Insect immune specificity in a host-parasite model

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Abstract

Ecological studies routinely show host-parasite genotype-genotype interactions in insect systems. The mechanisms behind these interactions are not clearly understood. Using the bumblebee *Bombus terrestris* / trypanosome *Crithidia bombi model system*, we have carried out a transcriptome-wide analysis of gene expression in bees during *C. bombi* infection. We have performed three analyses, comparing expression in infected and non-infected bees 24 hours after infection by *Crithidia bombi*, expression at 24 and 48 hours after *C.bombi* infection and most importantly searching for differential gene expression associated with the host-parasite genotype-genotype interaction at 24 hours after infection. We found a large number of genes differentially regulated related to numerous canonical immune pathways. These genes include receptors, signaling pathways and effectors. We discovered a possible interaction between the peritrophic membrane and the insect immune system in defense against *Crithidia*. Most interestingly we found differential expression of *Dscam* depending on the genotype-genotype interactions of the given bumblebee colony and *Crithidia* strain.

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Ecological studies with a number of natural, co-evolving host-parasite systems have shown that resistance to a parasite is highly variable in invertebrates, in part determined by the interaction of the genotypes of the host and the parasite (Schmid-Hempel 2001; Carius et al. 2001; Lambrechts 2010). Such interactions can lead to the evolution and maintenance of genetic variation in natural populations (Hamilton et al. 1981). Many human diseases use insects as vectors (Jacobslorena & Lemos 1995). Any effort at control will require a better understanding of this relationship between host and parasite. How this level of specificity is generated is unclear. Genotype-genotype interactions quantified by ecological measures of disease resistance (e.g., host mortality, fecundity and infection rate) cannot explicitly test whether the immune response produces this level of specificity (Hauton & Smith 2007). It may be that such outcomes can be explained by other processes. For example, the bumblebee, Bombus terrestris/trypanosome Crithidia bombi system is a well studied example of these ecological host x parasite genotype-genotype interactions (Schmid-Hempel et al. 1999; Schmid-Hempel & Reber-Funk 2004). Yet even here it has been shown that independent of host genotype, specific isolates of gut microbiota from different hosts are protective against particular parasite genotypes (Koch & Schmid-Hempel 2012). Still there is evidence that the immune system must have a role in generating this host-parasite specific response. A number of studies have found differential immune genes expression in response to Crithidia (Schlüns et al. 2010; Riddell et al. 2011; Brunner et al. 2013; Barribeau & Schmid-Hempel 2013). We found increased Crithidia loads in bees whose expression of antimicrobial peptides was knocked down by RNAi (Deshwal & Mallon 2014). We have even shown that bees from different host genotypes induce differential expression of antimicrobial peptides (AMPs), according to the strain of C. bombi they had been infected with (Riddell et al. 2009), that is we found specificity in the immune response itself.

 Here, we expand our previous study and carry out a transcriptome-wide analysis of gene expression in bees during *C.bombi* infection. We have carried out three analyses, comparing a) expression in infected and non-infected bees 24 hours after infection by *Crithidia bombi* b) expression at 24 and 48 hours after *C.bombi* infection and c) looking for differential gene expression associated with the host-parasite genotype-genotype interaction at 24 hours after infection. Enrichment analysis was also carried out on expression data to see which categories of molecules are differentially regulated during infection. The results confirm our previous findings of up-regulation in antimicrobial peptide expression and provide a comprehensive overview of changes in and the specificity of gene expression after exposure to 2 strains of *C.bombi*.

61 Methods The samples used during this experiment are taken from Riddell et al. 2009 (Riddell et al. 2009). 62 We have chosen samples that showed a reciprocal pattern of expression for the three antimicrobial 63 64 peptides (AMPs) tested in that paper. These were colony K5 (called K from now on) and Q1 (Q) and strains 6 and 8. K-8 showed a high AMP expression, Q-8 a low expression level, Q-6 a high 65 66 level and K-6 a low level of AMP expression. Sample Collection 67 68 Experiments were carried out on one commercially reared bumblebee colony from Koppert 69 Biological Systems U.K. (Colony K) and one colony from wild caught queens (Colony O). All parasite isolates used originated from wild queens collected in Spring 2008 in the botanical gardens, 70 71 University of Leicester. Experiments began when the colonies had a minimum of thirty workers, approximately four weeks old. Between observations, colonies were fed ad libitum with pollen 72 73 (Percie du sert, France) and 50% diluted glucose/fructose mix (Meliose – Roquette, France). Before and during the experiments colonies were kept at 26°C and 60% humidity in constant red light. 74 75 **Infections** 76 To prepare C. bombi isolates, faeces was collected from workers of naturally infected colonies, and mixed with 50% diluted Meliose to create a standardized dose of 500 Crithidia cells per ul of 77 78 inoculum. Previous studies had shown that such inocula, prepared from different colonies, are 79 genotypically different (Schmid-Hempel & Reber-Funk 2004) and generate specific responses in novel hosts (Schmid-Hempel et al. 1999). We infected a sample of workers from each of K and Q 80 81 bumblebee colonies (representing different host lines) with an inoculum of faeces from each of the 82 two wild infected colonies (6 and 8 Crithidia strain). We also collected uninfected controls. Bees were four days old at the time of infection. After infection bees were kept in colony x strain groups 83 84 (1–3 individuals depending on day collected) and fed ad libitum. 24 hours or 48 hours post infection

