Sensory gene identification in the transcriptome of the ectoparasitoid Quadrastichus mendeli

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Sensory genes play a key role in the host location of parasitoids. To date, the sensory genes that regulate parasitoids to locate gall-inducing insects have not been uncovered. An obligate ectoparasitoid, Quadrastichus mendeli Kim & La Salle (Hymenoptera: Eulophidae: Tetrastichinae), is one of the most important parasitoids of Leptocybe invasa, which is a global gall-making pest in eucalyptus plantations. Interestingly, Q. mendeli can precisely locate the larva of L. invasa, which induces tumor-like growth on the eucalyptus leaves and stems. Therefore, Q. mendeli–L. invasa provides an ideal system to study the way that parasitoids use sensory genes in gall-making pests. In this study, we present the transcriptome of Q. mendeli using high-throughput sequencing. In total, 31,820 transcripts were obtained and assembled into 26,925 unigenes in Q. mendeli. Then, the major sensory genes were identified, and phylogenetic analyses were performed with these genes from Q. mendeli and other model insect species. Three chemosensory proteins (CSPs), 10 gustatory receptors (GRs), 21 ionotropic receptors (IRs), 58 odorant binding proteins (OBPs), 30 odorant receptors (ORs) and 2 sensory neuron membrane proteins (SNMPs) were identified in Q. mendeli by bioinformatics analysis. Our report is the first to obtain abundant biological information on the transcriptome of Q. mendeli that provided valuable information regarding the molecular basis of Q. mendeli perception, and it may help to understand the host location of parasitoids of gall-making pests.

Sensory genes play a key role in the life of parasitoids, such as foraging, oviposition site selection, and mating partners. There are two major chemosensory mechanisms through olfaction and taste in which chemical signals are detected by one of the large multigene families that encode chemosensory proteins (CSPs), gustatory receptors (GRs), ionotropic receptors (IRs), odorant-binding proteins (OBPs), sensory receptors (ORs) and sensory neuron membrane proteins (SNMPs). The function of CSPs and OBPs is the first step in the recognition of chemical stimuli from the outside environment. Chemoreceptors (such as GRs, IRs and ORs) are involved in the recognition and identification of various chemical signals and environmental odors to modulate chemical perception. SNMPs are involved in cell signal transduction.

Some sensory genes of parasitoids in Hymenoptera have been identified, including Bethylidae, e.g., Scleroderma sp.2; Braconidae, e.g., Cotesia vestalis Haliday3, Cot. chilonis Matsumura4, Microplitis demolitor Wilkinson5, M. mediator Haliday6, Microcentrus cingulum Brischke7, Aphidiu sp. Haliday8, Ap. ervi Haliday9 and Meteorus pulchricornis Wesmael10; Encyrtidae, e.g., Anastatus japonicus Ashmead11 and Aenasius bambawalei Hayat12; Eupelmidae, e.g., Copidosoma floridanum Ashmead13; Ichneumonidae, e.g., Campeolosita chlorideae Uchida14; Trichogrammatidae, e.g., Trichogramma dendrolimis Matsumura15 and Tric. japonicum Ashmead16; Eulophidae, e.g., Ascodes hispinarum Boucek and Chouioia cunea Yang17,18. Previous studies have revealed that the sensory genes of parasitoids are involved in searching and locating wood-boring pests and leaf-mining pests. For example, Scleroderma sichuanensis Xiao can accurately find the location of their hidden host Monochamus alternatus Hope and then parasitize them. SsicOBP1 and SsicOBP2 are the basis for the behavior of the odor, which have shown a strong reaction with (−)-α-pinene, (+)-β-pinene, camphene, and (+)-3-carene19. However, the sensory genes of parasitoids used to locate gall-making pests have not yet been solved, which has aroused great interest. Understanding this information can provide potential molecular targets for research based on reverse chemical ecology.
**Materials and methods**

**Insects.** Branches of saplings damaged by *L. invasa* were collected from Guangxi University (108°29' E, 22°85' N), Nanning City, Guangxi Zhuang Autonomous Region, in October 2018. Specimens were placed in a glass container filled with water to retain freshness and transferred to a sealed net cage (length × width × height = 40 cm × 40 cm × 80 cm) with 70–80% relative humidity and a natural light photoperiod maintained at 27 ± 1 °C. The water in the glass container was replaced daily until the emergence of *Q. mendeli*. The emerged *Q. mendeli* were collected daily using 50-mL plastic tubes. One day later, the tubes were immediately placed into liquid nitrogen and stored at −80 °C. Six groups of female *Q. mendeli* adults (a group of twenty) were prepared for RNA extraction.

**RNA sequencing.** A NanoPhotometer spectrophotometer (Thermo Fisher Scientific, Massachusetts, USA) and the Nano6000 Assay Kit for the Agilent Bioanalyzer 2000 system (Agilent Technologies, California, USA) were applied to check the purity and integrity of the total RNA, respectively. After total RNA extraction, magnetic beads with Oligo dT (Thermo Fisher Scientific, Hampton, USA) were used to enrich mRNA, and then, a fragmentation buffer was added to make it a short fragment. The fragments were sequenced on an Illumina HiSeq4000 (Illumina, California, USA).

