Spatial Expression Analysis of Odorant Binding Proteins in Both Sexes of the Aphid Parasitoid *Aphidius gifuensis* and Their Ligand Binding Properties

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In China, *Aphidius gifuensis* is one of the most common endoparasitoids of the green peach aphid *Myzus persicae* and grain aphid *Sitobion miscanthi* in the field. Insect odorant-binding proteins (OBPs) play vital roles in odor perception during feeding, host searching, mating and oviposition. In addition, some OBPs are involved in other physiological processes such as gustation and reproduction. In the present study, a comparative antennal transcriptomic analysis was applied between male and female *A. gifuensis*. The spatial expression patterns among antennae, heads, thoraxes, abdomens and legs of OBPs in both sexes were further profiled. Fifteen *AgifOBPs* were predicted, and 14 of them were identified by gene cloning, including 12 classic OBPs and 2 min-C OBPs. As expected, all OBPs were mainly expressed at high levels in antennae, heads or legs which are sensory organs and tissues. Finally, ligand binding properties of 2 OBPs (*AgifOBP7* and *AgifOBP9*) were further evaluated. Female leg specifically expressed *AgifOBP9* displays a broad and high binding property to aphid alarm pheromones, plant green volatiles and aphid sex pheromones ($K_i < 10 \mu M$). However, female leg specifically expressed *AgifOBP7* displays poor affinity for all tested ligands except CAU-II-11 ((E)-3,7-dimethyl-octa-2,6-dien-1-yl-2-hydroxy-3-methoxybenzoate), a reported (E)-β-farnesene (EBF) analog with an exceptionally high binding affinity ($K_i = 1.07 \pm 0.08 \mu M$). In summary, we reported the spatial expression pattern of the OBP repertoire in *A. gifuensis*, and further studied the binding properties of OBP7 and OBP9, which are mainly expressed in female legs, laying the foundation for the dissection of the contribution of OBPs to chemosensation in *A. gifuensis*.

Keywords: *Aphidius gifuensis*, transcriptome, odorant binding protein, spatial expression pattern, fluorescence binding assay
INTRODUCTION

Aphidius gifuensis is one of the most common endoparasitoids of green peach aphid Myzus persicae and grain aphid Sitobion mitschulskyi in China. S. mitschulskyi is also habitually called Sitobion avenae in China (Zhang, 1999; Jiang et al., 2019), and is the undisputed dominant Chinese dominant pest of wheat. Aphids have long been the most damaging pest of crops and vegetables, causing yield and quality losses by stealing nutrients, transferring plant viruses, and excreting honeydew to block plant photosynthesis (Wu, 2002). In Yunnan and many other areas of China, M. persicae on tobacco has been successfully controlled by artificially released A. gifuensis as a powerful biocontrol tool (Ohita and Honda, 2010; Yang et al., 2009).