the bees were sacrificed by freezing in liquid nitrogen. They were then stored at -80°C.

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RNA sample preparation and sequencing

Total RNA was extracted from 23 individual homogenised abdomens using Tri-reagent (Sigma-Aldrich, UK). Any residual contaminants were removed from the RNA using the RNeasy mini kit (Qiagen, UK) and manufacturer's RNA clean-up protocol. To remove residual genomic DNA, RNA samples were treated with DNase (Sigma-Aldrich, UK). TruSeq RNA-seq libraries were made from the 23 samples at NBAF Edinburgh. Sequencing was performed on an Illumina HiSeq®2000 instrument (Illumina, Inc.) by the manufacturer's protocol. Multiplexed 50 base single-read runs were carried out yielding an average of 12M reads per sample. Statistical analysis The reference transcriptome was downloaded from http://www.nematodes.org/downloads/databases/Bombus terrestris/ 2011). (Colgan al.Functional annotation related to the transcriptome was obtained using the BLAST2GO package (Götz et al. 2008). Alignment was done using GSNAP (version 2012-07-20) (Wu & Nacu 2010). Only reads that mapped uniquely were selected for further analysis. Counts were generated per transcript for each sample. Differential expression analysis was performed using the edgeR (3.4.0) package (McCarthy et al. 2012) in R (3.0.1) (R Core Team 2013). Normalization factors were computed using the TMM technique, after which tagwise dispersions were calculated and subjected to a generalized linear model (GLM). Resulting p values were subjected to Benjamini-Hochberg multiple testing correction to derive FDRs; only transcripts with a FDR < 0.05 were considered for further analysis. Three separate GLMs were carried out, one looked for transcripts that are differentially expressed upon infection with *Crithidia* at 24 hours post-infection (~0+colony+infect(yes/no)) infect here are bees either infected with strain 6 or 8, one looking at the gene expression difference between 24 hours and 48 hours post strain 6 infection (~0+colony + time) and a further GLM that looked for

transcripts that were expressed in a specific pattern at 24 hours post-infection (\sim 0+colony*strain). Using Blast2Go, we then carried out an enrichment analysis (Fisher exact test) on each of these lists of differentially expressed genes to see which GO terms are overrepresented relative to the entire genome. We then used REVIGO to summarize and visualise these terms (Supek *et al.* 2011). For each of the lists of differentially expressed transcripts we also carried out a blastx analysis against the insect innate immunity database (IIID) (Brucker *et al.* 2012). We used the BLOSUM62 matrix with a word size of 3. The results were filtered to only contain hits with an *E*-value <1e-10, a bit score \geq 30,

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Results Genes differentially expressed at 24 hours post-infection 31,843 unique transcripts were mapped to the transcriptome. 489 transcripts were found to be differentially expressed 24 hours post-infection (FDR < 0.05), 324 were downregulated and 165 upregulated. Reannotating the transcripts using Blast2GO (blastx against the nr database with e < 0.001), 109 had no BLAST hits. A further 68 had uninformative BLAST hits (anonymous predicted protein). The remaining 312 were used in the enrichment analysis. Figure 1 shows a summary of the enriched GO terms found (Fisher's test p < 0.05). Defense response (GO:0006952, FDR = 0.047) and chitin metabolism (GO:0006030, FDR = 0.032) were the only processes significantly enriched at a more stringent level (FDR < 0.05). Peritrophic membrane: The peritrophic matrix (PM) forms a layer composed of chitin and glycoproteins that lines the insect midgut lumen (Kuraishi et al. 2011). The PM facilitates digestion and forms a protective barrier to prevent the invasion of ingested pathogens (Lehane 1997; Kuraishi et al. 2011). Fibrillin 1 (BTT14121 1), a venom protein precursor (BTT32193 1), Neurotrypsin (BTT07956 1), Peritrophin-1-like (BTT01709 1, BTT22959 1, BTT37215 1, BTT42262 1) and four chitinase transcripts (Chitinase 3: BTT23997 1 BTT38724 1, Chitinase 4 BTT20684 1, BTT23469 1) are downregulated upon infection. Fibrillins are extracellular matrix macromolecules, ubiquitous in the connective tissues (Isogai et al. 2003). BTT32193 1 was classed as a venom protein, but was also very similar to Chitinase 3 (blastx $e = 1e^{-16}$). Chitinases modulate the structure and porosity of the PM (Dinglasan et al. 2009). Neurotrypsin is a serine protease expressed in the nervous system (Gschwend et al. 1997). However in the protease domain it shares similarities with Sp22D, a chitin binding serine protease (Danielli et al. 2000). The chitin fibrils of the PM are assembled into a wide cross-hatched pattern connected by peritrophins (Dinglasan et al. 2009). A second group made up