**Transcriptome data analysis.** Reads obtained from the sequencing machines included dirty reads containing adapters or low-quality bases, which affected the subsequent assembly and analysis. De novo transcriptome assembly was carried out with the short read assembly program Trinity v3.0. Basic annotation of unigenes includes protein functional annotation, pathway annotation, COG/KOG functional annotation and Gene Ontology (GO) annotation. To annotate the unigenes, we used the BLASTx program (http://www.ncbi.nlm.nih.gov/BLAST) with an E-value threshold of 1e−5 for the NCBI nonredundant protein (Nr) database (http://www.ncbi.nlm.nih.gov), the Swiss-Prot protein database (http://www.expasy.ch/sprot), the Kyoto Encyclopedia of Genes and Genomes (KEGG) database (http://www.genome.jp/kegg), and the COG/KOG database (http://www.ncbi.nlm.nih.gov/COG). Protein functional annotations could then be obtained according to the best alignment results.

**Sequence alignment and phylogenetic analysis.** TransMembrane prediction using Hidden Markov Models 2.0 (http://www.cbs.dtu.dk/services/TMHMM)34. The signal peptides were predicted using SignalP 4.1 (http://www.cbs.dtu.dk/services/SignalP35). Amino acid sequence alignment was performed using the ClustalW method implemented in Mega v7.036. The resulting alignment was manually curated to remove gap-rich regions. Maximum-likelihood trees were constructed using IQ-TREE with the best-fitting substitution model38. Subsequently, trees were viewed and graphically edited in FigTree v1.4.339 and Adobe Illustrator CS6. Branch support was assessed using the bootstrap method based on 1000 replicates.

**Results**

**Transcriptome assembly and annotation.** To obtain high-quality clean reads, raw reads with adapters, low quality, and an N content greater than 10% were removed. The number of clean reads in female adults of *Q. mendeli* ranged from 18,733,252 to 25,259,462, and the sample GC content ranged from 45.06 to 53.01% (Table 1). At the same time, Q20 and Q30 ranged from 92.75 to 94.36% and 88.45 to 90.08% respectively (Table 1). In total, 31,820 transcripts were obtained and assembled into 26,925 unigenes (Additional file 1). A total of 42.10% of unigenes had a length greater than 2000 bp and an average length of 1369 bp (Table 2). The NR database (15,543, 57.73%) had the largest match. In general, the sequences had E-values between 0 and 1E−150.

**Materials and methods**
and the sequence retrieved, and the higher the score, the greater the degree of similarity between them. The transcripts of *Q. mendeli* were most similar to the sequences of *Nasonia vitripennis* Walker (26.82%), followed by the sequences of *Ceratosolen solmsi marchali* Mayr (7.49%), *Cop. floridanum* (3.77%), *Tric. pretiosum* (2.92%), and other species (44.27%) (Additional file 2). SwissProt (11,644, 43.25%) and KOG (10,924, 40.57%) shared similar quantities; KO (7524, 27.94%) showed the least match (Table 3).

In total, 14,735 were annotated into 52 subcategories belonging to three main GO categories: a ‘biological process’, ‘cellular component’ and ‘molecular function’ (Fig. 1a). There were 22 subcategories in the ‘biological process’, 18 subcategories in the ‘cellular component’, and 12 subcategories in the ‘molecular function’. The top ten subcategories were ‘catalytic activity’ (1580), ‘metabolic process’ (1552), ‘binding’ (1552), ‘cellular process’ (1539), ‘single-organism process’ (1312), ‘cell’ (932), ‘cell part’ (932), ‘membrane’ (639), ‘biological regulation’ (615) and ‘organelle’ (576) (Additional file 3). By KOG classifications, 4689 unigenes were classified functionally into 25 categories (Fig. 1b). The cluster of ‘general fractional prediction only’ was the largest group, which had 4855 unigenes. The ‘signal transduction mechanisms’ group was second with 3998 unigenes. The top 2 categories had 36.64% unigenes annotated to the KOG database (Additional file 4). In total, 5160 unigenes were functionally classified into 5 KEGG categories (Fig. 1c). They were ‘cellular processes’ (587 unigenes, 7.66% of the unigenes annotated to the KEGG database), ‘environmental information processing’ (739, 9.64%), ‘genetic information processing’ (1612, 21.03%), ‘metabolism’ (4504, 58.77%) and ‘organismal systems’ (222, 2.90%) (Additional file 4). Among the 31 subcategories, ‘Global and Overview’ (2123, 27.70%), ‘translation’ (664, 8.66%) and ‘Signal transduction’ (599, 7.82%) were the top 3 (Additional file 5).

### Identification of candidate chemosensory genes.

In this study, 3 putative unigenes encoding CSPs were identified, named QM_comp07737, QM_comp08732 and QM_comp26540 (Additional file 6). The lengths of these unigenes were 509 bp, 608 bp and 280 bp, respectively (Additional file 7). Among these unigenes, QM_comp07737 and QM_comp26540 were incomplete due to a lack of a 5’ or 3’ terminus (Additional file 7). QM_comp08732 sequences were full-length putative CSP genes because they had complete ORFs and 4 cysteines.

