Olfactory Proteins and Their Expression Profiles in the Eucalyptus Pest Endoclita signifier Larvae

Xiaoyu Zhang1,2, Zhende Yang1, Xiuhao Yang3, Hongxuan Ma4, Xiumei Liu4 and Ping Hu1,2*

1 Forestry College, Guangxi University, Nanning, China, 2 Forestry College, Central South University of Forestry and Technology, Changsha, China, 3 Guangxi Academy of Forestry, Nanning, China, 4 Guangxi Gaofeng National Forest Farm, Nanning, China

Endoclita signifier Walker (Lepidoptera: Hepialidae), a polyphagous insect, has become a new wood-boring pest in Eucalyptus plantations in southern China since 2007, which represents a typical example of native insect adaptation to an exotic host. After the third instar, larvae move from soil to standing trees and damage the plants with a wormhole. Although females disperse to lay eggs, larvae can accurately find eucalyptus in a mingled forest of eight species, which leads us to hypothesize that the larval olfactory system contributes to its host selection. Herein, we investigated the transcriptomes of the head and tegument of E. signifier larvae and explored the expression profiles of olfactory proteins. We identified 15 odorant-binding proteins (OBPs), including seven general OBPs (GOPBs), six chemosensory proteins (CSPs), two odorant receptors (ORs), one gustatory receptor (GR), 14 ionotropic receptors (IRs), and one sensory neuron membrane protein (SNMP). Expression profiles indicated that all olfactory proteins, except for EsigCSP1, were expressed in the head, and most were also detected in non-olfactory tissues, especially thorax tegument. Furthermore, EsigOBP2, EsigOBP8, EsigGOBP1, EsigGOBP2, EsigGOBP5, EsigCSP3, EsigCSP5, and EsigOR1 were expressed most strongly in the head; moreover, EsigCSP3 expressed abundantly in the head. EsigGR1 exhibited the highest expression among all tissues. Besides phylogenetic analysis shows that EsigGOBP7 probably is the pheromone-binding protein (PBP) of E. signifier. This study provides the molecular basis for future study of chemosensation in E. signifier larvae. EsigCSP3 and EsigGR1, which have unique expression patterns, might be factors that govern the host choice of larvae and worth further exploration.

Keywords: expression profiles, chemosensory proteins, transcriptome, odorant binding protein, Endoclita signifier

INTRODUCTION

The ghost moth Endoclita signifier Walker (Lepidoptera, Hepialidae) is a native polyphagous insect pest that is widely distributed in Japan, Korea, India, Thailand, Myanmar, and central, south, and southwest China (Yang et al., 2021). After Eucalyptus was planted in the south of China, E. signifier was discovered to have infested Eucalyptus in Guangxi in 2007.
This is an example of a native insect adapting to an exotic host, and it has resulted in great economic losses and major ecological impacts (Yang et al., 2016). In 2020, an infestation of *E. signifier* was found in 17.1% of the counties in Guangxi, where host plants included 51 species in 40 genera and 30 families (Yang et al., 2021). In Guangxi, *E. signifier* usually produces one generation a year and occasionally two. The adults eclose from mid-March through April and then mate and oviposit. Larvae hatch within 1 month and live in the soil. After the third instar stage, from July through August, the larvae move from soil to standing trees, where they feed on bark, bore into the stem, and weave wood pieces and silk over the hole entrance, constructing a home in which they reside until the following January before pupating in February (Figure 1). Some larvae spend two years in the tree, until January of the third year (Yang, 2013). Although female oviposition proceeds in a dispersed manner, larvae were shown to damage eight *Eucalyptus* species in mixed forests (Yang, 2017). Besides, the sensillum of *E. signifier* larva shows that only 16 sensilla in antennae, but a large number of sensilla in thoracic and abdominal tegument (Hu et al., 2021). Therefore, we hypothesized that the thoracic and abdominal tegument may have an olfactory function and help larvae to find their host.

In most insects, especially stem borers, females select the host. Olfaction plays a minor role in larvae. The small number of neurons in the *Drosophila* olfactory system makes it a very convenient system for olfactory studies (Bose et al., 2015), enabling the exploration of sensory coding at all stages of nervous system development (Bose et al., 2015). In *Drosophila* larvae, the epidermal growth factor receptor (Rahn et al., 2013) and serotonin signaling (Annina et al., 2017) are necessary for learning and memory. However, sub-circuits allow *Drosophila* larvae to integrate present sensory input into the context of past experience and elicit an appropriate behavioral response (Rahn et al., 2013). The responsiveness of larval sensilla to female-emitted sex pheromones is based on the same molecular machinery as that functioning in the antennae of adult males (Zielonka et al., 2016). Olfactory proteins that bind, transport, and degrade odor molecules play important roles in chemo-sensing in larvae and include odorant-binding proteins (OBPs), chemosensory proteins (CSPs), sensory neuron membrane proteins (SNMPs), odorant receptors (ORs), gustatory receptors (GRs), ionotropic receptors (IRs), and odor-degrading enzymes (ODEs) (Zhou, 2010). Behavior, physiology, and molecular result indicated olfactory proteins of larvae play a role in pheromone or volatile recognition. For example, four CSPs in *Chilo auricilius* larvae (Yi et al., 2019) were significantly regulated by α-pinene treatment. Pre-exposure larvae to single volatile organic compounds can upregulate ORs in *Spodoptera exigua* larvae strongly and non-specifically (Llopis-Gimenez et al., 2020). *S. exigua* larval OBP can bind to a major female sex pheromone component (Jin et al., 2015), whereas OBP8 of *Melipona scutellaris* is expressed and functioned in the larval mandible (Carvalho et al., 2017).