of Peritrophin-1 (BTT05886 1, BTT20661 1) and 3 further chitinase transcripts (Chitinase 2

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:BTT23246 1, Chitinase 3: BTT39163 1, Chitinase 4: BTT05313 1) is upregulated. Figure 2 shows the correlation of expression patterns between these sixteen transcripts related to chitin metabolism. There is some clustering, but not of any clear functional groups. Taken together however, this differential expression suggests an important role for the repair or restructuring of the peritrophic matrix in the bumblebees' response to *Crithidia*. When the BLAST searches against the IIID and nr databases are combined, eighty nine transcripts relate to canonical insect immune genes. We describe them in the order receptors, serine proteases, signalling pathways and effectors (Schmid-Hempel 2005). Receptors: The Down syndrome cell adhesion molecule (Dscam), a pattern recognition receptor has come to the forefront of research into insect immune specificity as it has been found to have thousands of different splice forms and is associated with insect immunity (Smith et al. 2011). We found five downregulated transcripts annotated as immunoglobulin superfamily (Dscam included in hit list) (BTT03519 1, BTT08682 1, BTT15814 1, BTT26724 1, BTT27678 1) and one upregulated transcript (BTT03519 1). Serine proteases: Serine proteases are important proteolytic enzymes in many molecular pathways. When these serine proteases are no longer needed, they are inactivated by serine protease inhibitors (Zhao et al. 2012). CLIP domain serine proteases mediate insect innate immunity (Zou et al. 2006). Twenty one transcripts related to serine proteases, serine protease homologues or serine protease inhibitors were differentially expressed upon infection (see Table 1). Lipophorin receptor 2 (downregulated BTT34617 1) binds with serpins to aid in their encytocytosis (Soukup *et al.* 2009).

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Signalling pathways: We found a transcript for Spatzle (BTT19738 1) downregulated at this time point. Activation of the Toll immune pathway requires the activation of Spatzle (Lemaitre & Hoffmann 2007). MyD88 (upregulated BTT15687 1) is a death domain-containing adaptor activated by Toll leading to the activation of *Pelle*. *Dorsal* (BTT25273 1) was also downregulated. The nuclear translocation of Dorsal, a member of the NF-kB family, in the Toll pathway induces the expression of many immune genes. We found an upregulated transcript (BTT09662 1) for *Helicase89B* part of the Toll and Imd Pathway. It is required downstream of NF-kB for the activation of AMP genes in Drosophila melanogaster (Yagi & Ip 2005). ird5 codes for a catalytic subunit of an IkappaB kinase that cleaves Relish. Relish (Imd pathway) is an essential regulator of antimicrobial peptide gene induction. We found *ird5* (BTT03904 1) to be downregulated 24 hours post-infection. In mammals semaphorins are crucially involved in various aspects of the immune response (Takamatsu & Kumanogoh 2012). A semaphorin-5A-like transcript (BTT01850 1) was downregulated 24 hours post-infection. Semaphorin regulates the activity of Ras-family small GTPases (Takamatsu & Kumanogoh 2012). A Ras-like protein11B transcript (BTT05368 1) was also down regulated. The Ras/MAPK pathway was found to be essential for the suppression of the Imd immune pathway in *Drosophila* (Ragab et al. 2011). The downregulated Drumstick (BTT13062 1) interacts with the JAK/STAT pathway during its' development role (Johansen et al. 2003), but we could not find any information about its immune role. Two transcripts (BTT11590 1, BTT14205 1) of *Puckered* were downregulated. *Puckered*, which codes for a dual specificity phosphatase, is a key regulator of the c-Jun-N-terminal kinase (JNK) immune pathway (Karkali & Panayotou 2012). Mpk2/p38a (downregulated BTT05769 1) is involved in the JNK Pathway and JAK/STAT Pathway. Heat-shock factor activation by p38 is a recently discovered part of antimicrobial reactions in flies (Chen et al. 2010). We found two heat

