Alternative splicing associated with phenotypic plasticity in the bumble bee *Bombus terrestris*

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Abstract

Phenotypic plasticity is when one genome can produce more than one phenotype. The caste system found in many social insects is an important example of plasticity. Several studies have examined gene expression in social insect developmental and caste differences. Changes in gene expression, however, are not the only source of phenotypic plasticity. Here we investigate the role of alternative splicing in the buff tailed bumble bee *Bombus terrestris*. We found that 5458 genes in *Bombus terrestris* (40%) express more than one isoform. Larvae have the lowest level of splicing events, followed by adults and then pupae. We found that when an isoform is expressed in a given caste in the larval stage, it tends to be expressed in all castes at the larval stage. The same is true at the pupal stage. However, we see more complicated interactions between the adult castes with reproductive females showing different isoform expression compared to non-reproductive females and male adults showing the most distinct patterns. We found 455 isoform switching genes, that is genes, where one developmental stage, sex or caste uses a specific isoform and another type uses a different isoform. Among genes displaying isoform switching are some involved in the ecdysteriod pathway, an important system in insect behaviour.
Introduction

A fundamental aim of evolutionary biology is to understand the link between genotype, phenotype and fitness (de Visser and Krug, 2014). A powerful way to understand this is to investigate how a single genome can give rise to multiple phenotypes. This question can be approached at the cellular level, for example, how a single genome can make both a skin cell and a blood cell, but also at the societal level, for example, how a single genome can make both the queen and worker castes in a eusocial insect colony. Indeed, recent models suggest that these different levels could even be functionally similar (Patalano et al., 2012).

In eusocial insect colonies, this differentiation between castes is fundamental, with queens carrying out most of the reproduction while workers carry out other tasks required of the colony e.g. brood care, foraging and nest maintenance. Queens are also generally bigger, live longer and have a host of behavioural and physiological specialisations compared to workers. These distinct morphological and behavioural castes are usually alternative expressions of the same genome. We previously found that the main difference in gene expression, in the buff-tailed bumble bee *Bombus terrestris*, was between castes at the adult stage rather than at larval or pupal stages (Harrison et al., 2015). A striking difference between *B. terrestris* and higher eusocial hymenopterans is that reproductive worker bees have gene expression profiles much more similar to queens than to their non-reproductive sister workers (Harrison et al., 2015).

Changes in gene expression are not the only way a single genome can give rise to multiple phenotypes. For example, the difference between head lice (harmless) and body lice, (disease vector) is caused by the alternative splicing of an identical genome not by differential gene expression (Tovar-Corona et al., 2015). Alternative splicing results in a single gene coding for multiple processed messenger RNA (mRNA), and potentially multiple proteins with distinct functions. Genome-wide analyses suggest alternative splicing as a source of phenotypic variation in eusocial insects, where a single genome must encode for numerous castes (honeybees (Foret et al., 2012; Li-Byarlay et al., 2013), possibly ants (Bonasio et al., 2012) and termites (Terrapon et al., 2014)). Bumble bees, as well as being one of the most ecologically and economically important pollinators (Woodard et al., 2015), are an interesting taxon to study the role of alternative splicing in eusociality as they possess both highly eusocial and more primitive characteristics (Harrison et al., 2015).

The diversity of forms and behaviours within a eusocial insect species is much greater than that found within solitary insect species. Yet eusocial insects have gene numbers within the range for solitary insects (Simola et al., 2013). Tovar-Corona et al. 2015 recently found the rate of alternative splicing in eight arthropod species ranging from 23% to 44%. Do bumble bees use alternative splicing more than other studied, non-social insects, in order to increase phenotypic diversity?
Here, using RNA-seq, we ask how alternative splicing is associated with caste, sex and development within the buff-tailed bumble bee, *Bombus terrestris*. We compare splicing patterns of larval (male versus workers), pupal (male versus workers) and adult (male versus non-reproductive worker versus reproductive worker versus queen) stages. Using the results of this analysis we answer a number of questions. Do bumble bees have increased rates of alternative splicing compared to studied non-social insects? Do splicing patterns follow those of gene expression with relatively fewer differences between juvenile stages compared to adults? Are reproductive workers’ splicing patterns more similar to queens’ or non-reproductive workers’? Do different castes, sexes or developmental stages use different isoforms of the same genes?
Methods