This study investigated the transcriptomes in the head and tegument of *E. signifier* larvae to determine the expression profiles of olfactory proteins in *E. signifier* larvae and evaluated the phylogenetic relationships between *E. signifier* OBPs and CSPs with those expressed in the larvae and adults of other species. This work illuminates the olfactory system in *E. signifier* larvae and provides a theoretical basis for further studies to explore ecologically relevant larval behaviors.

**MATERIALS AND METHODS**

**Ethics Statement**

The ghost moth *E. signifier* is a forestry pest in China, which were collected with the direct permission of the Guangxi forestry bureau. It is not included in the “list of endangered and protected animals in China.” All operations were performed according to ethical guidelines in order to minimize pain and discomfort to the insects.

**Insect and Tissue Collection**

*E. signifier* larvae were collected from damaging *Eucalyptus* plantation by cutting the tree during December 2019 to January 2020 and September to November 2020 in the Gaofeng forest station, Guangxi, China. Six 12th larvae and 18 5th larvae were taken indoors and stored at −80°C. Larval thoracic and abdominal tegument were obtained by using surgical scissors to cut open the larvae abdomen through the midline and using

---

**FIGURE 1** (a,b,d) Damage caused by *E. signifier*. (c) Larvae and (e) adults of *E. signifier*. 

---

Zhang et al.
tweezers to remove from the rest of the body (intestinal canal and fat body), and the teguments were cleaned in RNA-free ddH₂O.

**cDNA Library Construction and Illumina Sequencing**

Total RNA was extracted from six 12th larval head, thoracic, and abdominal tegument using TRIzol reagent (Ambion) and the RNeasy Plus Mini Kit (No. 74134; Qiagen, Hilden, Germany), following the instructions of the manufacturer, respectively. RNA quantity was detected using the NanoDrop 8000 (Thermo, Waltham, MA, USA). There is one replication, each replication with six larvae. RNA of 12th larval heads and thoracic and abdominal tegument were used to construct the cDNA libraries. cDNA library construction and Illumina sequencing of samples were performed at MajorBioCorporation (Shanghai, China). mRNA samples were purified and fragmented using the TruSeq RNA Sample Preparation Kit v2-Set A (No. RS-122-2001; Illumina, San Diego, CA, USA). Random hexamer primers were used to synthesize the first-strand cDNA, followed by synthesis of the second-strand cDNA using a buffer, dNTPs, RNase H, and DNA polymerase I at 16°C for 1 h. After end repair, A-tailing, and the ligation of adaptors, the products were amplified by PCR and quantified precisely using the Qubit DNA Br Assay Kit (Q10211; Invitrogen, Carlsbad, CA, USA). They were then purified using the MinElute Gel Extraction Kit (Qiagen, Cat No. 28604) to obtain a cDNA library. The cDNA library was sequenced on the HiSeq2500 platform.

**Assembly and Functional Annotation**

All raw reads were processed to remove low-quality and adaptor sequences using Trimmomatic (http://www.usadellab.org/cms/index.php?page=trimmomatic). Clean reads assembly was carried out with the short-read assembly program Trinity (Version: r2020-01-13), with the default parameters, after combining the heads and thoracic and abdominal tegument clean reads. The largest alternative splicing variants in the Trinity results were called “unigenes.” The annotation of unigenes was performed by NCBI BLASTX searches against the Nr protein database, with an E-value threshold of 1e-5. The blast results were performed by NCBI BLASTx searches against the Nr protein database, with an E-value threshold of 1e-5. The blast results were then imported into the Blast2GO pipeline for GO annotation. The candidate OBPs were searched for the presence of N-terminal signal peptides using SignalP4.0 (http://www.cbs.dtu.dk/services/SignalP/). Amino acid sequence alignment was performed using the muscle method implemented in the Mega v6.0 software package (Tamura et al., 2011). The phylogenetic tree was constructed using the neighbor-joining (NJ) method (Saitou and Nei, 1987) with the P-distances model and a pairwise deletion of gaps performed in the Mega v6.0 software package. The reliability of the tree structure and node support was evaluated by bootstrap analysis with 1,000 replicates. The phylogenetic trees were colored and arranged in FigTree (Version 1.4.2). The phylogenetic analyses of OBPs were based on Dastarcus helophoroides (Li et al., 2020), Chrysomya megacephala (Wang et al., 2015), Plutella xylostella (Zhu et al., 2016), S. exigua (Liu et al., 2015; Llopis-Gimenez et al., 2020), Helicoverpa armigera (Chang et al., 2017), and E. signifier. The CSPs tree was based on D. helophoroides (Li et al., 2020), C. megacephala (Wang et al., 2015), S. exigua (Llopis-Gimenez et al., 2020), H. armigera (Chang et al., 2017), and E. signifier. The gene name and the Genbank number of P. xylostella and H. armigera are listed in Supplementary Material 2; other genes sequences are available in the reference article.