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**Table 1.** Sequencing summary of the *Quadrastichus mendeli* transcriptome.

| Sample name | Raw reads | Clean reads | Clean data (Gb) | Q20 (%) | Q30 (%) | GC content |
|-------------|-----------|-------------|----------------|---------|---------|-------------|
| *Q. mendeli* 1 | 24,740,386 | 21,110,044 | 3.71 | 94.03 | 89.45 | 45.79 |
| *Q. mendeli* 2 | 22,581,498 | 19,278,188 | 3.39 | 94.13 | 89.65 | 45.22 |
| *Q. mendeli* 3 | 29,819,798 | 25,259,462 | 4.47 | 93.85 | 89.11 | 45.06 |
| *Q. mendeli* 4 | 25,051,518 | 18,733,252 | 3.76 | 92.75 | 88.45 | 53.01 |
| *Q. mendeli* 5 | 26,482,834 | 22,495,242 | 3.97 | 93.94 | 89.31 | 45.26 |
| *Q. mendeli* 6 | 26,557,838 | 22,736,816 | 3.98 | 94.36 | 90.08 | 45.18 |

**Table 2.** Number and length of transcripts and unigenes.

| Length range/bp | Contig | Transcript | Unigene |
|-----------------|-------|------------|--------|
| 0–300           | 442,916 (95.09%) | 7976 (25.07%) | 5859 (21.76%) |
| 301–500         | 5879 (1.26%) | 6191 (19.46%) | 4816 (17.89%) |
| 501–1000        | 5696 (1.22%) | 5739 (18.04%) | 4915 (18.25%) |
| 1001–2000       | 5660 (1.22%) | 5694 (17.89%) | 5315 (19.74%) |
| >2001           | 11,287 (2.43%) | 11,914 (37.44%) | 11,335 (42.10%) |
| Total number    | 465,788 | 31,820 | 26,925 |
| Total length (bp) | 66,378,092 | 39,430,660 | 36,871,436 |
| N50 length (bp)  | 1617 | 2356 | 2504 |
| Mean length (bp) | 143 | 1239 | 1369 |

**Table 3.** Unigenes annotated in different databases. NR NCBI non-redundant protein sequences, Swissprot A manually annotated and reviewed protein sequence database, KO KEGG Orthology, KOG Clusters of Orthologous Groups of proteins, Total no. total number of annotated unigenes, PCT (%) percentage (%).
which are characteristic of typical insect CSPs. QM_comp08732 with a molecular weight of 17 kDa had a signal peptide sequence of approximately 22 amino acids at the N-terminus (Additional file 8). Through a homology search with known proteins, the results showed that 73% of QM_comp26540 was orthologs of the proteins in *Tenebrio molitor* L., and the orthologs of other CSP sequences were also above 60% (Additional file 8). A phylogenetic tree based on the maximum likelihood method was constructed using the 3 CSP sequences of *Q. mendeli* along with 67 CSP sequences from 6 other species (i.e., *A. mellifera*, *B. mori*, *D. melanogaster*, *M. mediator*, *S. invicta* and *Trib. castaneum*) (Fig. 2 and Additional file 9). The phylogenetic tree showed that QM_comp26540 shares a high homology and is closely clustered with MmedCSP1, which has been functionally characterized, and QM_comp07737 and QM_comp08732 did not branch clusters with any other insects; they may be specific CSPs of *Q. mendeli* (Fig. 2).

**Identification of candidate gustatory receptors.** Ten candidate GR proteins were identified from the data sets (Additional file 6). Among these unigenes, QM_comp03300, QM_comp23544, QM_comp24536 and QM_comp26507 were incomplete due to the lack of a 5’ or 3’ terminus (Additional file 7). QM_comp00164, QM_comp03333, QM_comp11847, QM_comp15910, QM_comp22611 and QM_comp22814 sequences were full-length putative GR genes because they had complete ORFs. These unigenes had molecular weights that ranged between 4 and 56 kDa and had a signal peptide sequence that ranged between 15 and 41 amino acids at the N-terminus (Additional file 8). Through a homology search with known proteins, the results showed that 79% of QM_comp15910 were orthologs of the proteins in *Trichomalopsis sarcophagae* Gahan. A phylogenetic tree based on the maximum likelihood method was constructed using the 10 GR sequences of *Q. mendeli* along with 191 GR sequences from 9 other species (i.e., *A. mellifera*, *B. mori*, *C. floridanum*, *D. melanogaster*, *M. mediator*, *N. vitripennis*, *S. invicta*, *Trib. castaneum* and *Tric. pretiosum*) (Fig. 3 and Additional file 9). The phylogenetic
Identification of candidate ionotropic receptors. Twenty-one candidate IR proteins were identified from the data sets (Additional file 6). Among these unigenes, 13 unigenes were incomplete due to the lack of a 5’ or 3’ terminus (Additional file 7). Eight unigenes were full-length putative IR genes because they had complete ORFs. These unigenes had molecular weights that ranged between 5 and 12 kDa (Additional file 8). Through a homology search with known proteins, the results showed that 100% of the IRs in *Q. mendeli* were orthologs of the proteins in *Asbolus verrucosus* LeConte, and the orthologs of other IR sequences were also above 37% (Additional file 8). A phylogenetic tree based on the maximum likelihood method was constructed using the 21 IR sequences of *Q. mendeli* along with 131 IR sequences from 9 other species (i.e., *A. mellifera*, *B. mori*, *C. floridanum*, *D. melanogaster*, *M. mediator*, *N. vitripennis*, *S. invict*, *T. castaneum* and *T. pretiosum*) (Fig. 4 and Additional file 9). The 21 IRs of *Q. mendeli* along with 131 IRs from 9 other species (i.e., *A. mellifera*, *B. mori*, *C. floridanum*, *D. melanogaster*, *M. mediator*, *N. vitripennis*, *S. invict*, *T. castaneum* and *T. pretiosum*) were chosen to construct a phylogenetic tree based on the amino acid sequences (Additional file 9). The phylogenetic tree showed that all candidate IR proteins were clustered with at least one Hymenoptera ortholog (Fig. 4).