Colony sampling, RNA libraries and sequencing

Data for this study were obtained from Harrison et al. (2015). Reads were downloaded from the ENA under accession no. PRJEB9366. This contained a total of 469.3 million reads from 27 libraries. These libraries are made up of larval, pupal and adults stages of males, queens, workers (reproductive and non-reproductive). Whole bodies were used for sampling for two reasons. First, we had no prior assumptions regarding the tissues, within which genes would be alternatively spliced between castes, sexes and developmental stages. Second, this would allow us to detect as many alternatively spliced genes as possible across all comparisons. Most sample types are replicated over three colonies except for adult nonreproductive workers (two colonies), reproductive workers (four colonies) and larval, pupal and virgin queens (one colony). Sample production is described fully by Harrison et al. (2015). Due to the lack of replicates, queen larvae, pupae and virgin queen were not analysed in our current alternative splicing study.

Read processing and analysis

Raw reads were trimmed for adaptor sequence and low quality at the 5' and 3' end, and across a sliding window throughout the reads (PHRED >24) using Trimmomatic (v0.33) (Bolger et al., 2014), then visualised using FastQC (v0.11.5). Reads were aligned to the Bter_1.0 genome (Refseq accession no. GCF_000214255.1 (Sadd et al., 2015)) using Tophat (v2.1.0) (Trapnell et al., 2009), with an average of 93.3% efficiency (82.8%-95.8%). After mapping, the protocol for the tuxedo package was followed as outlined in Trapnell et al. (2012). In brief, cufflinks (v2.2.1) was performed on the alignments from tophat, then cuffmerge (v1.0.0) was used to create a consensus transcriptome. Finally cuffdiff (v2.2.1) was used to quantify isoform expression and identify statistically significant changes in isoform expression between the samples. For this study we have analysed isoforms at the CD level because they represent a change in the protein translated. Further downstream analysis of these isoforms was then performed in R 3.2.3 (core Team, 2016) using the cummeRbund (v2.16.0) package and custom R scripts (https://dx.doi.org/10.6084/m9.figshare.3201355). rMATs (v3.2.5) was run with the alignments from tophat to identify the types of alternative splicing present (Shen et al., 2014).

Isoform switching and Clustering

Relative isoform contributions for each gene were tested for differences pairwise between the samples utilising the jensen-shannon divergence test implemented by the cuffdiff program (FDR < 0.01). Allometric
differences in the amount of tissue in different castes, sexes or reproductive status could produce significant differences in isoform expression if judged solely on statistical testing. We therefore only consider isoforms with at least a log2-fold change of one in order to reduce the false-positive effects of tissue scaling (Montgomery and Mank, 2016). To identify those genes that display biologically significant isoform switching, the isoforms from significant genes identified by the jensen-shannon divergence test were subject to K-means clustering (Hartigan and Wong, 1979) (k=30-150 step of 10, Iterations = 500) based on their scaled relative abundance. The most distinct abundance patterns along with the fewest duplicated clusters were found at K of 110. Each cluster was then plotted separately to visualize isoform expression. As described by Aghamirzaie et al. 2015, clusters were then manually combined into super-clusters where they had similar relative abundance patterns.