**Sequence and Phylogenetic Analysis**

The candidate OBPs were searched for the presence of N-terminal signal peptides using SignalP4.0 (http://www.cbs.dtu.dk/services/SignalP/). Amino acid sequence alignment was performed using the muscle method implemented in the Mega v6.0 software package (Tamura et al., 2011). The phylogenetic tree was constructed using the neighbor-joining (NJ) method (Saitou and Nei, 1987) with the P-distances model and a pairwise deletion of gaps performed in the Mega v6.0 software package. The reliability of the tree structure and node support was evaluated by bootstrap analysis with 1,000 replicates. The phylogenetic trees were colored and arranged in FigTree (Version 1.4.2). The phylogenetic analyses of OBPs were based on Dastarcus helophoroides (Li et al., 2020), Chrysomya megacephala (Wang et al., 2015), Plutella xylostella (Zhu et al., 2016), S. exigua (Liu et al., 2015; Llopis-Gimenez et al., 2020), Helicoverpa armigera (Chang et al., 2017), and E. signifier. The CSPs tree was based on D. helophoroides (Li et al., 2020), C. megacephala (Wang et al., 2015), S. exigua (Llopis-Gimenez et al., 2020), H. armigera (Chang et al., 2017), and E. signifier. The gene name and the Genbank number of P. xylostella and H. armigera are listed in Supplementary Material 2; other genes sequences are available in the reference article.

**Expression Analysis by Fluorescence Quantitative Real-Time PCR**

Fluorescence quantitative real-time PCR was performed to verify the expression of candidate chemosensory genes. The total RNA of the 18 fifth instar larval head, thoracic, and abdominal tegument was extracted following the methods described above. NanoDrop 2008 and agarose gel electrophoresis examined the density and quality of the RNA. cDNA was synthesized from the total RNA, using the PrimeScript RT Reagent Kit with gDNA Eraser to remove gDNA (No. RR047A; TaKaRa, Shiga, Japan). Gene-specific primers were designed using Primer 3 (http://bioinfo.ut.ee/primer3-0.4.0/) (Supplementary Material 3). Eighteen SRNAs were identified, evaluated, and selected as a reference gene for qPCR (Chen and Hu, in press). A PCR analysis was conducted using the Roche LIGHT CYCLE 480II (USA). SYBRPremixExTaq™ II (No. RR820A; TaKaRa) was used for the PCR reaction under three-step amplification. Each PCR reaction was conducted in a 25-µl reaction mixture containing 12.5 µl of SYBR Premix Ex Taq II, 1 µl of each primer (10 mM), 2 µl of sample cDNA, and 8.5 µl of dH₂O. The RT-qPCR cycling parameters were as follows: 95°C for 30 s, followed by 40 cycles of 95°C for 5 s, 60°C for 30 s, and 65°C to 95°C in increments of 0.5°C for 5 s to generate the melting curves. To examine reproducibility, each qPCR reaction for each tissue was performed in three biological replicates (each replicate with six larvae) and three technical replicates. Negative controls without either template were included in each experiment. RocheLIGHT CYCLE 480II was used to normalize the expression based on ΔΔCq values, with GOBP3 in the head as control samples, and the 2−ΔΔCq method was used (Livak and Schmittgen, 2001). Before comparative analyses, the normal distribution and equal variance tests were examined, and all
taken logarithm data followed a normal distribution and with equal variances. The comparative analyses for every gene among six tissue types were assessed by a one-way nested analysis of variance (ANOVA), followed by Tukey's honestly significance difference (HSD) tests implemented in SPSS Statistics 18.0. Values are presented as means ± SE.

RESULTS

Transcriptome Sequencing and Sequence Assembly

We generated 53 million raw reads from a cDNA library derived from the tegument of *E. signifer* larvae, with q20 and q30 scores for 97.84% and 93.74% of the reads severally. The larval head transcriptome yielded 51 million raw reads, with q20 and q30 scores for 97.83% and 93.65% of the reads, respectively. After trimming the adapters, removing low-quality raw sequences, using Trimmomatic, blending the head and tegument sequences, splicing, and assembly (using Trinity), we obtained 44,104 transcripts, with an N50 of 1,707 bp, an average length of 986 bp, and a maximum length of 56,111 bp (Figure 2A). The *E. signifer* raw reads have been deposited in the NCBI Sequence Read Archive database under GenBank accession number PRJNA713545.

Homology Analysis and Gene Ontology Annotation

For 41.17% of the non-redundant protein database, using BLASTx with an E-value cutoff of 1e-5. The most frequent sequence matches were with *Eumeta japonica* (6.06%), followed by *Ostrinia furnacalis* (5.77%) and *Hyposmocoma kahamaanao* (5.60%) (Figure 2B). We used gene ontology (GO) annotations to classify the 10,177 transcripts into functional groups with BLAST2GO, with p-values calculated from the hypergeometric distribution test and an E-value threshold of < 1 × 10^{-5}. In the *E. signifer* transcriptome, molecular functions accounted for 36.75% of the GO annotations, followed by cellular components (33.96%) and biological processes (29.29%). In the molecular function category, the terms binding, catalytic activity, and transporter activity had the highest representation. In the biological process category, the terms cellular process, metabolic process, and biological regulation were most frequent. Membrane part, cell part, and organelle were the most common cellular component terms (Figure 3).