The behavioral response of insects to olfactory cues is essentially driven by feeding, reproduction and habitat selection (Pelosi et al., 2014). Molecular odorants enter the sensilla through pores and spread inside the hemolymph on the antennae due to odorant-binding proteins (OBPs) and/or chemosensory proteins (CSPs) (Pelosi et al., 2006; Leal, 2013). These odorants are then transported to olfactory receptors (ORs), ionotrophic receptors (IRs), or sensory neuron membrane proteins (SNMPs), from which the chemical signals will be transmitted into electrophysiological signals for the brain (Leal, 2013; Pelosi et al., 2018). Insect OBPs were initially discovered in antennae of the moth Antheraea polyphemus (Vogt and Riddiford, 1981). Their wide distributions in antennal sensilla indicated the first link of OBPs in the signal chain of odorant perception (Xu et al., 2009). OBPs are tiny, globular, water-soluble proteins with a molecular weight of 10–30 kDa (Pelosi et al., 2005). The presence of six highly conserved cysteine residues, which are paired in three interlocking disulfide bridges to maintain the protein’s tertiary structure, is a common feature of classical OBPs (Pelosi et al., 2014). OBPs act as shuttles for hydrophobic odor molecules, transporting them through the sensillum lymph to odorant receptors (Zhou et al., 2010). After initiating receptors, OBPs may also concentrate odorants in the sensillum lymph and swiftly destroy odorant molecules (Vogt and Riddiford, 1981; Leal, 2013). The prediction of the whole OB family in species became quite simple due to the availability of more insect genomes and transcriptomes using next-generation sequencing techniques. However, the number of OBPs in Hymenoptera varies; for example, Apis mellifera has 21 OBPs, Microplitis mediator has 18 OBPs, Pieris rapae has 14 OBPs, Spodoptera exigua has 34 OBPs, Cotesia vestalis has 20 OBPs, and 90 OBPs were predicted in Nasonia vitripennis (Forêt and Maleszka, 2006; Vieira et al., 2012; Liu et al., 2015; Peng et al., 2017; Li et al., 2020; Liu et al., 2020; Zhou et al., 2021). Insect OBPs not only are expressed in the chemosensory system, but also occur in nonsensory tissues and organs, such as the cornicles (Wang et al., 2021a), thoraces (Xue et al., 2016; Gao et al., 2018; Wang et al., 2019), reproductive organs (Li et al., 2008; Sun et al., 2012b), mandibular glands (Jovinella et al., 2011), salivary glands (Zhang et al., 2017), and wings (Calvello et al., 2003; Pelosi et al., 2005; Wang et al., 2021b). Some insect OBPs have physiological functions other than binding odorants. For example, the sperm carrier function of OBPs has been reported in the male reproductive apparatus of mosquitoes (Li et al., 2008). Moreover, one OBP expressed by male moths is found on the surface of fertilized eggs, which functions to avoid cannibalistic behaviors among larvae (Sun et al., 2012b). Therefore, spatial expression patterns would be helpful to classify and analyze the possible functions of OBPs.

Herbivore-induced volatiles (HIPVs), green leaf volatiles (GLVs), and pheromones such as the aphid alarm pheromone E-beta-farnesene (EBF) are used by natural enemies to find their prey during predation and parasitism (Dong et al., 2008; CMD De Moraes et al., 1998; Buitenhuys et al., 2004). A. gifuensis evolved a comprehensive chemosensory system to effectively detect the semiochemical cues of its host and plants (Yang et al., 2009). For example, A. gifuensis can distinguish healthy, mechanically damaged, and aphid-infested plants (Dong et al., 2008). Additionally, both female and male A. gifuensis were reported to present a positive electroantennogram (EAG) response to EBF and many tobacco volatiles, including trans-2-hexenal, methyl salicylate, benzaldehyde, cis-3-hexen-1-ol, and 1-hexanal (Song et al., 2021a). The volatile sex pheromone has also been shown to be released by female Aphidius, causing intense sexual orientation in males (Fan et al., 2018). OBPs, CSPs and chemosensory receptors in A. gifuensis have been widely predicted based on transcriptome data (Kang et al., 2017; Fan et al., 2018). However, there is still a paucity of information on the expression profiles of odorant binding proteins in various sensory organs of A. gifuensis. Sequence identification is critical for further functional studies, not to mention the mechanisms of host foraging and mating behavior which are completely unknown.

In the present study, we performed gene prediction, identification, expression profiling of AgifOBPs and further performed a ligand competitive binding test on their recombinant proteins expressed in a prokaryotic expression system to discover two leg-specifically expressed OBPs (AgifOBP7 and AgifOBP9) in A. gifuensis as follows: 1) used A. gifuensis antennal transcriptome to predict AgifOBPs; 2) we identified AgifOBPs and profiled their spatial expression patterns among tissues and organs of both sexes; and 3) we revealed a partial mechanism of olfactory perception based on the ligand competitive binding test.