Gene ontology (GO) analysis

We extracted the nucleotide sequence for all isoforms found in each one of the chosen super-clusters and searched for any matching sequence on NCBI using BLASTx (Altschul et al., 1990) with an E-value cutoff of 0.001. Using Blast2Go (Gotz et al., 2008), we identified gene ontology (GO) terms associated with these loci. We carried out an enrichment analysis (Fisher exact test) using a custom R script (https://dx.doi.org/10.6084/m9.figshare.3201355.v1) on each of these lists of super-cluster associated GO terms. This identified GO terms that are overrepresented (FDR < 0.1) relative to the entire transcriptome (https://dx.doi.org/10.6084/m9.figshare.3458828.v1).
Results

Isoform, CD and gene level quantification

We measured expression level in terms of fragments per kilobase of exon per million reads mapped (FPKM) of all isoforms present in published RNAseq libraries of several castes, sexes and developmental stages of *Bombus terrestris* (Figure 1). After removing genes with FPKM < 1 in all libraries there are 13548 distinct loci with expression in at least one sample. Alternative splicing produces different mRNA (isoforms) from the same gene. A subset of these isoforms will differ in their conserved protein domains (CD isoforms). At the isoform level, 8090 of the loci produce only one isoform, the remaining 5458 (40%) genes express 22786 isoforms (FPKM > 0.5 in at least one library), averaging 4.2 isoforms per gene. At the CD level 2833 (20.9%) show expression of more than one CD isoform in at least one sample, totalling 7726 CD isoforms (2.7 CD isoforms/gene). To avoid problems with assembly and annotation previously well described in alternative splicing studies (Hiller et al., 2009; Jiang and Wong, 2009), we concentrated on the 2833 genes expressing more than one CD isoform (see Table 1).

CD isoform expression varies mainly between developmental stages, but also sex and caste

Expression (FPKM) for each of the 7726 CD isoforms is highly correlated between replicates of each biological type (mated queen, reproductive adult worker, non-reproductive adult worker, worker pupae, worker larvae, male adult, male pupae and male larvae) (Figure 1a) indicating that the patterns we observe here are of biological origin.

Principle component analysis of the 7726 CD isoform’s expression (Figure 1b) shows that there is variance in the expression of these isoforms which is associated primarily with developmental stage but also sex and caste. PC1 principally describes variance in the isoform expression of different developmental stages by separating adult and pupal replicates on the left and larval replicates on the right, this difference explains 36% of the variance. PC2 is explained also by differences in developmental stages but this time separating the pupae samples from the rest (22% variance). PC3 displays the variance that exists between the sexes in the adult stages by dividing male adults and the female adult samples but also separating the female adult samples (13% variance).

Alternative splicing events can come in different forms; exon skipping, mutually exclusive exons, alternative 5' splice junction, alternative 3' splice junction and intron retention (Black, 2003). The proportions of the splicing event forms stay relatively constant between samples with only a small increase in skipped exon splice events for non-reproductive workers (see Figure S1). We observed differences in the numbers of spliced genes and the number of isoforms expressed in our samples (Table 1). Male and
Figure 1: Global splicing analysis. a. Correlogram of Pearson correlations of expression (FPKM) between replicates for the 7726 coding isoforms from multiple isoform loci. Clustering is produced based on Euclidean distances. This clustering separates replicates into stage, sex and reproductive status (coloured bars on top). b. A principal component analysis (PCA) of the 7726 isoforms from multiple isoform loci. QR = Queen Reproductive, WR = adult Worker Reproductive, WNR = adult Worker Non-Reproductive, WP = Worker Pupae, WL = Worker Larvae, MA = Male Adult, MP = Male Pupae, ML = Male Larvae.

Table 1: Total numbers of expressed features in the eight biological types examined