Olfactory Proteins

We identified 15 transcripts-encoding putative OBPs in *E. signifer*, of which six were general OBPs (GOBPs). According to the FPKM value of the unigenes, EsigGOBP3 and EsigOBP7 were expressed less strongly in the head, and ten OBPs were not expressed in the tegument (Table 1). We identified six transcript-encoding putative CSPs, of which four were expressed more strongly in the head, whereas CSP1 and CSP2 were expressed more strongly in the tegument (Table 1). One transcript encoded a putative SNMP and was strongly expressed in the tegument (Table 1). Two identified ORs were strongly expressed in the head (Supplementary Material 4).

Olfactory Protein Expression Profiles

We characterized the expression profiles of all the OBPs, CSPs, ORs, GRs, and SNMPs in the head, thoracic, and abdominal tegument of fifth *E. signifer* larvae. Of the OBPs, EsigOBP5 was the most strongly expressed OBPs in all larval tissues (10- to 1,000-fold higher values). EsigOBP2, EsigOBP8, EsigGOBP1, EsigGOBP2, and EsigGOBP5 were most strongly expressed in the head, and EsigOBP8 and EsigGOBP5 were expressed at significantly lower levels in the thorax than in the head. EsigGOBP2 exhibited significant head-biased expression. In non-olfactory tissues, EsigOBP3, EsigOBP5, EsigOBP6, EsigGOBP3, and EsigGOBP7 were most highly expressed in the thoracic tegument, with a significant thorax-biased expression of EsigGOBP3. EsigGOBP7 was expressed at significantly higher levels in the thorax and abdomen than in the head. EsigOBP1, EsigOBP7, EsigGOBP4, and EsigGOBP6 were most strongly expressed in the abdominal tegument, especially EsigOBP7. Significantly, more EsigGOBP6 was expressed in the head and abdomen than in the thorax (Figure 4). Of the CSPs, EsigCSP3 and EsigCSP6 were the most strongly expressed CSPs, with levels 100- to 1,000-fold of other CSPs, and EsigCSP3 was only expressed in the head, but EsigCSP6 was strongly expressed in all larval tissues. EsigCSP5 and EsigCSP3 were most strongly expressed in the head, especially EsigCSP5. The expression of EsigCSP2, EsigCSP4, and EsigCSP6 was highest in the abdomen, especially for EsigCSP2 and EsigCSP4. EsigCSP4 expression differed significantly from the others in the three tissues, and EsigCSP1 was highly expressed in the thorax (Figure 4). EsigGR1 was expressed most strongly in the abdomen (Figure 4). EsigOR1 was expressed only in the head, whereas, significantly, more EsigOR2 was expressed in the abdomen than in other tissues. We also detected significantly higher EsigSNMP1 expression in the abdomen compared with the other tissues (Figure 4).

Phylogenetic Analysis of OBPs and CSPs

In the phylogenetic tree of OBPs (Figure 5), the no-expression clade (blue) included EsigGOBP3, EsigOBP6, EsigGOBP2, EsigGOBP7, EsigOBP1, EsigOBP3, EsigOBP4, and EsigOBP8. The OBPs not expressed in larvae included SexiOBP3, SexiOBP7, SexiOBP9, SexiOBP17, SexiOBP36, SexiOBP39, SexiOBP42, - SexiOBP46, SexiOBP47, SexiPBP1, SexiPBP2, SexiPBP4, DhelOBP1, and DhelOBP9. The expression clade (green) included EsigGOBP6, EsigGOBP5, EsigOBP2, EsigGOBP4, EsigOBP5, EsigOBP7, with SexiOBP8, SexiOBP21, SexiOBP24, SexiOBP25, SexiOBP26, SexiOBP27, SexiOBP28, SexiOBP29, SexiOBP31, SexiOBP32, SexiOBP33, and SexiOBP6 expressed only in larvae. The PBP clade (red circle) contained EsigGOBP7; four PBPs of *S. exigua*; PBP1, G0BP1, and G0BP2 of *P. xyllostella*; and HarmGOBP2 (Figure 5). In the phylogenetic tree with CSPs, the no-larval-expression clade (blue) included SexiCSP1, SexiCSP2, and SexiCSP4, SexiCSP23, and SexiCSP24 (Figure 6).
DISCUSSION

Larval survival contributes to its host selection. From the head and tegument transcriptomes, we identified 39 olfactory proteins, including 15 OBPs, 6 CSPs, 2 ORs, 1 e GR, 1 SNMP, and 14 IRs. This is the first report of the separate larval head and tegument transcriptomes in Hepialidae. The number of olfactory proteins identified in E. signifer is less than that reported for most adult antennae transcriptomes, such as Conogethe spinicolalis (Jing et al., 2020). Additionally, larvae tend to have a shorter squirm range and a less complicated survival environment than adults and may consequently express a smaller set of olfactory genes than adults, as observed for Spodoptera littoralis (Poivet et al., 2013). The number of olfactory proteins in E. signifer is considerably fewer than the 26 OBPs and 21 CSPs identified in the transcriptomes of H. armigera larval antennae and mouthparts (Chang et al., 2017); the 20 OBPs, 11 CSPs, 9 ORs, 11 IRs, 7 GRs, and 4 SNMPs in the transcriptomes of newly hatched Dastarcushelophoroides larvae (Li et al., 2020); and the 58 ORs, 20 GRs, and 21 IRs in the transcriptomes of the antennae of males and females and the head tissue of neonates of Cydia pomonella (Walker et al., 2016). This reflects the scarcity of olfactory proteins of the original Lepidoptera group in the NCBI database, such as Hepialidae, resulting in less annotation of olfactory proteins in E. signifer.