Identification of candidate odorant binding proteins. Fifty-six candidate OBP proteins were identified from the data sets (Table 4; Additional file 6). Among these unigenes, 10 unigenes were incomplete due to the lack of a 5’ or 3’ terminus (Additional file 7). Forty-six unigenes were full-length putative OBP genes because they had complete ORFs. These unigenes had molecular weights ranging between 10 and 17 kDa and had a signal peptide sequence ranging between 16 and 23 amino acids at the N-terminus (Additional file 8). Insect OBPs can be classified into classical OBPs (six-cysteine conserved signature) and Minus-C (missing C2 and C5) and Plus-C (carries an additional conserved cysteine located between C1 and C2 and after C6). QM_comp02388, QM_comp07285, QM_comp08846, QM_comp10855, QM_comp21133, QM_comp21238 and QM_comp24139 had four conserved cysteines, which were minus-COBPs. Other OBPs had six conserved cysteines, which are classic OBPs. Plus-COBPs were not found in hymenopteran species. Through a homology search with known
proteins, the results showed that 88% of QM_comp19239 were orthologs of the proteins in *N. vitripennis*, and the orthologs of other OBP sequences were also above 42% (Additional file 8). A phylogenetic tree based on the maximum likelihood method was constructed using the 56 OBP sequences of *Q. mendeli* along with 209 OBP sequences from 9 other species (i.e., *A. mellifera*, *B. mori*, *C. floridanum*, *D. melanogaster*, *M. mediator*, *N. vitripennis*, *Q. mendeli*, *Trib. castaneum*, and *Tric. pretiosum*) (Fig. 5 and Additional file 9). The phylogenetic tree showed that all candidate OBP proteins were clustered with at least one Hymenoptera ortholog (Fig. 5).

**Identification of candidate odorant receptors.** Thirty candidate OR proteins were identified from the data sets (Additional file 6). Among these unigenes, 21 unigenes were incomplete due to the lack of a 5' or 3' terminus (Additional file 7). Nine unigenes were full-length putative OR genes because they had complete ORFs. These unigenes had molecular weights ranging between 4 and 53 kDa and had a signal peptide sequence ranging between 16 and 23 amino acids at the N-terminus (Additional file 8). Through a homology search with known proteins, the results showed that 95% of QM_comp14333 were orthologs of the proteins in *C. cunea*, and the orthologs of other OR sequences were also above 32% (Additional file 8). A phylogenetic tree based on the maximum likelihood method was constructed using the 30 OR sequences of *Q. mendeli* along with 235 OR sequences from 9 other species (i.e., *A. mellifera*, *B. mori*, *C. floridanum*, *D. melanogaster*, *M. mediator*, *N. vitripennis*, *S. invict*, *Trib. castaneum* and *Tric. pretiosum*) (Fig. 5 and Additional file 9). The phylogenetic tree showed that all candidate OR proteins were clustered with at least one Hymenoptera ortholog (Fig. 5).

**Identification of candidate sensory neuron membrane receptors.** Two candidate SNMP proteins were identified from the data sets (Additional file 6). QM_comp21591 was incomplete due to the lack of a 3' terminus (Additional file 7). QM_comp09081 was a full-length putative SNMP gene because it had complete ORFs.
QM_comp09081 had a signal peptide sequence of 13 amino acids at the N-terminus, and the molecular weight of QM_comp09081 was 59 kDa (Additional file 8). According to sequence similarity, SNMP is divided into two SNMP subtypes, SNMP1 and SNMP2. Through a homology search with known proteins, the results showed that 67% of QM_comp09081 were orthologs of the proteins in *N. vitripennis* (Additional file 8). A phylogenetic tree based on the maximum likelihood method was constructed using the 2 SNMP sequences along with 25 SNMP sequences from 9 other species (i.e., *A. mellifera*, *B. mori*, *C. floridanum*, *D. melanogaster*, *M. mediator*, *N. vitripennis*, *S. invict*, *Trib. castaneum* and *Tric. pretiosum*) (Fig. 7 and Additional file 9). The phylogenetic tree showed that QM_comp21591 fell into the same clade as the insect SNMP1 group, and QM_comp09081 fell into the same clade as the insect SNMP2 group (Fig. 7).

**Discussion**

In this study, the major sensory genes (i.e., CSPs, GRs, IRs, OBPs, ORs and SNMPs), which perhaps regulate *Q. mendeli* to locate its host *L. invasa*, are first reported, providing valuable information for exploring how parasitoids use sensory genes to locate gall-making pests.