<table>
<thead>
<tr>
<th>Feature</th>
<th>QR</th>
<th>MA</th>
<th>ML</th>
<th>MP</th>
<th>WR</th>
<th>WNR</th>
<th>WL</th>
<th>WP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genes</td>
<td>9954</td>
<td>10043</td>
<td>9685</td>
<td>10756</td>
<td>10249</td>
<td>10418</td>
<td>9762</td>
<td>9969</td>
</tr>
<tr>
<td>Genes With Multiple Isoforms</td>
<td>5233</td>
<td>5295</td>
<td>5226</td>
<td>5338</td>
<td>5282</td>
<td>5311</td>
<td>5250</td>
<td>5265</td>
</tr>
<tr>
<td>Isoforms from Genes with multiple isoforms</td>
<td>16210</td>
<td>16840</td>
<td>16048</td>
<td>16844</td>
<td>16909</td>
<td>17090</td>
<td>16071</td>
<td>16274</td>
</tr>
<tr>
<td>Genes with multiple CD isoforms</td>
<td>2792</td>
<td>2801</td>
<td>2760</td>
<td>2800</td>
<td>2812</td>
<td>2818</td>
<td>2772</td>
<td>2782</td>
</tr>
<tr>
<td>CD isoforms from genes with multiple CD isoforms</td>
<td>6265</td>
<td>6326</td>
<td>6092</td>
<td>6349</td>
<td>6446</td>
<td>6472</td>
<td>6107</td>
<td>6208</td>
</tr>
</tbody>
</table>

Worker larvae have the lowest number of transcripts expressed in each category (genes, multiple isoform genes, multiple isoforms, multiple CD genes, multiple CD isoforms). The pupae shows a discrepancy between the sexes with the male pupae expressing more transcripts of each type than the worker pupae. Male pupae express the most genes and the most multiple isoform genes but adult workers (WR and WNR) express the most isoforms, multiple CD isoform genes and the most CD isoforms.

455 genes with varied functions display significant isoform switching

The 2833 multiple isoform loci equate to the total amount of alternative splicing at the CD level observed in our samples but not all of these loci express different proportions of isoforms in different samples, that is, isoform switching. To find these loci, we tested the 2833 multiple isoform loci pairwise for differences...
in their relative isoform abundance between samples (FDR < 0.01). 455 genes, 16% of the multiple isoform loci (2833) display significant isoform switching between samples. These 1303 CD isoforms from 455 genes were analysed further to identify biologically significant isoform switching events.

We used K-means clustering to identify clusters of isoforms displaying similar expression patterns across biological types. 297 (from 1303) were not clustered because they have small changes in their relative abundance between samples (range < 0.2) and would be misrepresented when their isoform abundance was scaled before clustering. K-means clustering (K = 110) of the 1006 remaining isoforms achieved separation of most isoforms by their relative abundance (Figure S2). By manually combining clusters with the same pattern of relative abundance (Figure 2) we identified 26 super-clusters that show a high degree of sample-specific expression (658 isoforms). The remaining 348 isoforms showed cryptic abundances and could not be assigned to a super-cluster, these isoforms are included in the variable cluster (Figure 2). 22 of the 26 super-clusters form antagonistic pairs, where both super-clusters represent the high and low abundance isoforms for that particular sample or group of samples. Most super-clusters (491 isoforms) show preferential high or low expression in samples(s) from one developmental stage and four clusters (167 isoforms) have preferential expression in samples from more than one developmental stage. The majority of isoforms show adult sample(s) specific expression (374) with 105 for pupae and 76 for larvae (supplementary data: isoform_cluster_database). Of the 455 isoform switching genes; 328 have multiple isoforms in multiple super-clusters, 39 have isoforms solely in the variable cluster, 25 genes have one or zero isoforms in a super-cluster and 63 are uncharacterised loci. From the 328 genes that have multiple isoforms in multiple super-clusters 80 genes have their isoforms clustered in antagonistic pairs.