MATERIALS AND METHODS

Insect Rearing and Tissue Collection

The laboratory population of Aphidius gifuensis was the same as that previously described by Fan et al., 2018. The mummies were collected and placed separately in Petri dishes (3.5 cm in diameter). Newly emerged (within 0–12 h) Aphidius were transferred to larger Petri dishes (9 cm in diameter and 2 cm in height) for another 24 h, and the two groups were divided by sex. Cotton balls dipped in a 25% aqueous solution of sucrose were constantly supplied as the diet for adult wasps. Approximately 500 pairs of antennae from each sex were collected for RNA sequencing. In total, for each replication of qRT-PCR, 100 antennae, 50 heads, 50 thorax, 50 abdomens, and
300 legs were collected. Three replicates were conducted for sampling. The dissected tissues were immediately frozen in liquid nitrogen and stored at −80°C.

**Total RNA Extraction and Synthesis of the First Chain of cDNA**

Total RNA was extracted using TRIzol reagent and combined with micro total RNA extraction kit (Tianmo, Beijing, China) following the manufacturer’s instructions. The frozen tissues were homogenized with a liquid nitrogen cooled mortar and ground with a pestle into very fine dust. Homogenized tissues were treated with 1 ml of TRIzol reagent (Invitrogen, Carlsbad, CA, United States). RNA degradation and contamination were monitored on 2% agarose gels. RNA purity was checked using a Nanodrop ND-1000 spectrophotometer (NanoDrop products, Wilmington, DE, United States). The RNA concentration was measured using a spectrophotometer RNA Nano 6000 Assay Kit of the Agilent Bioanalyzer 2,100 system (Agilent Technologies, CA, United States). Individual total RNA was isolated and cDNA was synthesized using the TRUEscript RT kit (LANY Science & Technology, Beijing, China) following the manufacturer’s protocol.

**Transcriptome Sequencing, Assembly and Functional Annotation**

A total of 3 μg of RNA sample with standard quality ratios (1.8 < OD260/280 < 2.1) was purified using poly-T oligo-attached magnetic beads. Divalent cations under elevated temperature in NEBNext First Strand Synthesis Reaction Buffer (5x) were used for fragmentation. Single-stranded (ss) cDNA was synthesized using a random hexamer primer, M-MuLV Reverse Transcriptase and DNA Polymerase I and RNase H (NEB, United States). The 3’ ends of the DNA fragments were adenylated and the NEBNext Adaptor was ligated to the fragments for hybridization. The library fragments were purified with the AMPure XP system (Beckman Coulter, Beverly, MA, United States) to size select cDNA fragments ~150–200 bp in length. Then, 3 μl of USER Enzyme (NEB, United States) was used with size-selected, adaptor-ligated cDNA at 37°C for 15 min followed by 5 min at 95°C prior to PCR. PCR was performed with Phusion High-Fidelity DNA polymerase, Universal PCR primers and Index (X) Primer. The products were purified (AMPure XP system), and library quality was assessed using the Agilent Bioanalyzer 2,100 system (Agilent Technologies, CA, United States). Clustering of the index-coded samples was performed on a cBot Cluster Generation System using TruSeq PE Cluster Kit v3-cBot-HS (Illumina, China) according to the manufacturer’s instructions. The library preparations were sequenced on an Illumina HiSeq 2,500 platform and paired-end reads (the sequencing strategy was PE125) were generated after cluster generation. After sequencing, the raw reads were processed to remove low quality and adaptor sequences by ng_qc, and then assembled into unigenes using Trinity r20140413p1 min_kmer_cov:2 and other default parameters (Grabherr et al., 2011). Then the unigenes were annotated using seven databases, including the nonredundant protein sequence (Nr, e-value = 1e−5), nonredundant nucleotide (Nt, e-value = 1e−5), Pfam (e-value = 0.01), Clusters of Orthologous Groups (KOG/COG, e-value = 1e−5), Swiss-Prot (e-value = 1e−5), Kyoto Encyclopedia of Genes and Genomes (KEGG, e-value = 1e−10) and Gene Ontology (GO, e-value = 1e−6) databases.

