above for the initial 0 h time point. We generated three replicate, pooled samples (10 heads per pool) for each population (apple and haw) at each time point, yielding 18 samples in total.

Library preparation, sequencing and informatics

Libraries were prepared for RNA sequencing at the Notre Dame Genomics and Bioinformatics core facility using the TruSeq RNA Sample Preparation v2 Kit (Illumina, Inc., San Diego, CA, USA). Quality control identified one library (an apple 24 h sample) of poor quality that was excluded from further analysis, leaving 17 total samples. Libraries were sequenced (100 bp paired end) across two lanes (9 and 8 samples multiplexed per lane) on an Illumina HiSeq 2000 at the Beijing Genomics Institute. After de-multiplexing, we excluded one additional sample (a haw 0 h

sample) with very low read counts from further analysis. All statistics thus reflect an analysis of 16 total samples, three replicate pools per population per treatment except for apple 24 h and haw 0 h, each of which had only two replicate pools. The generalized linear models that we describe below are valid with imbalanced replicates, though imbalance does reduce statistical power. Excluding the poor samples, we were left with 380,831,464 total paired-end reads.

Though a complete genome sequence is not currently available for *R. pomonella*, there is a published transcriptome based on 454 sequence data (Schwarz et al., 2009). We also generated additional 454 transcriptome sequence data in the current study based on pools of 100 adult apple and hawthorn fly heads (50 male, 50 female for each race) sampled directly from host trees at a sympatric site in Urbana, IL, USA. Live adult flies were individually aspirated from unsprayed apple trees at the University of Illinois Pomology area and hawthorn trees near the University of Illinois married student housing. Heads were removed, pooled, and DNA was extracted using a phenol–chloroform procedure. Libraries were generated using a Roche Lib-L emPCR kit and XL+ sequencing kit (Roche, Indianapolis, IN, USA) and sequenced on a single Roche/454 GS FLX Titanium plate (one-quarter plate per sex/host race combination). The resulting data were used to estimate allele frequency differences for SNPs between apple and hawthorn fly populations in nature, and were combined with the current Illumina RNAseq data and the previously published 454 data to produce an updated transcriptome assembly. In order to use the Trinity assembler (Haas et al., 2013), which is optimized for short reads, we first simulated 100 bp paired-end reads from the 454 data using a custom script (deposited in the Dryad archive – see ‘Data availability’, below). These simulated reads were pooled with the Illumina reads, and Trinity was run on the combined pool using default parameters. The assembly yielded 212,600 isogroups or clusters of transcripts likely to represent single transcripts with alternative splicing. We used TGICL (Pertea et al., 2003) to identify redundant sequences, but this yielded negligible improvement (only 4368 redundant sequences). Thus, we started all of our analyses with 212,600 isogroups prior to filtering (see below). Compared with the previously published assembly that contained only 27% of a conserved, Benchmarking Universal Single-Copy Ortholog set of genes compiled for Arthropoda (BUSCO set, see Simão et al., 2015), the new assembly contains 79% of arthropod BUSCOs, a substantial improvement. All sequences were annotated using blastx searches against FlyBase and UniProt databases (expect $\leq 1 \times 10^{-4}$). Analyses requiring annotations used only the FlyBase annotations, though all annotations are provided in supplement S1.2 (see supplement S1.3 for all assembly statistics; ‘Data availability’, below).

For differential expression analysis (see below), all retained Illumina reads were mapped to the new reference transcriptome using RNA star (Dobin et al., 2013) with default parameters. We then used RSEM (Li and Dewey, 2011) to count reads per isogroup (total across all possible alternative splice variants within an isogroup). Transcripts not represented by at least one count in 50% of the samples were filtered out, leaving a total of 66,235 of the initial 212,600 transcripts for downstream analysis.