According to the OBP expression profiles in larvae, distinct no-larval-expression or larval-specific-expression clades were apparent in the neighbor-joining tree based on OBPs, with

FIGURE 2 | (A) Length distribution of E. signifer unigenes and (B) BLASTx comparison of unigenes in the E. signifer transcriptome with those of other species.
EsigGOBP3, EsigOBP6, EsigGOBP2, EsigGOBP7, EsigOBP1, EsigOBP3, EsigOBP4, and EsigOBP8 placed in the no-larval-expression clade. EsigCSP1 and EsigCSP2 belonged to the no-larval-expression clade in the CSP phylogenetic tree, and EsigCSP1 was not expressed in the larval head. However, EsigGOBP4, EsigGOBP5, EsigGOBP6, EsigOBP2, EsigOBP5, and EsigOBP7 were placed in the larva-specific-expression clade, and EsigOBP5 and EsigOBP7 were highly expressed in larvae compared with the other genes. The small number of proteins known to be specifically expressed in larvae may cause false positives in the larva-specific clade. EsigGOBP7 was placed in the PBP clade, suggesting that it is an *E. signifer* PBP and that PBPs expressed in larvae function as sex pheromones binding (Zielonka et al., 2016).

The expression profile of olfactory proteins in *E. signifer* larvae showed that all were expressed in at least one tissue, verifying the olfactory proteins identified in the head and tegument transcriptomes. EsigCSP6, EsigOBP5, EsigGOBP1, EsigOBP6, EsigOBP7, and EsigGR1 were highly expressed in all tissues of the fifth instar larvae, and the high expression of olfactory proteins indicates that *E. signifer* larvae require many olfactory proteins to support olfactory recognition, especially the transfer from soil to standing trees. Additionally, we observed the expression in all tissues, except the head, thorax, and abdomen.
of borers, for some proteins with olfactory functions, while some olfactory proteins with no olfactory functions, such as OomCSP12 of *Ophraella commun*ua, are expressed in female ovaries, and silencing of OomCSP12 results in significantly reduced ovipositing by females (Ma et al., 2019). Eating is the main behavior in larvae, which may explain why EsigGR1 exhibited the highest expression among all tissues, and EsigGR1 may correlate with the feeding habits of *E. signifier* larvae and reflect gustatory preferences.

The head is the center of sensation. EsigCSP3 was strongly expressed only in the head, and CSPs are known to contribute to mediating responses to plant volatility in *Eucalyptus*-derived compounds in *E. signifier* antennae. Importantly, eight genes were expressed most strongly in the head (EsigOBP2, EsigOBP8, EsigGOBP1, EsigGOBP2, EsigGOBP5, EsigCSP3, EsigCSP5, and EsigOR1), with the expression of EsigGOBP2, EsigCSP5, and EsigOR1 biased to the larval head, consistent with the observation of 50 *S. exigua* ORs expressed in larval heads (Llopis-Gimenez et al., 2020) and larvae (Liu et al., 2015). For *S. exigua* OBPs, expression levels are higher in the larval head than in the larval body (Liu et al., 2015). In the larval head, many *S. littoralis* OBPs and CSPs exhibit organ-specific transcription in caterpillar antennae and maxillary palps, suggesting the complementary involvement of these two organs in larval chemosensory detection (Poivet et al., 2013). Moreover, *H. armigera* expressed more OBPs and CSPs in the larval antennae than in the mouthparts (Chang et al., 2017). Therefore, it is necessary to examine the expression of the eight most strongly expressed olfactory genes in the antennae and mouthparts of *E. signifier* to determine their roles in smelling and tasting the odors of the host plant (Jin et al., 2015; Di et al., 2017; Waris et al., 2020).