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shock protein transcripts (BTT23758 2, BTT37030 1) and one other (BTT17701 1) that were downregulated and upregulated respectively. These are all involved in the JAK/STAT pathway. Effectors: Our previous paper (Riddell et al. 2011) found that antimicrobial peptides were upregulated at 24 hours post-infection. We would expect the same to be true here. Indeed, we found 5 transcripts for defensin (BTT06274 2, BTT8490 1, BTT10405 1, BTT14019 1, and BTT42034 1) and 3 transcripts for hymenoptaecin (BTT18071 1, BTT24170 1, BTT24170 2), all upregulated. An apidaecin precursor (BTT33652 1) was downregulated. Apidaecin has recently been shown to be expressed in bumblebees (Colgan et al. 2011). The downregulated beta-amyloid-like protein (BTT20240 1) has been shown to be an antimicrobial peptide in mammals (Soscia et al. 2010). Hemolectin (BTT15326 1, upregulated) is a clotting protein known to have a role against gram negative bacteria (Lesch et al. 2007). Reactive oxygen species (ROS) are generated by respiration in the mitochondria or as part of the immune response (Molina-Cruz et al. 2008). P450 cytochromes are oxidases acting terminally in monooxygenase systems (Felix & Silveira 2012). Some are regulated in response to infection possibly either as direct immune responders (Vlachou et al. 2005), producing nitric oxide (NO) or other reactive oxygen radicals or as part of the host detoxification process decreasing oxidative stress after an infection (Molina-Cruz et al. 2008). A number of cytochromes P450 were differentially expressed 24 hours post infection. Ten cytochrome p450 transcripts (Cvp4p3: BTT05294 1, BTT20848 1, BTT22253 1, BTT23317 1, BTT32674 1, cytochrome P450 4g15: BTT23811 1, BTT32459 1, cytochrome P450 6k1: BTT35547 1, BTT40653 1, cytochrome P450 6a14: BTT38445 1) were found to be downregulated. Three other cytochrome P450 transcripts (Cyp4p3: BTT21216 1, BTT35543 1, cytochrome P450 315a1: BTT26726 1) were upregulated. Several other cytochromes (cytochrome b: BTT20524 1, BTT39776 1, BTT41896 1, and

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cytochrome c: BTT05255 2) were downregulated. Numerous other actors in the production of ROS were found to be differentially expressed. TPX4 (BTT13285 1), coding for a Thioredoxin-dependent peroxidase, was downregulated. This gene was found be differentially expressed during *Plasmodium* infection in *Anopheles gambiae* (Baton et al. Thioredoxin-dependent peroxidase detoxifies H₂O₂. Calcineurin BTT26273 1) was found to be downregulated 24 hours post-infection. This agrees with our previous findings (Riddell et al. 2011). In infected D. melanogaster larvae, NO signals are enhanced by Calcineurin to promote induction of strong, robust immune responses via the Imd signalling pathway (Dijkers & O'Farrell 2007). We found downregulation of sortilin-related receptor-like (BTT31654 1). In mammals, sortilin aids in phagocytosis (Wähe et al. 2010). Two downregulated transcripts (BTT35021 1, BTT08756 1) were matched to *croquemort*. Croquemort, which codes for a scavenger receptor is a key part of the Imd pathway but in its apototic phagocytosis role not its immune one (Franc et al. 1999). Annexin IX (downregulated BTT02025 1) has been shown to be induced by septic injury in *Drosophila*. It is thought to encode for an anticoagulant (Gregorio *et al.* 2001). Miscellaneous: Major royal jelly protein (BTT05317 2, BTT36365 1 upregulated) has been shown to have antimicrobial properties and to be expressed in response to bacterial infection in honeybees (Scharlaken et al. 2007; Buttstedt et al. 2013). Vitellogenin (downregulated BTT36006 1) is a potent regulator of the immune response in honeybees (Amdam et al. 2004). Several orthologs of putative *Drosophila* immune loci were found to be differentially expressed 24 hours post-infection (CG12505: BTT00934 1, CG18348: BTT04397 1, CG7296: BTT15035 1, BTT18395 1, CG8791: BTT18908 1, CG5527: BTT35653 1, Fst: BTT11511 1). The downregulated CG4393