Differential expression analysis

We used the edgeR package to apply a generalized linear model to the read count data, assuming a negative binomial distribution (Robinson et al., 2010). Scale factors calculated using the weighted trimmed mean of M-values (TMM) method were incorporated into the model, correcting for differences in library composition

(Robinson and Oshlack, 2010). We started by fitting a full model to each transcript that included an interaction term:

$$y_{ijk} = \mu + H_i + T_j + (HT)_{ij} + e_{ijk},$$

where host (H) and time point (T) are factors, and HT is the interaction term. Likelihood ratio tests were used to test the statistical significance of each term, applying a Benjamini and Hochberg false discovery rate (FDR) threshold of 0.05 as a significance cutoff. For all transcripts with a non-significant interaction term, we applied a reduced model excluding the interaction effect. We then used linear contrasts to test the following null hypotheses: (1) no difference between the host races, (2) no difference between the 0 and 24 h treatments, and (3) no difference between the 24 and 48 h treatments. Overlaps of these sets were visualized in an area-proportional Venn diagram using eulerAPE (Micallef and Rodgers, 2014) and tested for significance using Fisher’s exact tests.

Our test to distinguish the during-winter versus after-winter hypotheses centered on the host and host by time point interaction terms in the generalized linear model. Specifically, models for transcripts with a significant main host term but a non-significant interaction term are consistent with the during-winter hypothesis (Fig. 1A). Moreover, we predicted that such genes, if they are involved in diapause termination, should be significantly up- or down-regulated in apple versus hawthorn pupae at the 0 h treatment and that this host-associated difference should remain constant (or show the same directionality) in the subsequent 24 and 48 h samples following warming. We tested for significant differences between the host races at the 0, 24 and 48 h sampling time points using linear contrasts. In comparison, models for transcripts that have a significant interaction term are consistent with the after-winter hypothesis for a host race difference occurring after the transfer to warmer conditions. Moreover, for the after-winter hypothesis to be correct, not only should there be a significant interaction term but also transcript levels should not differ significantly between the host races at the 0 h point. Rather, they should vary between apple and hawthorn flies 24 and 48 h after the temperature shift, and in the same direction for both comparisons (Fig. 1B).

We also hypothesized that if differentially expressed sets of transcripts exhibiting host race differences during winter or after heating were involved in diapause development, they should demonstrate enrichment or over-representation of functional categories related to growth and development. To test for such enrichment, we submitted annotated lists of genes identified in the tests above to DAVID (Huang et al., 2008) for functional category (e.g. GO, KEGG, INTERPRO) enrichment analysis using the statistical procedure described at https://david.ncifcrf.gov/helps/functional_annotation.html#fisher.

454 SNP analysis

The 454 sequencing of separate pools of 100 heads each from apple- and hawthorn-infesting population at the Urbana site allowed testing for SNP allele frequency differences between the host races, using read counts to estimate population allele frequencies (Futschik and Schlötterer, 2010). We identified SNPs using the Genome Analysis Toolkit (GATK) Unified Genotyper (v3.3; McKenna et al., 2010), removing duplicate reads and filtering for a minimum phred-scaled SNP probability of 21 for bi-allelic SNPs with at least 10× coverage, and a minimum alternative allele frequency of 0.05. We then applied a Fisher’s exact test (with Benjamini and Hochberg FDR correction) to the read counts to test for allele frequency differences. Given our

filtering criterion, we applied the test to 65,793 total SNPs across 4262 transcripts. We then submitted the list of transcripts containing SNPs with significant FDR values to the DAVID tool to assess functional enrichment.

RESULTS

Differential expression between the host races

Expression patterns supported the during-winter hypothesis that the host races differ in diapause development during winter, with overwintering apple pupae appearing to be developmentally advanced relative to hawthorn pupae. Out of 66,235 total analyzed transcripts, 3002 (4.5%) showed a significant main host effect ($FDR < 0.05$) and were differentially expressed in the same direction between apple and hawthorn pupae across all three measured time points (0, 24 and 48 h; Fig. 2; supplement S1.2). These 3002 significant transcripts represent a maximum of 2823 different genes (=total transcripts minus transcripts with the same annotation). In contrast, only 371 transcripts demonstrated a significant host race \times time point interaction, as predicted by the after-winter hypothesis. Moreover, of these 371 transcripts, 142 displayed a significant difference between the host races in the 0 h treatment, suggesting that they were already differentially expressed in apple versus hawthorn flies prior to pupal heating, consistent with the during-winter hypothesis. Only 166 of the transcripts having significant interaction terms were differentially expressed between the host races only in the 24 and 48 h samples, as expected under the after-winter hypothesis (supplement S1.2 and S1.4). Thus, a total of 166 transcripts displayed a pattern that was consistent with the after-winter hypothesis compared with 3144 ($N=3002+142$) that supported the during-winter hypothesis.