CSPs are widespread in the antenna and other chemical sensory organs of insects and are involved in the chemical perception and related behavior of insects. Compared to the total number of insects in the world, CSPs have been identified in only a few species of insects to different degrees, such as Coleoptera, Hemiptera, Hymenoptera, Lepidoptera and Orthoptera, whose numbers show interspecific diversity. For example, the number of CSPs varies from 4 CSPs in *Drosophila melanogaster* to 22 in *B. mori*. In this study, 3 candidate CSPs were identified, which are less than those in the parasitoids *Ch. cunea* (11), *Scleroderma* sp. (10), *Me. pulchricornis* (8) and *Tric. dendrolimi* (7) and greater than those in the parasitoids *Cot. chilonis* (2) and *Ap. ervi* (2). Previous research confirmed that generalists seem to be specifically suited for the processing of odorant mixtures, and they respond in a similar manner to plant volatiles. For example, *Ch. cunea* is a generalist and...
| Unigene ID  | Unigene length (bp) | ORF length (aa) | Complete ORF | 5' or 3' terminus lost | Signal peptide (aa) | Cysteine number | FPKM (Mean) | Homology search with known proteins |
|------------|---------------------|----------------|--------------|------------------------|---------------------|----------------|-------------|------------------------------------|
| QM_comp01352 | 728                 | 132            | Yes          | –                      | Yes                 | 18             | 8           | 14.67 | 37 2e–20 C. florida-num  | XP_014204137.1 |
| QM_comp02170 | 549                 | 129            | Yes          | –                      | Yes                 | 19             | 7           | 234.80  | 53 2e–21 N. vitripennis  | XP_001600573.1 |
| QM_comp02233 | 411                 | 129            | No           | 3'                     | Yes                 | 21             | 8           | 0.32    | 34 2e–07 C. florida-num  | XP_014212211.1 |
| QM_comp02388 | 1415                | 125            | Yes          | –                      | No                  | –              | 5           | 1.67    | 58 1e–150 Habropoda laboriosa | KOC59862.1 |
| QM_comp02394 | 438                 | 130            | Yes          | –                      | Yes                 | 19             | 8           | 17.91   | 43 5e–27 N. vitripennis  | XP_001600573.1 |
| QM_comp02616 | 679                 | 157            | Yes          | –                      | No                  | –              | 9           | 203.66  | 68 2e–46 N. vitripennis  | XP_001601182.1 |
| QM_comp02693 | 407                 | 118            | No           | 3'                     | Yes                 | 19             | 7           | 5.73    | 52 7e–26 N. vitripennis  | XP_001600573.1 |
| QM_comp03741 | 518                 | 129            | Yes          | –                      | Yes                 | 19             | 7           | 1.12    | 40 6e–18 N. vitripennis  | XP_001600573.1 |
| QM_comp04039 | 793                 | 133            | Yes          | –                      | Yes                 | 18             | 9           | 5.00    | 34 1e–08 C. florida-num  | XP_014204137.1 |
| QM_comp04767 | 321                 | 41             | No           | 3'                     | No                  | –              | 0           | 0.58    | 48 2e–09 C. florida-num  | XP_014204137.1 |
| QM_comp05191 | 915                 | 113            | Yes          | –                      | Yes                 | 20             | 10          | 10.24   | 41 7e–12 C. florida-num  | XP_014204137.1 |
| QM_comp05947 | 727                 | 133            | Yes          | –                      | Yes                 | 19             | 6           | 35.75   | 41 5e–14 N. vitripennis  | XP_001600573.1 |
| QM_comp06244 | 433                 | 127            | No           | 3'                     | Yes                 | 17             | 8           | 8.89    | 63 5e–45 T. dew-droloim  | ANG08504.1 |
| QM_comp06765 | 917                 | 150            | Yes          | –                      | Yes                 | 23             | 7           | 3.90    | 52 2e–41 Ceratosolen solmsi marchali | XP_011505749.1 |
| QM_comp07285 | 729                 | 92             | Yes          | –                      | No                  | –              | 4           | 1.40    | 86 1e–09 C. florida-num  | XP_014206764.1 |
| QM_comp08027 | 538                 | 140            | Yes          | –                      | Yes                 | 19             | 8           | 113.56  | 49 2e–39 N. vitripennis  | XP_001600573.1 |
| QM_comp08037 | 736                 | 134            | Yes          | –                      | Yes                 | 19             | 9           | 192.11  | 37 9e–24 N. vitripennis  | XP_001600573.1 |
| QM_comp08573 | 1211                | 133            | Yes          | –                      | Yes                 | 21             | 6           | 4.55    | 36 2e–06 C. florida-num  | XP_014212211.1 |
| QM_comp08613 | 542                 | 113            | Yes          | –                      | Yes                 | 20             | 9           | 21.22   | 44 2e–12 C. florida-num  | XP_014204137.1 |
| QM_comp08638 | 476                 | 130            | Yes          | –                      | Yes                 | 20             | 7           | 311.84  | 51 2e–25 C. florida-num  | XP_014204137.1 |
| QM_comp08676 | 507                 | 135            | Yes          | –                      | Yes                 | 17             | 7           | 275.44  | 69 6e–63 N. vitripennis  | XP_001601068.1 |
| QM_comp08846 | 492                 | 125            | Yes          | –                      | Yes                 | 20             | 7           | 62.90   | 38 6e–08 C. florida-num  | XP_014204137.1 |
| QM_comp08899 | 580                 | 128            | Yes          | –                      | Yes                 | 18             | 7           | 43.75   | 35 1e–12 N. vitripennis  | XP_001606346.1 |
| QM_comp08900 | 484                 | 129            | Yes          | –                      | Yes                 | 18             | 7           | 125.85  | 36 2e–13 N. vitripennis  | XP_001606346.1 |
| QM_comp09538 | 510                 | 131            | Yes          | –                      | Yes                 | 19             | 11          | 59.68   | 42 4e–23 N. vitripennis  | XP_001600573.1 |
| QM_comp09339 | 507                 | 131            | Yes          | –                      | Yes                 | 19             | 9           | 176.04  | 51 3e–19 N. vitripennis  | XP_001600573.1 |
| QM_comp09356 | 647                 | 145            | Yes          | –                      | Yes                 | 22             | 6           | 4.84    | 53 5e–51 N. vitripennis  | XP_001603472.2 |
| QM_comp09551 | 688                 | 108            | Yes          | –                      | Yes                 | 20             | 9           | 9.27    | 43 1e–08 C. florida-num  | XP_014204137.1 |
| QM_comp10209 | 523                 | 137            | Yes          | –                      | Yes                 | 17             | 7           | 36.04   | 45 5e–30 N. vitripennis  | XP_001601290.1 |
| QM_comp10426 | 498                 | 124            | Yes          | –                      | Yes                 | 19             | 9           | 7.66    | 33 1e–08 N. vitripennis  | XP_001606053.1 |
| QM_comp10855 | 879                 | 144            | Yes          | –                      | No                  | –              | 4           | 8.33    | 28 7e–07 N. vitripennis  | XP_001601068.1 |
| QM_comp11668 | 700                 | 126            | Yes          | –                      | Yes                 | 18             | 8           | 45.36   | 34 7e–12 C. florida-num  | XP_014204137.1 |
| QM_comp12532 | 510                 | 135            | Yes          | –                      | Yes                 | 16             | 7           | 3.32    | 74 6e–57 N. vitripennis  | XP_001601068.1 |