**OBP Gene Prediction and Identification**

The available sequences of OBPs from Hymenoptera species were used as “query” sequences to identify candidate unigenes that code OBPs in the *A. gifuensis* antennal transcriptome with the TBLASTn program with an e-value threshold of 10−5. The sequences that fit the criteria were considered candidate OBPs. The open reading frames were searched by ORF finder (http://www.ncbi.nlm.nih.gov/orf finder/). The putative N-terminal signal peptides were predicted using the SignalP V4.1 program (http://www.cbs.dtu.dk/services/SignalP-4.1/) following the default parameters. Alignments of amino acid sequences were performed with Clustal Omega (https://www.ebi.ac.uk/Tools/msa/clustalo/) and edited using DNAMAN (Lynnon Biosoft San Ramon, CA, United States) software. According to the DEG results, the mean FPKM values for each gene in the antennae of males and females were then log-transformed ["log2 (FPKM +1)"]. A heat map was generated using TBTools (Chen et al., 2020). A phylogenetic tree was constructed by MEGA11 using the maximum likelihood method with a LG + mode to analyze the relationship of OBPs among species and reveal clues of their function (Tamura et al., 2021). Values indicated at the nodes are bootstrap values based on 1,000 replicates presented with 95% cutoff. The orthologous protein sequences from the genomes and transcriptomes of the following Hymenoptera species were used in the analysis: *Apis mellifera* (Forêt and Maleszka, 2006); *M. mediator* (Zhang et al., 2009; Peng et al., 2017); *M. palmaricorin*s (Sheng et al., 2017) and *Aulacocentrum confusum* (Li et al., 2021). The amino acids of the sequences used are listed in [Supplementary Material S1](#). A circular phylogenetic tree was then generated and taxonomically color-coded using the online tool iTOL (https://itol.embl.de/iitol.cgi). To identify the sequences of all candidate AgiOBPs, gene-specific primers ([Supplementary Table S3](#)) were designed with Primer 5.0 (http://frodo.wi.mit.edu/primer5/). Polymerase chain reactions were conducted on an Eppendorf Mastercycler® gradient PCR machine using 2xFastPfu PCR SuperMix (Trans, Beijing, China) and antennal cDNA as a template. An initial denaturation step at 95°C for 5 min followed by 35 cycles of 95°C for 35 s, 58°C as a melting temperature for 35 s, and 72°C for 45 s, and a final extension at 72°C for 10 min. The PCR products were electrophoresed on 2% agarose gels and stained with ethidium bromide to ensure that the correct products were amplified. All targeted PCR products were purified using the AxyPrep PCR clean up Kit (CORING, Jiangshu, China), and then cloned into the pEASY Blunt clone vector (Trans, Beijing, China). After transformation of *Escherichia coli* DH5α competent cells with the ligation products, positive colonies were selected by PCR using the plasmid primers M13 F and M13 R and...
sequenced at San Bo Biotech (Beijing, China). Individual clones confirmed to contain the desired sequence were incubated in LB/ampicillin medium.