The 3144 differentially expressed transcripts displaying a main host effect ($N=3002$) or host \times time point interaction accompanied by a race difference in the 0 h sample ($N=142$) were significantly

enriched for growth-related functional categories (e.g. neuron development, cell motion, gland development), suggesting that they represent differences directly involved in diapause progression (Table 1). As discussed in Materials and methods, all flies in the 24 and 48 h groups remain in diapause. Hence, the transcription differences we observed during winter likely represented preparation for or the initiation of cell differentiation and proliferation in anticipation of diapause termination, rather than overt post-diapause development. The directionality of expression of the 3144 differentially expressed transcripts further supported the during-winter hypothesis of greater preparatory or developmental activity in apple flies. Only the subset of transcripts up-regulated in apple relative to hawthorn flies ($N=1809$) was enriched for the developmentally related categories of loci; there was no enrichment for the transcripts down-regulated in apple fly pupae ($N=1335$; Table 1).

SNP analysis

From the 454 dataset comparing pools of adult apple and hawthorn flies, we identified a total of 42 ($FDR < 0.05$) and 79 transcripts ($FDR < 0.1$) that contained at least one SNP displaying significant frequency differences between the host races at the Urbana site (supplement S1.5). These transcripts were significantly enriched for genes related to oxidation-reduction functions (supplement S1.6). However, there was a potential detection bias for the SNPs displaying host differences, which had an average of 3 times greater coverage compared with non-significant SNPs. The set of transcripts differentially expressed between apple and hawthorn flies at $FDR < 0.05$ was marginally enriched for SNPs showing significant allele frequency differences between the host races at Urbana ($P=0.05$, Fisher's exact test). All told, these transcripts annotated to seven unique FlyBase genes (*hui*, *CG3902*, *Mgst1*, *Ddx1*, *CG13639*, *Non1*, *CG9917*), including two loci involved in embryonic and imaginal disc development (*hui*, *Ddx1*) and one unnamed gene (*CG3902*) exhibiting physical interactions with PI3K, a major mediator of insulin-regulated events (supplement S1.7).

Post-winter transcriptional responses

Following warming, both races undergo marked changes in expression for many transcripts that do not exhibit a host race \times time interaction. Out of the total of 66,235 transcripts analyzed in this study, 22,803 (34%) were differentially expressed between the 0 and 24 h samples. For the subset of 7557 of these 22,803 transcripts showing the most pronounced expression differences (≥ 2 -fold), particularly those up-regulated at the 24 h time point ($N=5633$), there was no evidence for any enrichment for a functional category directly related to stress (Table 1), which is known to elicit massive transcriptional responses (Gasch et al., 2000; Sorensen et al., 2005). Rather, these transcripts were highly enriched in categories related to developmental progression and cell cycling that are typically observed in *Rhagoletis pomonella* when pupal diapause ends and morphogenesis resumes (Table 1; Ragland et al., 2011). However, metabolic rate measurements taken for each fly included in the 24 and 48 h samples indicated that all pupae remained in diapause. Thus, diapause does not end immediately in either host race upon exposure to warmer temperature conditions, despite the pronounced transcriptomic response.