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(BTT05817 1) is weakly analogous to TNF receptor associated factor 3 (TRAF3) that mediates signal transduction involved in mammalian immune responses. Downregulated BTT37289 1 codes for a putative fatty acyl-CoA reductase. Genes differentially expressed between 24 hours post-infection and 48 hours post-infection 43 transcripts were found to be differentially expressed between 24 hours post-infection and 48 hours post-infection. Of these 17 had no BLAST hits. A further six had uninformative BLAST hits (anonymous predicted protein). The remaining 20 were used in the analysis. Defense response was the only GO term significantly enriched (FDR= 0.00015), with seven transcripts. Three transcripts correspond to Hymenoptaecin (BTT18071 1, BTT24170 1,BTT24170 2). They were all upregulated. This suggests a continuing strong AMP production 48 hours after infection. This agrees with other immune assays in bumblebees (Korner & Schmid-Hempel 2004). Argonaute-2, a RNA-silencing endonuclease, is involved in antiviral defense in insects (downregulated BTT02484 1) (van Rij et al. 2006). GstD8, a glutathione S-transferase, is involved in the detoxification process (upregulated BTT04810 1) (Gerardo et al. 2010). Dopa decarboxylase (upregulated BTT28048 1) converts L-dopa to dopamine during the melanisation process (Ferdig et al. 2000). SCR-B9 (upregulated BTT40924 1) codes for a scavenger receptor protein. Scavenger receptor proteins have been found to be microbial pattern recognition receptors in flies (Rämet et al. 2001). Genes differentially expressed depending on host genotype – parasite genotype interactions There were 591 differentially expressed transcripts (FDR < 0.05). Reannotating the transcripts using Blast2GO (blastx against the nr database with e < 0.001), 150 had no BLAST hits. A further 64 had uninformative BLAST hits (anonymous predicted protein). There were 109 transcripts that

had previously been found to be differentially expressed at 24 hours post infection. Figure 3 shows

a multidimensional scaling (MDS) plot of the samples based on the expression of these 591 genes.

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It can be clearly seen that the 11 samples are grouped into their colony-strain interaction. Of the 591 transcripts, 132 were upregulated and 459 were downregulated. Up or downregulation does not have the same meaning here as in the infected versus uninfected model were there was a clear baseline (uninfected). Depending on how you order the GLM we could get the reciprocal result. Our model used colony K strain 8 as the final contrast. From our previously published qPCR data (Riddell et al. 2009), we know the colony K strain 8 interaction displayed the highest levels of AMPs (effectors). Therefore when we say a transcript is upregulated, we mean it is upregulated in this high immune response interaction. As with the infection data, we combined the BLAST searches against the IIID and nr databases. Ninety transcripts correspond to canonical insect immune genes. We again describe them in the order receptors, serine proteases, signalling pathways and effectors (Schmid-Hempel 2005). Receptors: Two transcripts were associated with gram negative binding proteins (upregulated GNBP, BTT03533 1 and downregulated GNBP1-2 BTT35513 1) Although, as their name suggests, GNBPs are most associated with defense against gram negative bacteria, they have been show to have a role in respond to *Plasmodium* infections (Tahar et al. 2002). C-type lectins (CTLs) bind carbohydrates and mediate processes including pathogen recognition (Cirimotich et al. 2010). CTL4 is agonist to *Plasmodium* infections in mosquitoes (Cirimotich et al. 2010). A CTL4 transcript (BTT29328 1) was found to be downregulated. One downregulated transcript was related to *Dscam* (BTT12755 1). A further fourteen downregulated transcripts were part of the Ig superfamily (IGFn3-1: BTT05561 1, BTT05581 1, BTT08682 1, BTT12655 1, BTT13442 1, BTT14516 1, BTT18750 1, BTT21156 1,

305 BTT22598 1, BTT22819 1, BTT23339 1, BTT24070 1, IGFn3-7: BTT08109 1, BTT09498 1) 306 and one was upregulated (IGFn3-8: BTT03519 1). Dscam and most of the other Ig superfamily 307 transcripts cluster together in the top right of figure 4, suggesting they are similarly expressed. 308 309 Serine proteases: 310 28 transcripts related to serine proteases, serine protease homologues or serine protease inhibitors 311 were differentially expressed (see Table 2). 312 313 Signalling pathways: 314 The Toll-like receptor 18Wheeler (BTT35732 1) was upregulated as was Toll 10 (BTT09386 1). 315 18Wheeler has been shown to be important in the anti gram-negative immune response in 316 Drosophila larvae (Ligoxygakis et al. 2002). Dorsal IA (BTT04010 1), a transcription factor that 317 is a fundamental part of the Toll pathway, was downregulated. A transcript for Spatzle 1-2 was 318 downregulated (BTT10679 1). 319 320 The tyrosine kinase Pvr (BTT04822 1), which inhibits JNK activation (Bond & Foley 2009) was 321 downregulated. Jun, a transcription factor of the JNK pathway was downregulated (BTT13636 1). Mpk2/p38a (downregulated BTT16580 1) and MAPKKK9 (downregulated BTT04404 1) are 322 323 mitogen-activated protein kinases involved in the JNK Pathway and JAK/STAT pathways. We 324 found two heat shock protein transcripts (BTT17371 1, BTT22195 1) and one other (BTT17701 1) 325 that were downregulated and upregulated respectively. These are all involved in the JAK/STAT 326 pathway. Akt 1 (downregulated BTT14188 1) is part of the insulin/insulin-like growth factor 1 327 signaling (IIS) cascade. IIS plays a critical role in the regulation of innate immunity. Activation of 328 Akt signaling leads to a decrease in malaria infection intensity in mosquitoes (Corby-Harris et al. 329 2010). 330