Spatial Expression Pattern of AgifOBPs
To explore the expression characteristics of the AgifOBPs, RT-qPCR with an ABI 7500 real-time PCR system (Applied Biosystems Fosters City CA, United States) was conducted with cDNAs prepared from each tissue of male and female *Aphidius*. Briefly, 0.6 µl of both forward primer (10 µmol/L) and reverse primer (10 µmol/L) (Supplementary Table S4) were used in a 20-µl reaction containing 10 µl of 2x SuperReal PreMix Plus, 2 µl of cDNA (from 250 ng of total RNA), 0.4 µl of 50x ROX reference dye, and 6.4 µl of ribonuclease-free ddH_2_O following the instructions provided with the SuperReal PreMix Plus (SYBR Green) kit (FP205) (Tiangen, Beijing, China). The PCR program was as follows: initial 15-min step at 95°C, 40 cycles of (Tiangen, Beijing, China). The PCR program was as follows: initial 15-min step at 95°C, 40 cycles of denaturation at 95°C for 15 s, annealing at 60°C for 32 s and elongation at 72°C for 1 min and finally a 10-min step at 72°C. For melting curve analysis, a dissociation step cycle was added automatically. The amplification efficiency was calculated using the equation: E = [10^(-1/slope)-1] ×100%, in which the slope was derived by plotting the curve threshold (Ct) value against five 2-fold serial dilutions. Only primers with 95–105% amplification efficiencies were used for subsequent data analysis. Relative quantification was performed according to the 2^ΔΔCt method (Livak and Schmittgen, 2001). β-Actin and (nicotinamid adenine dinucleotide) NADH were used as reference genes to normalize the data. All qRT-PCR analyses were performed in three technical and biological replications.

Prokaryotic Expression and Purification of AgifOBP7 and AgifOBP9
The prokaryotic expression and purification procedures were consistent with previous studies (Prestwich, 1993; Wang et al., 2021a). Gene-specific primers were designed to clone the full-length cDNAs encoding mature AgifOBP proteins. The PCR products were first cloned into the pEASY-T1 clone vector (TransGen Biotech, Beijing, China), and then excised and subcloned into the bacterial expression vector pET28a (+) (Novagen, Madison, WI) between the Nde I and EcoR I restriction sites, and reconstructed plasmids were verified by sequencing. The recombinant AgifOBP7 and AgifOBP9 in the present study contain no histidine-tagged peptide at the N-terminus.

Protein expression was induced by adding isopropyl-1-thio-b-D-galactopyranoside (IPTG) to a final concentration of 1 mM when the culture reached an OD_600 value of 0.6. Cells were incubated for an additional 12 h at 28°C and then harvested by centrifugation and sonicated at a low temperature (ice-water mixture). After centrifugation, the bands obtained were checked by 15% SDS–PAGE for their correspondence to the predicted molecular masses of the proteins. They were solubilized according to protocols for the effective rebuilding of the recombinant OBPs in their active forms (Prestwich, 1993). The soluble proteins were then purified by anion-exchange chromatography with RESOURCE Q15 HP column (GE HEALTH CARE, United States) and gel filtration [Superdex 75 10/300 GL column (GE HEALTH CARE, United States)], The crude extracts were passed over a pre-equilibrated RESOURCE Q15 HP column (20 mM Tris–HCl, pH 8.5), and then washed and eluted with Buffer B (20 mM Tris–HCl, 1 M NaCl, pH 8.5). And finally with two rounds of gel filtration through a Superdex 75 10/300 GL column, those eluted proteins were collected and analyzed by 15% SDS-PAGE, and then, several successive dialyzes were performed: 1) at 4°C for 3 h, against 2 L of storage buffer (20 mM Tris–HCl, pH 8.5), 2) at 4°C for 3 h, against 2 L storage buffer (20 mM Tris–HCl, pH 8.5), and 3) at 4°C against 2 L of storage buffer (20 mM Tris–HCl, pH 8.5) overnight. Finally, the desalted protein samples were ultra-centrifuged for 30 min using 3-kDa ultrafiltration at 4°C, and 5,000 rpm. Protein samples were analyzed by SDS-PAGE after every purification step, the concentration of purified protein was determined by a Protein Assay kit (Qubit™ Protein Assay kit, Q33211, Invitrogen), and the purified AgifOBPs were analyzed by mass spectrometry (LC–MS). The purity and concentration of the soluble proteins were evaluated using SDS–PAGE. Finally, stock solutions of AgifOBP7 and AgifOBP9 were collected and kept at −20°C in Tris–HCl (50 mM, pH 7.4).