DISCUSSION

Winter diapause development

Transcriptome-wide gene expression data support our during-winter hypothesis (Fig. 1) that regulatory differences ultimately dictating

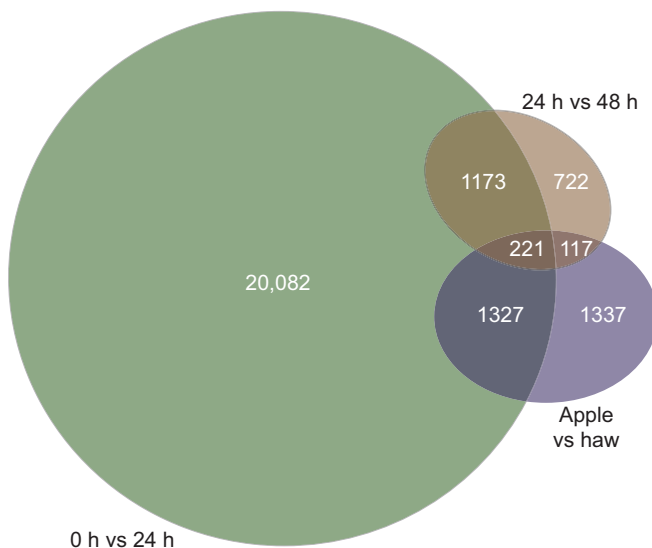


Fig. 2. Venn diagram depicting the number of transcripts differentially expressed at different time points in apple versus hawthorn flies.

Significant differential expression (false discovery rate, $FDR < 0.05$, as determined from linear models) was identified in three pairwise comparisons as inferred from generalized linear models of transcript counts including parameters for sampling time point and host race but no interaction; two comparisons between sampling time points (0 h versus 24 h and 24 h versus 48 h; Fig. 1) and one comparison between the apple and hawthorn host races (populations). Overlapping regions represent sets of transcripts that were differentially expressed in two or more comparisons.

Table 1. Functional categories enriched in sets of transcripts significantly differentially expressed (FDR<0.05) in single comparisons or pairs of comparisons

Comparison	Direction	Category	FDR	N	
Apple/haw	Both	Neuron development	0.02	53	
		Cell motion	0.06	43	
		Gland development	0.02	29	
	Up	Ribosome biogenesis	0.02	16	
		Neuron development	0.01	36	
		Gland development	0.01	22	
		Insect cuticle protein	0.08	17	
		Respiratory system development	0.02	20	
		Ectodermal gut development	0.02	10	
		Imaginal disc development	0.02	38	
	Down	None	NA	NA	
	24 h/0 h (LFC>1)	Both	Cell morphogenesis	<0.01	98
			DNA binding	<0.01	175
Transcription			<0.01	93	
Up		Imaginal disc development	<0.01	83	
		Zinc finger, C2H2-like	0.02	76	
		Regulation of cell cycle	0.01	40	
		Gland development	<0.01	41	
		DNA binding	<0.01	167	
		Imaginal disc development	0.22	69	
		Cell morphogenesis	<0.01	98	
		Transcription	<0.01	91	
		Zinc finger, C2H2-like	0.15	72	
		Regulation of cell cycle	0.01	40	
		Gland development	0.01	40	
		Cell–cell signaling	0.01	48	
		Down	Endopeptidase inhibitor activity	<0.01	17
		Aging	0.06	15	
		MADF domain	0.02	13	
Apple/haw and 24 h/0 h		Both	Neuron development	<0.01	39
			Respiratory system development	0.06	18
			Stem cell division	0.04	10
	Gland development		0.01	23	
	Ribosome biogenesis		0.02	12	
	Ribosome biogenesis		<0.01	32	
24 h/0 h and 48 h/24 h	Both	mRNA binding	<0.01	27	
		Nucleotide binding	<0.01	83	
		Helicase activity	<0.01	17	
		RNA methyltransferase activity	<0.01	8	
		Developmental growth	0.02	13	
		Ribosome biogenesis	<0.01	32	
		mRNA binding	<0.01	24	
		Helicase activity	<0.01	16	
		Nucleotide binding	0.01	64	
	Opposite	RNA methyltransferase activity	<0.01	7	
		None	NA	NA	
		Same	None	NA	NA