| Name                | Nr description                                | Species                                      | Acc. NO.       | Tegument FPKM | Head FPKM | Tegument vs. Head |
|---------------------|-----------------------------------------------|----------------------------------------------|----------------|---------------|-----------|------------------|
| EsigOBP1            | Odorant binding protein LOC100307012 precursor | *Bombyx mori*                                | NP_001159621.1 | 0             | 1.39      | Down             |
| EsigOBP2            | Odorant binding protein 7                     | *Grapholita molesta*                        | AVZ44706.1     | 0             | 4.32      | Down             |
| EsigOBP3            | Odorant binding protein 8                      | *Bombyx mori*                                | NP_001159621.1 | 0             | 8         | Down             |
| EsigOBP4            | Odorant binding protein 10                     | *Bombyx mori*                                | NP_001159621.1 | 0             | 4.04      | Down             |
| EsigOBP5            | Odorant binding protein                        | *Eogystia hippophaeolus*                    | AOG12872.1     | 0             | 2.97      | Down             |
| EsigGOBP1           | General odorant-binding protein 70-like        | *Amylos translata*                           | XP_013201142.1 | 0             | 2.97      | Down             |
| EsigGOBP2           | General odorant-binding protein 56d-like       | *Hyposmocoma kahamanoa*                     | XP_026319068.1 | 0.45          | 4.87      | Down             |
| EsigGOBP3           | General odorant-binding protein 82a-like       | *Plutella xylostella*                        | XP_011154700.1 | 1.48          | 0.39      | Up               |
| EsigGOBP4           | General odorant-binding protein 19d-like       | *Papilio xuthus*                             | XP_013173035.1 | 0             | 4.47      | Down             |
| EsigGOBP5           | General odorant-binding protein 19d            | *Eumeta japonica*                            | GBP31818.1     | 0             | 4.98      | Down             |
| EsigGOBP6           | General odorant-binding protein 28a-like       | *Hyposmocoma kahamanoa*                     | XP_026330999.1 | 0             | 3.07      | Down             |
| EsigOBP6            | Odorant-binding protein 16                     | *Ectropis obliqua*                           | AL503864.1     | 18.31         | 107.65    | Down             |
| EsigOBP7            | Putative odorant-binding protein A10 isoform X2| *Zeugodacus cucurbitae*                      | XP_011177223.1 | 101.91        | 98.21     | Down             |
| EsigOBP8            | Odorant binding protein                        | *Conogethes punctiferalis*                  | APG32543.1     | 0.07          | 30.95     | Down             |
| EsigGOBP7           | General odorant-binding protein 1              | *Athetis dissimilis*                         | ALJ93806.1     | 0             | 13.48     | Down             |
| EsigCSP1            | Chemosensory protein 10                        | *Carposina sasakii*                         | AYD42214.1     | 1.82          | 0.52      | Up               |
| EsigCSP2            | Chemosensory protein 24                        | *Cnaphalocrocis medinalis*                  | ALT31606.1     | 181.82        | 2.78      | Up               |
| EsigCSP3            | Chemosensory protein 5                         | *Agrotis ipalor*                             | AGR39575.1     | 9.65          | 10.58     | Down             |
| EsigCSP4            | Chemosensory protein CSP14                     | *Lobesia botrana*                            | AXF48711.1     | 3.16          | 9.35      | Down             |
| EsigCSP5            | Chemosensory protein 5                         | *Emopoasca onuki*                            | AWG68022.1     | 0             | 71.07     | Down             |
| EsigCSP6            | Chemosensory protein                            | *Cnaphalocrocis medinalis*                  | AIX97837.1     | 10.33         | 218.45    | Down             |
| EsigOR1             | Odorant receptor 28, partial                   | *Locusta migratoria*                        | ALD51442.1     | 4.17          | 0         | Up               |
| EsigOR2             | Odorant receptor OR4                           | *Rhyacophila nubila*                        | AYN64394.1     | 0.98          | 0.94      | Up               |
| EsigGR1             | Gustatory receptor                             | *Eogystia hippophaeolus*                    | AOG12970.1     | 17.74         | 76.58     | Down             |
| EsigSNMP1           | Sensory neuron membrane protein 2 isoform X1   | *Neodiprion lecontei*                       | XP_015517411.1 | 1.87          | 0.16      | Up               |

**TABLE 1** Best blastx hits for odorant-binding proteins (OBPs), chemosensory proteins (CSPs), odorant receptors (ORs), gustatory receptors (GRs), and sensory neuron membrane proteins (SNMPs) of *Endoclita signifier*. 

---

*Zhang et al.* Olfactory Proteins of *Eucalyptus* signifer. July 2021 | Volume 12 | Article 682537
Furthermore, the expression profiles of olfactory proteins in *S. exigua* larvae indicated that they are also expressed in non-olfactory tissues, such as the larval body (Liu et al., 2015; Llopis-Gimenez et al., 2020). In *E. signifer* larvae, most olfactory proteins were highly expressed in thoracic and abdominal tegument, consistent with the large number of sensilla on the thorax and abdomen (Hu et al., 2021). This establishes the molecular basis for the head and the other main sensor tissues. Borers need to use non-olfactory tissues, such as the thorax and abdomen, to sense the wood hole environment and adapt to wood hole survival. It is necessary to explore the functions of olfactory proteins that are highly expressed in non-olfactory tissues, such as EsigGOBP3, EsigOBP7, EsigCSP2, EsigOR2, and EsigSNMP1. The patterns of olfactory protein expression during larval development have also been studied. For example, an OBP (Cmeg33593_c0) in *C. megacephala* is increasingly expressed from the first to the third instar larval stages, and the larval olfactory protein expression profile indicates that some proteins are expressed only in larvae (Wang et al., 2015). Two binding proteins appear to be larva specific in *S. littoralis* (Poivet et al., 2013) and green SexiOBPs in the OBP phylogenetic tree (Llopis-Gimenez et al., 2020), as well as six OBPs and four CSPs are larval tissue specific in *H. armigera* (Chang et al., 2017). More larval olfactory proteins are expressed in both larvae and adults of other species, as demonstrated for several *C. pomonella* ORs that exhibit sex-biased expression in adults, as well as larva-enriched transcription (Walker et al., 2016). Based on these results, the expression profile of *E. signifer* olfactory proteins should be...
further explored in the antennae and mouthparts and at various developmental stages.