For each super-cluster and groups of super-clusters relating to each developmental stage we tested for enriched gene ontology (GO) terms against their background value in the transcriptome. No discrete cluster had significant enriched GO terms at the FDR <0.1. However, twenty three GO terms were associated with isoform switching genes found in adults (supplementary data: cluster_GO_terms). Two of these, steroid hormone mediated signalling pathway (GO:0043401) and steroid hormone receptor activity (GO0003707) relate to important functions in social insect biology. A number of isoform switching genes are associated with these terms, including; ecdysone inducible protein 74EF (LOC100651988), E75 (LOC100644185), E78 (LOC100646780), ecdysone receptor (LOC100646757) and farnysl pyrophosphatase synthase-like (LOC00165005). The majority of these genes differ in their isoform composition between one or more adult sample. As an example of one of these genes, we show the expression, exon map and protein domains for ecdysone inducible protein 75 (Figure 3). In this gene, transcript variant X1 (XM_003398944.2) has high levels of expression especially for non-reproductive females compared to transcript variant X2 (XM_003398942.2) (see Figure 3b). Expression plots and exon maps
**Figure 2:** Super-clusters showing abundance patterns around particular biological sample types. K-means clustering for the 1006 isoforms from differentially spliced genes. Y-axis displays scaled relative abundance (-2 to 2). QR = Queen Reproductive, WR = adult Worker Reproductive, WNR = adult Worker Non-Reproductive, WP = Worker Pupae, WL = Worker Larvae, MA = Male Adult, MP = Male Pupae, ML = Male Larvae. The labels in the banners above figures identify the biological types in which the cluster is differentially expressed in. For example, Male Pupae (second column, second row) shows two clusters, one where there is a spike in expression in male pupae and one where there is a drop.
(10.6084/m9.figshare.5208940) are available in the supplementary data for all 455 isoform switching genes along with a database containing their annotation (supplementary data: isoform_cluster_database), their isoform’s super-cluster membership (supplementary data: Clusters in super clusters) and GO terms (supplementary data: cluster_GO_terms).
Figure 3. a) Expression (FPKM) of the two transcripts of Ecdysone-inducible protein E75 over the different castes and developmental stages. Circles represent the means, the error bars are standard errors of the mean. QR = Queen Reproductive, WR = adult Worker Reproductive, WNR = adult Worker Non-Reproductive, WP = Worker Pupae, WL = Worker Larvae, MA = Male Adult, MP = Male Pupae. b) Exon map of the two transcripts. c) Protein domains encoded by the two transcripts. NR_LBD_REV_ERB = ligand binding domain of REV-ERB receptors, NR_DBD_REV_ERB = DNA-binding domain of REV-ERB receptors.
Discussion

Alternative splicing is a way to increase the variety of proteins produced by a given number of genes. One aim of this paper was to ask do bumble bees have increased rates of alternative splicing to account for the increased phenotypic diversity within a eusocial insect species despite them having gene numbers within the range for solitary insects (Simola et al., 2013). We observed 2833 genes that have expression of more than one CD isoform in the samples examined. This corresponds to 20% (2833/13548) of the bumble bee genes producing different proteins from the same genotype. To compare Bombus terrestris with current literature we also looked at the number of genes that express more that one isoform regardless of the protein produced (Tovar-Corona et al., 2015). Compared to the rate of alternative splicing in eight arthropod species recently studied (23% to 44%) (Tovar-Corona et al., 2015), bumble bees’ rate of 40% is at the higher end. However this rate is equivalent to those of two other insects (Drosophila melanogaster 42% and the pea aphid, Acyrthosiphon pisum 44%). Further work in other insects will be needed to confirm this, but our data indicates that an increased rate of alternative splicing is not required by bumble bees.

We then examined how the bumble bee uses these isoforms. We found four hundred and fifty five genes express different proportions of isoforms in different samples, that is, isoform switching. We have annotated and ranked these genes according to their super cluster membership, conserved domain changes in their isoforms, GO terms and gene expression status, providing a rich resource for future studies in this and related species. Patterns of alternative splicing follow those previously found for gene expression (Harrison et al., 2015). This is not due to the effect of overall expression itself, as we only looked at genes that have expression of at least two isoforms, removing almost 80% of the observed genes (13548 - 2,833). Even still, within the 455 isoform switching genes, 45% also show gene expression changes. This exemplifies how alternative splicing studies can both add to the knowledge of genes known to be differentially expressed and also find novel source of plasticity that could not be found by only examining gene level expression.