For single comparisons [$\log(\text{Group1}/\text{Group2})$], the direction column indicates whether the test was performed on all differentially expressed transcripts ('both') or just the subsets that were up- or down-regulated. 'Same' and 'opposite' refer to tests performed on subsets of differentially expressed transcripts with the same or opposite signs for the indicated comparisons. Because of the very large number of differentially expressed transcripts in the 24 h/0 h comparison, enrichment tests were only performed on the significant subset having absolute values of log fold-changes greater than 1 (2-fold differential expression). FDR, false discovery rate.

seasonal timing during the summer occur during the winter, long before diapause is terminated. Moreover, signals of relative up-regulation of growth and development in apple compared with hawthorn flies suggest that diapause development is generally more advanced in apple compared with hawthorn fly pupae during winter, consistent with the apple race terminating diapause earlier than the hawthorn race. Though some transcriptional differences may be the result of geographic and temporal differences between the East Lansing and Notre Dame collection sites unrelated to diapause, we consider geographic variation to be an unlikely explanation for the marked enrichment in growth and developmental transcripts observed for differentially expressed genes between the host races. Likewise, differences in diet may

influence transcription. For example, insulin signaling (discussed below) is clearly altered by diet. However, given that: (1) diet does not contribute substantially to host race differences in diapause duration (Dambroski and Feder, 2007; Smith, 1988), (2) genetic variation accounts for a large proportion of variation in eclosion timing in *R. pomonella* (Feder et al., 1993; Michel et al., 2010) and (3) a myriad of developmental processes related to morphogenesis are clearly up-regulated in apple relative to hawthorn flies (Table 1, and see following two paragraphs), we suggest that many of the transcriptional differences that we observed influence diapause duration and are underlain by genetic differences between host races. However, future time series experiments will be necessary to test whether hawthorn pupae differentially regulate the same

24 h time points tended to change direction and become down-regulated between the 24 and 48 h time points, and vice versa. Genes displaying such reversals were enriched for several functional categories related to transcription and translation (Table 1). For example, every representative member of the GO category ‘ribosome biogenesis’ was up-regulated 24 h after warming, but by 48 h displayed expression levels approaching those of 0 h (Fig. 4A). Members of the GO category ‘nucleotide binding’, which includes several important transcription factors involved in insulin signaling and cell cycle regulation connected to growth and development, showed a similar pattern (Fig. 4B). The group of transcripts that decreased in abundance 24 h after the shift to warm temperatures and then showed a rebound after 48 h also contains *pepck*, a key enzyme in gluconeogenesis that is almost universally up-regulated in the diapause responses of insects, and subsequently down-regulated following diapause termination (Poelchau et al., 2013; Ragland et al., 2010). Here, *pepck* is initially down-regulated, as would be expected at the end of diapause, but by 48 h *pepck* transcript abundance has returned to the same levels as observed during diapause at 4°C. Overall, the strong overarching pattern of ‘bounce back’ in gene expression agrees well with observed initial increases followed by gradual decreases in metabolic rate following 4°C to 23°C shifts in other *R. pomonella* diapause experiments (T.H.Q.P., D.A.H. and G.J.R., unpublished data).

Active repression of development and metabolism has significant implications for energy expenditure over the course of winter. Even during metabolic depression, diapausing insects may exhaust fuel stores vital for successful completion of development following winter (Hahn and Denlinger, 2011). Events that increase winter temperatures, such as transient warm fronts that melt insulating snow layers, may be particularly important, causing short periods of intense metabolic demand (Irwin and Lee, 2000; Williams et al., 2012a,b). However, there is evidence for active suppression of temperature-elevated metabolism during winter in insects (Williams et al., 2015). In addition to increasing metabolic demand, transient warming exposes diapausing pupae to conditions permissive for diapause termination. Thus, the most drastic fitness outcome from winter warming would likely be premature termination of diapause during a transient event that will be followed by additional days or weeks of cold, unfavorable temperatures. Suppression of growth

and development despite transiently permissive conditions is therefore critical for overwinter survival and may reflect the reversal in expression patterns of growth-related transcripts that we observed in the current study.