CONCLUSION

We identified 39 olfactory proteins in *E. signifer* larvae, with *EsigOBP2, EsigOBP8, EsigGOBP1, EsigGOBP2, EsigGOBP5, EsigCSP3, EsigCSP5,* and *EsigOR1* expressed most strongly in the head. CSP3 was expressed only in the head, where it plays key roles in sensing *Eucalyptus-*derived compounds, whereas *EsigGOBP2, EsigCSP5,* and *EsigOR1* exhibited biased expression. *EsigGR1* exhibited the highest expression among all tissues, which may correlate with the feeding habits of *E. signifer* larvae based on gustatory preferences. The functions of *EsigGR1* and *EsigCSP3* in larval olfactory
and gustatory recognition should be explored further. Most olfactory proteins were highly expressed in thoracic and abdominal teguments, establishing the molecular basis for the head as the center of sensation and explaining how borers use the thorax and abdomen to sense the wood hole environment.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ncbi.nlm.nih.gov/, PRJNA713545.
AUTHOR CONTRIBUTIONS

XYZ carried out the molecular genetic studies, performed the sequence alignment, experiments and drafted the manuscript. XHY, XHM, and XML collected all samples and participated in some experiments. PH and ZDY designed and conceived of the study. PH also helped to draft the manuscript. All authors read and approved the final manuscript.

FUNDING

This work was supported by the National Natural Science Foundation of China (Grant No. 32001321), the Guangxi Natural Science Foundation under (Grant No. 2020JJA130068), the Special Fund for Science and Technology Bases and Talents of Guangxi Zhuang Autonomous Region (Grant No. 2020AC19057).

ACKNOWLEDGMENTS

We thank Yao Shan and Changxiong Wu for insect collection.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphys.2021.682537/full#supplementary-material

REFERENCES

Annina, H., Melanie, E., Nazli, G., Collins, K. a. n., Kathrin, B. P., Lyubov, P., et al. (2017). Anatomy and behavioral function of serotonin receptors in Drosophila melanogaster larvae. PLoS ONE 12:e0181865. doi: 10.1371/journal.pone.0181865

Bose, C., Basu, S., Das, N., and Khurana, S. (2015). Chemosensory apparatus of Drosophila larvae. Bioinformation 11, 185–188. doi: 10.6026/97320630011185

Carvalho, W. J., Fujimura, P. T., Bonetti, A. M., Goulart, L. R., Cloanon, K., Da Silva, N. M., et al. (2017). Characterization of antennal sensilla, larval morphology and olfactory genes of Melipona scutellaris stingless bee. PLoS ONE 12:e0174857. doi: 10.1371/journal.pone.0174857

Chang, H., Ai, D., Zhang, J., Dong, S., Liu, Y., and Wang, G. (2017). Candidate odorant binding proteins and chemosensory proteins in the larval chemosensory tissues of two closely related noctuidae moths, Helicoverpa armigera and H. assulta. PLoS ONE 12:e0179243. doi:10.1371/journal.pone.0179243

Chen, X., and Hu, P. (in press). Reference gene for tissue and instar of Endoclita signifier larvae. Chin. J. Appl. Entomol. 1–7. doi: 10.19668/j.cnki.cn11-1688/20210008

Di, C., Ning, C., Huazng, L. Q., and Wang, C. Z. (2017). Design of larval chemical attractants based on odorant response spectra of odorant receptors in the cotton bollworm. Insect. Biochem. Molec. 84, 48–62. doi:10.1016/j.ibmb.2017.03.007

Hu, P., Yang, X. H., and Yang, Z. D. (2021). Morphology and distribution of sensillum in antennae, thoracic and abdominal tegument of Endoclita signifier larva. J. Beijing Forestry Univ. in press.

Jin, R., Liu, N.-Y., Liu, Y., and Dong, S.-L. (2015). A larval specific OBP able to bind the major female sex pheromone component in Spodoptera exigua (Hübner). J. Integr. Agr. 14, 1356–1366. doi: 10.1007/s10295-3119(14)60849-2

Jing, D., Zhang, T., Bai, S., He, K., Prabu, S., Luan, J., et al. (2020). Sexual-bisexual gene expression of olfactory-related genes in the antennae of Conogethes pinicolalis (Lepidoptera: Crambidae). BMC Genomics 21:244. doi: 10.1186/s12864-020-6664-3

Li, B., and Dewey, C. N. (2011). RSEM: accurate transcript quantification from RNA-Seq data with or without a reference genome. BMC Bioinformatics 12, 93–99. doi: 10.1186/1471-2105-12-323

Li, X., Dong, G., Fang, J., Liu, H., Guo, W., and Yin, H. (2020). Identification of putative olfactory genes in newly hatched larvae of a Coleopteran ectoparasitoid Dastarcus helophoroides (Fairmaire) (Coleoptera: Bothrideridae) by transcriptome analysis. Entomol Res 50, 329–342. doi: 10.1111/1748-5967.12436

Liu, N. Y., Zhang, T., Ye, Z. F., Li, F., and Dong, S. L. (2015). Identification and characterization of candidate chemosensory gene families from Spodoptera exigua developmental transcriptomes. Int. J. Biol. Sci. 11, 1036–1048. doi: 10.7150/ijbs.12020

Liva, K. J., and Schmittgen, T. D. (2001). Analysis of relative gene expression data using real-time quantitative PCR and the 2−ΔΔCT method. Methods 25, 402–408. doi: 10.1006/meth.2001.1262