We found relatively little differentiation between the sexes at the larval stage when compared to adult stages (see Figures 1 and 2). These findings suggest a comparatively low number of isoforms are associated with the creation of distinct morphological sexes compared to the high number involved in distinct behaviours between adult males and females. Reproductive workers show more similarity to queens than to their fellow, but non-reproductive, workers (see horizontal dendrograms Figure 1a). We have hypothesised that this is due to the plastic nature of bumble bee workers which allow them to compete with the queen at the end of the annual colony cycle (Harrison et al., 2015). Male adults are the most distinct group in terms of isoform expression (Figure 1), a finding that reflects their similarly
distinct behaviour, physiology and life history.

No GO terms were found to be significantly associated with any of the super clusters. However, twenty
three GO terms were associated with isoform switching genes found in adults. Two of these, steroid hor-
mone mediated signalling pathway (GO:0043401) and steroid hormone receptor activity (GO0003707)
relate to the ecdysteroid pathway, an important system in insect biology (Niwa and Niwa, 2016). Ecdys-
teroids are key compounds involved in ovary activation, regulating agonistic behaviour and establishing
the dominance hierarchy in bumble bee workers and queens (Geva et al., 2005).

In the results, as an example of an isoform switching gene, we expanded our analysis for one ecdysone
related gene (Ecdysone-inducible protein E75 (LOC100644185)). Ecdysone-inducible protein E75 is a
transcription factor in the ecdysone cascade (Li et al., 2015). Transcript variant X1 (XM_003398944.2)
has high levels of expression especially for non-reproductive females compared to transcript variant X2
(XM_003398942.2) (see Figure 3b). In Drosophila melanogaster, different isoforms of E75 have oppo-
site roles in controlling female reproduction with, for example, DmE75A inducing apoptosis in the egg
chamber, while DmE75B inhibits DmE75A function allowing egg development (Terashima and Bownes,
2006). The difference between the transcripts is a number of exons encoding for binding domains of the
nuclear receptor REV-ERB (Figure 3c). REV-ERB receptors are transcriptional regulators belonging
to the nuclear receptor superfamily. They regulate a number of physiological functions including the
circadian rhythm, lipid metabolism, and cellular differentiation. The ligand binding domain (LBD) of
REV-ERB receptor-like, found in both isoforms, is responsible for coactivator interaction with steroid
hormones (Raghuram et al., 2007). The DNA-binding domain (DBD), found only in transcript variant
X2, interacts with specific DNA sites upstream of the target gene and modulates the rate of transcrip-
tional initiation. This truncation of a binding domain is a very simple way for a gene to go from
being expressed specifically to a more global pattern of expression. We found this truncation in two
other genes we examined in depth, the CXXC zinc finger domain in Methylcytosine dioxygenase TET2
(LOC100642293 Figure S3) and the RNA recognition motif in Insulin-like growth factor 2 mRNA-binding
protein 1 (LOC100647990 Figure S4). Further study of our data set and others could test if this is a
common phenomenon.

We have shown that alternative splicing of genes is important for bumble bee biology. How is this
alternative splicing is controlled? Alternative splicing in some social insect species has been shown to be
affected by DNA methylation (Bonasio et al., 2012; Foret et al., 2012). Methylation has also been shown
to play a role in social insect biology (Yan et al., 2015) including bumble bee reproductive division of
labour (Amarasinghe et al., 2014). Together these observations suggest that a fruitful area of research
for social insect biologists would be in understanding the role of methylation in controlling alternative
splicing.
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References


Data Accessibility,

All sequence data for this study are archived at European Nucleotide Archive (ENA); Accession no. PRJEB9366 (http://www.ebi.ac.uk/ena/data/view/PRJEB9366). GO-analysis results and lists of isoforms are available as Supporting Information.

Author contributions

All authors developed the project idea and were involved in the interpretation of data and finalization of the manuscript. JP and EBM analyzed the data. EBM and JP drafted the manuscript.