Comparison with other diapause studies

A rich literature on the comparative physiology of insect diapause establishes a clear connection between three major hormones, ecdysteroids, juvenile hormone (JH) and diapause hormone, and the regulation of diapause (Denlinger et al., 2005). Diapause is a physiologically dynamic process that proceeds through stages often categorized as initiation, maintenance and termination (Kostal, 2006). Hormones appear to regulate transitions between all of these stages, although the particular developmental roles and molecular interactions mediated by these hormones are often lineage specific (Denlinger, 2002). Among this clear diversity of mechanisms, however, are some commonalities in the functionally related sets of genes that have repeatedly been associated with diapause regulation across taxa.

Insulin is an important component of diapause regulation in several fly and mosquito species, where it regulates developmental arrest and nutrient provisioning. In flies (*D. melanogaster*) and mosquitoes (*Culex pipiens*), insulin signaling is an upstream regulator of juvenile hormone production, where knockdown of insulin signaling suppresses JH activity (Sim and Denlinger, 2013). A previous study in *R. pomonella* did not detect differential regulation of insulin-signaling genes during the rapid transcriptional changes associated with the end of diapause (Ragland et al., 2011). However, the regulation of diapause termination, which culminates in the end of diapause, likely involves a distinct set of genes that may or may not also change after the transition to post-diapause development. Here, we detected host race differences in the abundance of the insulin receptor *InR* and in *Tsc1*, which mediates cross-talk between the insulin and Tor pathways and has known effects on cell growth (Wullschlegel et al., 2006). After 23 weeks at 4°C, both midwestern apple- and hawthorn-infesting populations remain in diapause (Feder et al., 1997b). Therefore, the observed differences in expression between the apple and hawthorn host races occur during the maintenance and/or termination phases of diapause, and thus may reflect regulatory differences that influence when the end of diapause

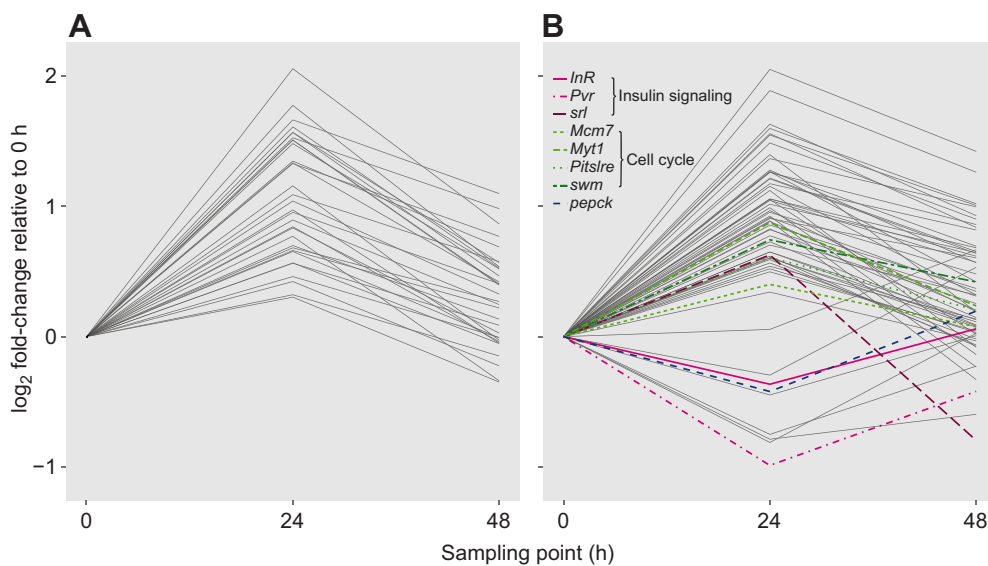


Fig. 4. Time series of transcript abundance relative to expression at the first sampling point (0 h) for transcripts consistent with a pattern of post-winter suppression.