Continued
has multiple hosts (e.g., *Stilpnotia salicis* L., *Ivela ochropoda* Eversmann, *Clostera anachoreta* Fabricius, *Semiothosa cinerea* Bremer & Gray and *Clania variegata* Snellen), while *Cot. chilonis* mainly parasitizes larvae of the genus *Chilo* Zincken48. It can be deduced that the number of CSPs relates to their host range. For the specialist *Q. mendeli*, use of CSPs is expected to cope with the variability in host availability46. Previous studies also revealed that CSPs could be involved in the solubilization of hydrocarbons in the stratum corneum to recognize *Q. mendeli*′s host trees. A similar story has been confirmed for *M. mediator* and its shelter host eucalyptus indicating that QmenCSPs may function in the chemical sensing of *L. invasa* and its host range.

**Table 4.** Detailed information on the OBP ungenes of *Quadrastichus mendeli*.

| Unigene ID | ORF length (bp) | Complete ORF | 5′ or 3′ terminus lost | Signal peptide length (bp) | Signal peptide (aa) | Cysteine number | FPKM (Mean) | Identity (%) | E value | Species | Protein ID |
|------------|-----------------|--------------|------------------------|---------------------------|---------------------|----------------|-------------|-------------|---------|----------|------------|
| QM_comp12533 | 669             | Yes          | –                      | Yes                       | 16                  | 7              | 21.25       | 74          | 2e–56   | N. vitripennis | XP_001601068.1 |
| QM_comp13483 | 1445            | Yes          | –                      | Yes                       | 19                  | 8              | 144.20      | 44          | 4e–14   | C. floridanum  | XP_014204137.1 |
| QM_comp18897 | 568             | Yes          | –                      | Yes                       | 20                  | 9              | 2.74        | 26          | 4e–6    | N. vitripennis | XP_001600573.1 |
| QM_comp20903 | 596             | Yes          | –                      | Yes                       | 20                  | 11             | 13.86       | 27          | 1e–5    | C. floridanum  | XP_014206340.1 |
| QM_comp21133 | 508             | Yes          | –                      | No                        | 19                  | 6              | 0.95        | 29          | 3e–5    | Ceratosolen solmis marceli | XP_011505723.1 |
| QM_comp21238 | 482             | Yes          | –                      | No                        | 19                  | 5              | 0.59        | 32          | 1e–6    | N. vitripennis | XP_001600573.1 |
| QM_comp21371 | 321             | Yes          | –                      | Yes                       | 19                  | 6              | 1.40        | 36          | 3e–9    | N. vitripennis | XP_001600573.1 |
| QM_comp21830 | 1294            | Yes          | –                      | Yes                       | 23                  | 9              | 73.07       | 79          | 9e–65   | N. vitripennis | XP_001600769.1 |
| QM_comp21900 | 238             | No           | 3′                     | –                         | 1                   | 0.67          | 51          | 32         | 1e–17   | N. vitripennis | XP_001600573.1 |
| QM_comp23080 | 223             | No           | 3′                     | No                        | 19                  | 5              | 0.59        | 32          | 1e–6    | N. vitripennis | XP_001600573.1 |
| QM_comp23536 | 487             | Yes          | –                      | Yes                       | 19                  | 5              | 0.99        | 28          | 6e–7    | T. pretiosum   | XP_014221963.1 |
| QM_comp23819 | 378             | No           | 3′                     | Yes                       | 22                  | 4              | 0.65        | 49          | 4e–29   | N. vitripennis | XP_001603472.2 |
| QM_comp24139 | 1063            | Yes          | –                      | Yes                       | 18                  | 4              | 197.63      | 26         | 3e–4    | C. floridanum  | XP_014206340.1 |
| QM_comp24881 | 456             | Yes          | –                      | Yes                       | 17                  | 8              | 3.80        | 26         | 3e–4    | C. floridanum  | XP_015603838.1 |
| QM_comp24882 | 483             | Yes          | –                      | Yes                       | 17                  | 8              | 1.79        | 28         | 2e–4    | T. pretiosum   | XP_014221963.1 |
| QM_comp26560 | 572             | Yes          | –                      | Yes                       | 22                  | 6              | 40.28       | 69          | 5e–66   | N. vitripennis | XP_016842824.1 |
| QM_comp26834 | 545             | Yes          | –                      | Yes                       | 17                  | 8              | 249.66      | 73         | 8e–49   | AHE40949.1    | XP_001603472.2 |
| QM_comp30395 | 562             | Yes          | –                      | Yes                       | 20                  | 7              | 12.82       | 46         | 6e–28   | C. floridanum  | XP_014208150.1 |
| QM_comp401156 | 497            | No           | 5                      | Yes                       | 20                  | 9              | 93.54       | 30          | 2e–10   | N. vitripennis | XP_016845238.1 |
| QM_comp6538 | 520             | Yes          | –                      | Yes                       | 20                  | 7              | 2.40        | 77          | 8e–67   | T. denivoli     | ANG08495.1    |
| QM_comp6539 | 593             | Yes          | –                      | Yes                       | 20                  | 7              | 43.82       | 73          | 1e–52   | T. pretiosum   | XP_014220461.1 |
| QM_comp10678 | 1344            | Yes          | –                      | Yes                       | 20                  | 6              | 21.55       | 51          | 4e–41   | T. sar- caphalae | OXU16757.1 |
| QM_comp18896 | 1057            | Yes          | –                      | Yes                       | 20                  | 9              | 5.18        | 41          | 4e–25   | C. floridanum  | XP_014208127.1 |

**Table 4.** Detailed information on the OBP ungenes of *Quadrastichus mendeli*.
QM_comp26540 may function as a chemosensor, which is involved in the process of recognizing plant volatiles from eucalyptus when *Q. mendeli* searches its host, *L. invasa*.