331 Effectors: 332 Five transcripts relate to the AMPs defensin (BTT06274 2, BTT42034 1) and hymenoptaecin 333 (BTT18071 1, BTT24170 1, BTT24170 2). They were all upregulated. An apidaecin precursor 334 (BTT20828 1) was upregulated. Hemolectin had three downregulated transcripts (BTT14194 1, 335 BTT17013 1, BTT26614 1) and one upregulated (BTT15326 1). Argonaute-2, a RNA-silencing 336 endonuclease, is involved in antiviral defense in insects (downregulated BTT02374 1) (van Rij et 337 al. 2006). 338 339 Eater encodes for a transmembrane receptor involved in phagocytosis in Drosophila (Kocks et al. 340 2005). A transcript (BTT11132 1) relating to *Eater* was upregulated. The melanisation process 341 component Dopa decarboxylase (BTT19093 1) was upregulated. Another enzyme involved in 342 melanisation, laccase was found to be downregulated (BTT20241 1, BTT33633 1) (Arakane et al. 343 2005). 344 345 Cyp4p3 transcript BTT40653 1 was upregulated. Two previously unseen Cyp4p3 transcripts 346 (BTT05254 1, BTT20622 2) were upregulated and one (BTT36257 1) downregulated. TPX4 347 (BTT13285 1) that codes for a Thioredoxin-dependent peroxidase was downregulated. 348 349 Miscellaneous: 350 A small number of transcripts were related to chitin metabolism. SCRASP1 has a chitin-binding 351 domain that has been hypothesized to sense chitin in response to injury and to transduce signals via 352 the serine protease domain (Blumberg et al. 2013). We found an upregulated transcript related to 353 SCRASP 1 (BTT41923 1). A peritrophin precursor was also upregulated (BTT10727 1). As was a 354 chitinase 3 transcript (BTT23246 1). 355 356 Retinoid and fatty-acid-binding protein (RfaBp) (BTT07678 1) was downregulated. RfaBp was

found to be upregulated upon injection of LPS in *Drosophila* during a proteomic study (Vierstraete *et al.* 2004). *Notch* (upregulated BTT09545_1) is involved in the specification of crystal cells in *Drosophila melanogaster* (Mukherjee *et al.* 2011). Several orthologs of putative *Drosophila* immune loci were found to be differentially expressed (CG5527: BTT08512_1, CG12505: BTT00934_1, CG13323: BTT38025_1, BTT38087_1, CG17560: BTT02877_1 downregulated, BTT05845_1 upregulated, CG18348: BTT20843_1)

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Discussion We present a comprehensive transcriptomic analysis of gene expression in this important model host-parasite system. We have identified a large number of bumblebee genes whose expression are changed upon infection with Crithidia. We have also found a large number of genes whose expression depends on the interaction between host and parasite genotypes, that is show specificity. We confirmed the importance of antimicrobial peptides in the specific defense against Crithidia (Riddell et al. 2009, 2011; Deshwal & Mallon 2014). It is also clear that several other effectors including ROS and phagocytosis may be important. Several immune pathways seem to be important in the anti-Crithidia response. These include the Toll, Imd and JAK/STAT pathways. Toll especially seems to be important in a specific immune response. There are a larger proportion of receptor transcripts found in the specificity analysis (3.2% 19/591) compared to the infection analysis (1.2% 6/489). This is not surprising, as we would expect a specific immune response to a given strain to be based mainly on how it is recognised. Although several receptors, including GNBPs and lectins, are differentially expressed, the most exciting discovery is the large number of transcripts related to *Dscam*. The Down syndrome cell adhesion molecule (Dscam), a pattern recognition receptor has come to the forefront of research into insect immune specificity as it has been found to have thousands of different splice forms and is associated with insect immunity (Smith et al. 2011). In the fruit fly Drosophila, silencing of Dscam retards the insect's capacity to engulf bacteria by phagocytosis (Watson et al. 2005). In Anopheles, the Dscam splice forms produced in response to parasite exposure differs between bacteria and Plasmodium and between Plasmodium berghei and Plasmodium falciparum (Dong et al. 2006). This has been tempered by the finding that Dscam diversity does not increase with exposure to increasing heterogeneity of *Plasmodium* falciparum genotypes (Smith et al. 2011). Recently it has been shown that *Dscam* specificity is mediated by specific splice-factors transcription downstream