Llopis-Gimenez, A., Carrasco-Oltra, T., Jacquin-Joly, E., Herrero, S., and Crava, C. M. (2020). Coupling transcriptomics and behaviour to unveil the olfactory system of Spodoptera exigua larvae. J. Chem. Ecol. 46, 1017–1031. doi: 10.1007/s10886-020-01224-z

Ma, C., Cui, S., Tian, Z., Zhang, Y., Chen, G., Gao, X., et al. (2019). OcomCSP12, a chemosensory protein expressed specifically by ovary, mediates reproduction in Ophryoxis communis (Coleoptera: Chrysomelidae). Front. Physiol. 10:1290. doi: 10.3389/fphys.2019.01290

Mortazavi, A., Williams, B. A., Mccue, K., Schaeffer, L., and Wold, B. (2008). Mapping and quantifying mammalian transcriptomes by RNA-Seq. Nat. Methods 5, 621–628. doi: 10.1038/nmeth.1226

Poivet, E., Gallot, A., Montagne, N., Glaser, N., Legeai, F., and Jacquin-Joly, E. (2013). A comparison of the olfactory gene repertoires of adults and larvae in the noctuid moth Spodoptera littoralis. PLoS ONE 8:e60623. doi: 10.1371/journal.pone.0060263

Rahn, T., Leippe, M., Roeder, T., and Fedders, H. (2013). EGFR signaling in the brain is necessary for olfactory learning in Drosophila larvae. Learn Mem. 20:194–200. doi:10.1016/j.nlm.2013.11.002

Saitou, N., and Nei, M. (1987). The neighbor-joining method: a new method for reconstructing phylogenetic trees. Mol. Biol Evol. 4, 406–425

Tamura, K., Peterson, D., Peterson, N., Stecher, G., Nei, M., and Kumar, S. (2011). MEGA5: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. Mol. Biol. Evol. 28, 2731–2739. doi: 10.1093/molbev/msr121

Walker, W. B. 3rd, Gonzalez, F., Garczynski, S. F., and Witzgall, P. (2016). The chemosensory receptors of codling moth Cydia pomonella expression in larvae and adults. Sci. Rep. 6:23518. doi:10.1038/srep23518

Wang, X., Xiong, M., Lei, C., and Zhu, F. (2015). The developmental transcriptome of the synaptorphic fly Chrysomya megacephala and insights into olfactory proteins. BMC Genomics 16:20. doi: 10.1186/s12864-014-1200-y

Waris, M. I., Younas, A., Ameen, A., Rasool, F., and Wang, M. Q. (2020). Expression profiles and biochemical analysis of chemosensory protein 3 from Nilaparvata lugens (Hemiptera: Delphacidae). J. Chem. Ecol. 46, 363–377. doi: 10.1007/s10886-020-01166-6

Yang, X., Luo, Y., Wu, Y., Zou, D., Hu, P., and Wang, J. (2021). Distribution and damage of endoclita signifier walker, as an important wood borer pest insect on forest. Forest Pest Dis 1–7.

Yang, X., Yang, X., Xue, D., and Han, H. (2016). The complete mitochondrial genome of Endoclita signifier (Lepidoptera, Hepialidae). Mitochondrial DNA A DNA Mapp. Seq. Anal. 27, 4620–4621. doi: 10.3109/19401736.2015.1101585

Yang, X. H. (2013). Studies on the Biological and Ecological Characteristics of Endoclita signifier. Beijing: Beijing Forestry University.

Yang, X. H. (2017). Biological Ecology and Control Techniques of Endoclita signifier, an Important Pest of Eucalyptus. Beijing: China Forestry Publishing House
Zhang et al. Olfactory Proteins of Endoclita signifier

Yi, X., Shi, S., Wang, P., Chen, Y., Lu, Q., Wang, T., et al. (2019). Characterizing potential repelling volatiles for “push-pull” strategy against stem borer: a case study in Chilo auricilius. BMC Genomics 20:751. doi: 10.1186/s12864-019-6112-4

Younas, A., Waris, M. I., Tahir ul Qamar, M., Shaaban, M., Prager, S. M., and Wang, M. Q. (2018). Functional analysis of the chemosensory protein MsepCSP8 from the oriental armyworm Mythimna separata. Front. Physiol. 9:872. doi: 10.3389/fphys.2018.00872

Zhou, J.-J. (2010). Chapter ten-odorant-binding proteins in insects. Vitam Horm. 83, 241–272. doi: 10.1016/S0083-6729(10)8010-9

Zhu, J., Ban, L., Song, L. M., Liu, Y., Pelosi, P., and Wang, G. (2016). General odorant-binding proteins and sex pheromone guide larvae of Plutella xylostella to better food. Insect. Biochem. Mol. Biol. 72, 10–19. doi: 10.1016/j.ibmb.2016.03.005

Zielonka, M., Gehrke, P., Badeke, E., Sachse, S., Breer, H., and Krieger, J. (2016). Larval sensilla of the moth Heliothis virescens respond to sex pheromone components. Insect. Mol. Biol. 25, 666–678. doi: 10.1111/imbo.12353

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2021 Zhang, Yang, Yang, Ma, Liu and Hu. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.