GRs are widespread in gustatory organs of insects that respond to various taste-related soluble compounds, and cuticular hydrocarbons and odorants, such as sugars, amino acids, salts, bitter compounds, CO₂ and pheromones, can be recognized and combined by GRs. To date, GRs in some model insects with genome reports have been identified, such as *A. gambiae* (76), *B. mori* (69), *D. melanogaster* (68) and *N. vitripennis* (58). In this study, 10 candidate GRs were identified, which was similar to other parasitoids, such as *An. japonicus* (8), *Sclerodermus* sp. (6), and *M. mediator* (6). This could be attributed to the sequencing depth and species-functional specificity of GRs. DmelGR5 and DmelGR64 in *D. melanogaster* are receptor proteins for sweet taste and are used to detect glucose, sucrose, maltose, maltitol and cottonseed sugar. For *Q. mendeli*, females that were fed a honey solution or honey solution + young eucalyptus leaves lived for a longer time than those who underwent other treatments, including flowers, gall leaves, water, galled leaves + honey solution, no food and young leaves. Therefore, GRs in *Q. mendeli* should play a key role in recognizing sugar and fresh eucalyptus leaves via various soluble compounds. The phylogenetic tree showed that QM_comp00164 and QM_comp22611 were the same clade as the proteins in *Trib. castaneum*, QM_comp03300, QM_comp22814, and QM_comp26507 were the same clade as the proteins in *D. melanogaster*, and other proteins in *Q. mendeli* were the same clade as the proteins in *N. vitripennis*. Thus, GRs of *Q. mendeli* may share high homology and closely cluster with the proteins in *D. melanogaster*, *N. vitripennis*, and *Trib. castaneum*. Interestingly, QM_comp11847 was in the same clade as the sugar receptor NvitGR1, which was used to recognize the only source of nutrients from host *Lucilia caesar* L. for the offspring of *N. vitripennis*. Thus, QM_comp11847 may be involved in recognizing host organisms and sugars, which helps *Q. mendeli* to quickly access energy from these molecules.

IRs, which evolve from the ionotropic glutamate receptor (iGluR), are a new class of sensory proteins mainly in taste organs/sensilla that respond to food components, such as sugars, salts, water and bitter compounds, and detect small temperature differences. In this study, 21 candidate IRs were identified, which was more than in the parasitoids *M. pulchricornis* (19), *C. cunea* (10), *M. mediator* (6), *Sclerodermus* sp. (3), *Mi. cingulum* (3) and *An. japonicus* (3). Physiological recordings from taste sensilla in *D. melanogaster* and other insects have revealed responses of taste neurons to salts, sugars, water, bitter compounds and a large diversity of other

Figure 5. Phylogenetic tree of odorant binding proteins (OBPs) from *Quadrastichus mendeli* and other insects based on the maximum likelihood method. Included are OBPs from *Apis mellifera* (Amel), *Bombyx mori* (Bmor), *Copidosoma floridanum* (Cflo), *Drosophila melanogaster* (Dmel), *Microplitis mediator* (Mmed), *Nasonia vitripennis* (Nvit), *Quadrastichus mendeli* (Qmen), *Tribolium castaneum* (Tcas), and *Trichogramma pretiosum* (Tpre). The specific clades are marked. Node support was assessed with 1000 bootstrap replicates.
tastants. Taste sensilla are widely distributed on the antennae of Q. mendeli, suggesting that QmenIRs may function as taste receptors. The phylogenetic tree showed that QmenIRs were spread across the tree branches and clustered with homologous IRs from other species, which suggested that QmenIRs may be functionally conserved. QM_comp21031 was located in the same clade as BmorIR21a of B. mori, DmelIR21a of D. melanogaster and TcasIR21a of Trib. castaneum, indicating that QM_comp21031 has the closest relationship with insect IR21a, which can mediate cool sensing in Drosophila. Thus, QM_comp21031 may perceive changes in temperature since insect IR21a can achieve both heat avoidance and heating. Taking the oviposition features into consideration, we deduced that female Q. mendeli may be capable of sensing surface heat on galls related to L. invasa damage, which requires further exploration.