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of activation of the Toll and Imd pathways (Dong et al. 2012). Our results suggest that Dscam may be important in differentiating strains of the trypanosome *Crithidia bombi*. We found a number of genes associated with chitin metabolism. The peritrophic matrix may be fundamental in the bee's defense against Crithidia. The peritrophic matrix acts as an immunological barrier against trypanosomes. Tsetse flies with an underdeveloped PM have lower levels of refractoriness to trypanosome infections (Weiss et al. 2013). This is due to a premature immune response; the trypanosomes get through the PM quicker and stimulate the immune response at an earlier stage compared to refractory flies. A similar study has recently been published (Barribeau *et al.* 2014). Although they find geneotype x genotype interactions in gene expression, it is in a much smaller number of genes. None of these genes offer a potential model for generating specificity. We hypothesise that our much larger catalogue of genes, including Dscam, is due to our experimental design. We used a 2x2 analysis of preselected reciporcal host-parasite interactions. This increased our likelihood of detecting differential expression using RNA-seq compared to a less directed sample collection. Given that we have found that the bees own physiology, especially its immune response is vital in both the defense against Crithidia and in explaining the host-parasite specificity, how do we incorporate recent findings that the bees gut microbiota are important in exactly these phenomena (Koch & Schmid-Hempel 2011, 2012). Gut microbiota impact the condition of the PM and gut epithelium generally (Buchon et al. 2009a; Weiss et al. 2013). It has recently been suggested that the components of the peritrophic matrix may be under the control of various immune pathways, Imd (Buchon et al. 2009b) and STAT (Narasimhan et al. 2014) explicitly. Gut microbiota stimulate these pathways keeping the PM intact. The intact peritrophic matrix then acts as a physical barrier to colonization by parasites. Future work will focus on understanding the interactions of this

- 416 triumvirate of host genotype, parasite genotype and gut microbiota and their effect on disease
- 417 outcome.

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427

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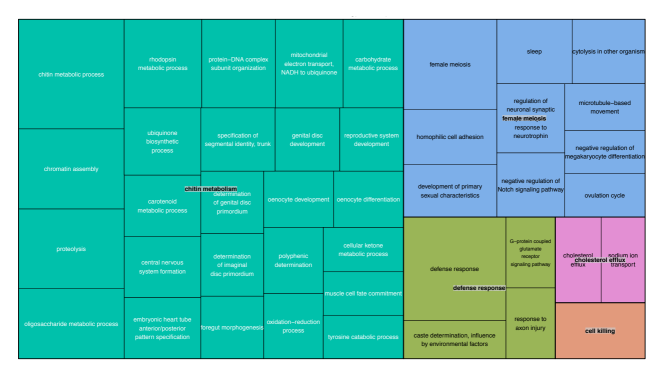


Figure 1 A summary of the enriched GO terms (based on Blast2Go annotation) found for differentially expressed genes at 24 hours post-infection compared to uninfected samples. This figure was produced using Revigo (Supek et al. 2011)

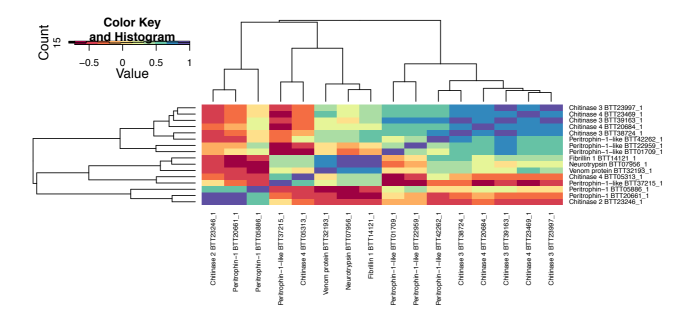


Figure 2. A heatmap showing the correlations of the expression patterns of the transcripts labelled as chitin metabolism genes that where differentially expressed twenty four hours after infection.

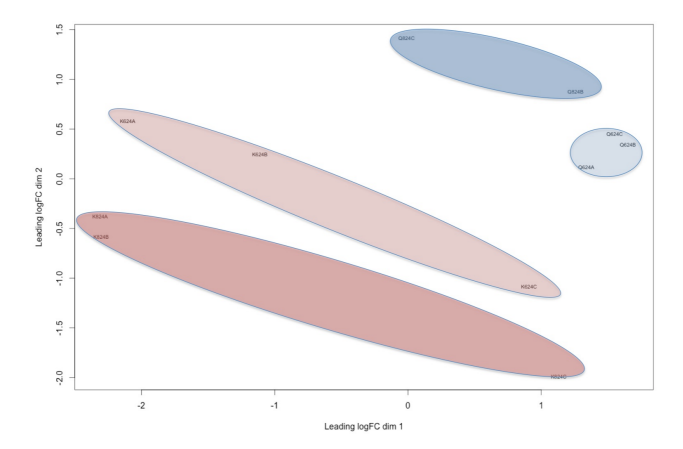


Figure 3. A multidimensional scaling (MDS) plot of the 11 samples used in the specificity analysis based on the expression of 591 differentially expressed transcripts. There are two colonies (K (red) and Q (blue)) and two *Crithidia* strains (6 (light) and 8 (dark)). Dimension 1 is the direction that best separates the samples. Dimension 2 is the next best direction, uncorrelated with the first, that separates the samples. The samples are clearly grouped into their colony-strain interactio