(A) Ribosome biogenesis. (B) Nucleotide binding. Values are estimates of log₂ (expression at time point/expression at 0 h) obtained via *post hoc* contrasts of GLM parameters. These include transcripts from two functional categories that were enriched in the set of all transcripts that were oppositely differentially expressed (up-regulated at 24 h but down-regulated at 48 h, or vice versa) between (1) 0 and 24 h and (2) 24 and 48 h.

occurs and adults emerge. We observed a decrease in the expression of these insulin signaling-related genes in the apple host race, which by all other transcriptional indicators seems to be developmentally advanced relative to the hawthorn host race. In *C. pipiens* mosquitoes, an increase in insulin signaling regulates the timing of diapause termination (Sim and Denlinger, 2008). But, *C. pipiens* undergoes adult ovarian arrest, illustrating that apparently conserved diapause-related genes may act in a species-specific and stage-specific manner.

Many developmental events are coordinated by the Wnt pathway, a conserved, master regulator of tissue proliferation and patterning across animals (Reya and Clevers, 2005; Wodarz and Nusse, 1998) that has also been previously connected to diapause development. In *D. melanogaster*, *wg*, a gene in the multi-member *wnt* family, sits at the head of the so-called ‘canonical’ and calcium-dependent Wnt signaling pathways that play a central role throughout embryogenesis and morphogenesis (Wodarz and Nusse, 1998). Various studies of diapause developmental arrest implicate down-regulation of the *wnt*-related pathways as either a regulator or result of the cessation of cell differentiation and proliferation (Wadsworth and Dopman, 2015; Wodarz and Nusse, 1998). In *R. pomonella*, calcium-dependent Wnt signaling is up-regulated 1 week after transfer from 4 to 23°C, suggesting a regulatory role upstream of pupal–adult apolysis and adult morphogenesis (Ragland et al., 2011). Here, we have detected increased transcript abundance mainly of the hub gene *wg* and downstream genes of the canonical Wnt pathways (the β -catenin-dependent pathway) in apple compared with hawthorn pupae. This is consistent with developmental advancement of the apple host race, which completes diapause and emerges earlier than the hawthorn host race in the field, and mirrors the positive relationship between developmental progression and Wnt signaling in other diapause studies (Chen and Xu, 2014; Lin et al., 2009).

Conclusions

Clear patterns of differential regulation of genes and pathways implicated in growth and development suggest that: (1) regulatory differences that lead to differences in summer emergence timing across host races of *R. pomonella* likely begin during winter, in either the diapause maintenance or early termination phases, and (2) upon transfer from cold, winter-like conditions to warm, summer-like conditions, diapausing pupae appear to repress temperature-driven increases in growth and development, remaining firmly in diapause despite the permissive conditions. Key developmental genes involved in insulin and Wnt signaling may play a role in the microevolution of seasonal timing in *R. pomonella* and in other insects. While whole-transcriptome analyses are excellent at identifying large-scale physiological patterns, they can only nominate, but not confirm, individual candidate genes. Further, many transcriptional differences between populations may reflect downstream responses to upstream, unobserved regulatory events because of the difficulty in sampling at precise time points at which environmental signals are transduced. Thus, additional functional testing combined with genotype–phenotype association tests will be necessary to confirm the roles of candidate genes in diapause development and termination.

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Competing interests

The authors declare no competing or financial interests.

Author contributions

P.J.M., G.J.R., T.H.Q.P., D.A.H. and J.L.F. conceived the gene expression study, P.J.M. and G.J.R. designed the gene expression experiments, S.H.B., H.M.R. and G.J.R. conceived the 454 SNP study, P.J.M. performed the gene expression experiments, S.H.B., H.M.R. and K.K.O.W. performed the 454 SNP study, P.J.M., G.J.R., S.H.B., H.M.R. and K.K.O.W. analyzed the data, and all authors contributed to manuscript drafting.

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Data availability

All raw sequence reads have been deposited in the NCBI Short Read Archive, accessible through BioProject Accession PRJNA324814. The transcriptome assembly is deposited in the NCBI Transcriptome Shotgun Assembly archive under accession GETQ000000000.1. Supplementary tables and scripts are deposited in Dryad: <http://dx.doi.org/10.5061/dryad.51db2>.

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