OBPs are crucial in insect olfactory perception and are the first step in the recognition of chemical stimuli from the outside environment. In some model insects, OBPs have been identified to different degrees, such as B. mori (57), D. melanogaster (51), Trib. castaneum (46) and A. gambiae (44). In this study, 56 candidate OBPs were identified, which was more than in the parasitoids Ae. bambawalei (54), Ch. cunea (25), Tric. dendrolimi (24), M. mediator (20), Me. pulchricornis (16), T. japonicum (15), Ap. ervi (15), Sclerodermus sp. (10), Cop. floridanum (8), Cot. chilonis (8) and As. hispinarum (8). The number of Q. mendeli OBPs identified was less than that in the parasitoids Cot. vestalis (74). Previous studies revealed that OBPs in parasitoids play a key role in binding and transporting hydrophobic odorants from the environment to sensory receptors. A significant behavioral response to the gall volatiles d-limonene and decanal was observed (unpublished data). Relevant QmenOBPs function in chemical sensing of these volatiles characterizing L. invasa and its shelter host eucalyptus trees, which bioinformatics analysis could help to target. The phylogenetic tree showed that 15 QmenOBPs were the same clade as 5 DmelOBPs of D. melanogaster, and 15 QmenOBPs were the same clade as NvitOBP1 of N. vitripennis, and 3 CfloOBPs of C. floridanum, which suggested that most OBPs of Q. mendeli may share high homology and closely cluster with the proteins in C. floridanum, D. melanogaster, and N. vitripennis.

The evolutionary relationship of Q. mendeli OBPs, as inferred in the phylogenetic tree, indicated that they are orthologous sequences due to the absence of monophyletic groups. Our results showed that QM_comp21238 is the same clade as MmedOBP10 of M. mediator, which is involved in the process of recognizing β-ionone and Nonanal when they find the location of their hidden host A. segetis. Thus, QM_comp21238 may also
be involved in the process of recognizing similar odorants or ligands when *Q. mendeli* locates its shelter host, *L. invasa*.

ORs are thought to play critical roles in the perception of chemosensory stimuli by insects. The number of ORs in parasitoids vary greatly. In this study, 30 candidate ORs were identified, which was more than in the parasitoid *Tric. dendrolimi* (9) and *Mi. cingulum* (9)2. Previous studies revealed that the OR of *M. mediator* play an important role in recognizing plant volatiles, such as nonanal and farnesene, which provided a key start to manipulate and develop ORs in wasps to find hosts and use them as biological tools for pest control. The phylogenetic tree showed that QmenORs were spread across the tree branches and clustered with homologous ORs from other species, which suggested that QmenORs may be functionally conserved. The RNAi investigation of the role of MmedOrco, the *M. mediator* ortholog of *Drosophila* Or83b, supported the assumption that this highly conserved gene plays a similar role in insects. QmenORs may function as chemoreceptors to recognize plant volatiles from eucalyptus. Our results showed that QM_comp20892 is in the same clade as DmelOR10a of *D. melanogaster*, which plays a role in responding to odorants such as methylsalicylate and acetophenone. For *Q. mendeli*, QM_comp20892 may be involved in the process of recognizing similar odorants or ligands.

SNMPs are involved in cellular signal transduction and play a role in odor detection. Two SNMPs are normally broadly identified in different insects, e.g., parasitoids *Ch. cunea* and *Sclerodermus* sp.2. It has been reported that SNMP1 and SNMP2 are both expressed in antennae sensilla and have different expression patterns. In *Ch. cunea*, CcunSNMP1 is a morphine receptor of neurons, and CcunSNMP2 is mainly expressed in supporting cells and the lymph of antennal sensilla, while the location and expression patterns of SNMPs in *Q. mendeli* should be further studied since this information should be associated with their functions. SNMP1 of *D. melanogaster* is involved in pheromone detection and enhances the Ca\(^{2+}\) responses served in signal transduction. SNMP1 of *M. mediator* was determined to participate in both pheromone and general odor detection. In contrast, the general functional mechanism of SNMP2 in parasitoids is still poorly understood. The phylogenetic tree showed that QM_comp21591 was the same clade as TcasSNMP1 of *Trib. castaneum* and QM_comp09081 was the same clade as CflosSNMP2 of *C. floridanum*, suggesting that QM_comp21591 and QM_comp09081 shall be SNMP1 and SNMP2 in *Q. mendeli* respectively. In addition, *Q. mendeli* is a uniparental...
parasitoid that is not required in the male search for mating. Thus, the function of QM_comp21591 and QM_comp09081 may include an oviposition pheromone receptor rather than a sex receptor and a membrane protein with unknown functions, which needs to be further explored.

Chemical detection involves a series of complicated processes that require participation and interactions by multiple cascades of sensory proteins. Insect sensory proteins are capable of functional cooperation and division. Firstly, OBPs and CSPs are both chemically binding proteins to various odorants and can also respond to the same chemicals, e.g., MmedOBP10 and MmedCSP1 are involved in the process of recognizing β-ionone and nonanal when they find the location of their hidden host. Nonanal and β-ionone are involved in the process of recognizing odor substances. For example, IRs are better at detecting long-lasting odor pulses, and they are less sensitive, suggesting that they are better at close-range odor detection. In contrast, ORs are more sensitive and better at resolving brief (low molecular flux) pulsed stimuli. Moreover, features of functional organization have emerged between behavioral response profiles of OBPs and electrophysiological response profiles of ORs. Therefore, the sensory genes in Q. mendeli should systematically act on the process of locating their gall-making host, and the biological functions of these genes and their products are still poorly known. Overall the sensory genes of the wasp reported here provide valuable insight into the molecular mechanisms of olfaction, which help pave the way for the host location of Q. mendeli in gall-making pests.

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Author contributions
This experiment was conceived and coordinated by Z.Y.H., X.Y.W., W.L. and X.L.Z. in this study. Sampling was performed by Z.Y.H. and X.L.Z. Analysis of transcriptome data and sequence was performed by Z.Y.H. and X.Y.W. Z.Y.H., X.Y.W. and X.L.Z. drafted the manuscript. All authors read and approved the final version of the manuscript.

Competing interests
The authors declare no competing interests.

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