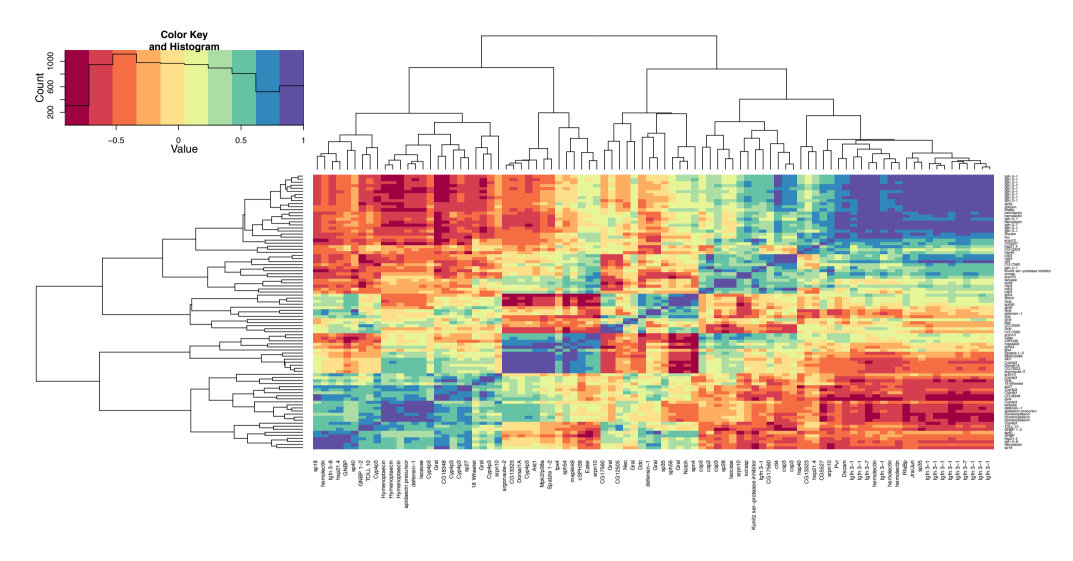


Figure 4. A heatmap showing the correlations of the expression patterns of the 90 transcripts labelled as immune genes in the analysis identifying genes differentially expressed depending on host genotype – parasite genotype interactions.

Serine Proteases	Upregulated	Downregulated
CLIPA6		BTT20125_1
CLIPA7		BTT07313_1, BTT31897_1
CLIPD5		BTT10579_1, BTT10912_1, BTT18247_1 BTT25711 1, BTT06803 1
SP24	BTT03436_1	
SP27		BTT08108_1, BTT38696_1
SP35	BTT05300_1	
Serine protease homologues	Upregulated	Downregulated
SPH54	BTT01977_1	BTT06125_1
Serine protease inhibitors	Upregulated	Downregulated
NEC-like		BTT31997_1
spn4		BTT04130_1, BTT40693_1, BTT41025_1, BTT41461_1
SRPN10		BTT04508_1, BTT20259_1

Table 1. List of transcriptions associated with serine proteases and serine protease inhibitors found to be differentially expressed twenty four hours post infection with Crithidia bombi.

Serine Proteases	Upregulated	Downregulated
cSp3	BTT35293_1	BTT10579_1, BTT10912_1, BTT18247_1, BTT25711_1
Sp18	BTT20808_1	
Sp27	BTT40251_1	
Sp28		BTT20637_1
Sp35	BTT05300_1	BTT10155_1
Sp40	BTT15256_1	
Tequilla/GRAL/Sp23	BTT01709_1, BTT05886_1, BTT09081_1, BTT20661_1, BTT20725_1, BTT24359_1, BTT25071_1	
Serine protease homologues	Upregulated	Downregulated
cSPH39		BTT21868_1
Sph54		BTT27769_1
Sph56	BTT17814_1	
Serine protease inhibitors	Upregulated	Downregulated
Kunitz ser-protease inhibitor		BTT14993_1
necrotic (nec)	BTT35742_1	
Spn 4		BTT04130_1
SRPN10		BTT02607_1, BTT4508_1, BTT20259_1, BTT40693_1

Table 2. List of transcriptions associated with serine proteases and serine protease inhibitors found to be differentially expressed in the specificity analysis.