Fluorescence Competitive Binding Assays
To investigate the ligand-binding property of two AgifOBPs, five groups of competitive ligands were used: 1) aphid alarm pheromone components, including EBF, (-)-α-pinene, (-)-β-pinene and (+)-limonene which are released by other aphids following natural enemy predation or physical damage (Francis et al., 2005; Song et al., 2021b), 2) main components of the aphid sex pheromone: (4aSR,7SR,7aRS)-Nepetalactone; 3) green leaf volatiles of wheat: (Z)-3-hexen-1-ol; 4) aphid-induced plant volatiles (methyl salicylate, and 6-methyl-5-hepten-2-one); and 5) an EBF derivative artificial chemical, namely CAU-II-11, ((E)-3,7-dimethylocta-2,6-dien-1-yl-2-hydroxy-3-methoxybenzoate), which showed a high affinity for aphid EBF- binding proteins (OBP3/7/9, Qin et al., 2020), and was used to investigate the binding properties of purified AgifOBP7 and AgifOBP9. The classes, CAS numbers and purity of the chemicals used in this study are listed in Table 1.

Fluorescence intensity was recorded in a right angle configuration on a Lengguang 970CRT spectrofluorimeter (Shanghai Jingmi, China) at room temperature using a 1 cm light path fluorimeter quartz cuvette. A slit width of 10 nm was selected for both excitation and emission. The measured fluorescence intensities were corrected for both blank signals due to protein emission and scattered excitation light. The spectral data were processed using the software 970CRT 2.0L. Fluorescence binding experiments were conducted in 50 µM Tris–HCl buffer, pH 7.4, at room temperature. The binding affinity for N-phenyl-1-naphthylamine (1-NPN) was determined by adding aliquots of a 1 mM stock solution of 1-NPN dissolved in
spectrophotometric grade methanol into a 2 μM protein sample. The fluorescence of 1-NPN was excited at 337 nm, and emission was recorded between 350 and 500 nm. Spectra were recorded with a high-speed scan. All ligands used in competitive experiments were dissolved in spectrophotometric grade methanol. In competition assays, aliquots of the competing ligands were added into a 2 μM protein solution in the presence of a given concentration of 1-NPN. To estimate the binding affinities of each AgiOBP for a variety of different ligands, we monitored the decrease in 1-NPN fluorescence due to the ability of different odorants to displace 1-NPN and determined the Ki value for each compound. To determine the dissociation constants, the intensity values corresponding to the maximum fluorescence emissions were plotted against the cumulative 1-NPN concentration. The amount of bound ligand was calculated from the fluorescence intensity values by assuming that the protein was 100% active, with a stoichiometry of 1:1 protein: ligand at saturation. The curves were linearized using Scatchard plots. The value of K1-NPN was estimated on a nonlinear plot by a direct plot with an equation corresponding to a single binding site using Prism 7 (GraphPad Software, Inc., United States), and the IC50 was defined as the concentration of a competitor that caused a 50% reduction in fluorescence intensity. The dissociation constants of the inhibitors (Ki) were calculated according to the formula \( Ki = \frac{IC50}{1+([1-NPN]/K1-NPN)} \), in which [1-NPN] represents the free 1-NPN concentration and K1-NPN represents the dissociation constant for AgiOBPs/1-NPN (Ban et al., 2003; Zhong et al., 2012; Sun et al., 2016; Fan et al., 2017; Qin et al., 2020; Wang et al., 2021). All fluorescence competitive binding assays were performed in three independent replicates, and Ki dates are present as means ± SD.

### RESULTS

#### Overview of Transcriptomes

A total of 2.22 and 2.30 million raw reads were obtained from *A. gifuensis* antennae libraries from females and males, respectively. The data presented in the study that was deposited in the Sequence Read Archive (SRA) repository, the accession number are SRR18251541 and SRR18251542. After removal of low-quality, adaptor, and contaminating sequences, 3.31 and 3.03 million clean reads were retained (Supplementary Table S1) and assembled into 81,235 distinct transcripts (mean length = 661 bp) and 65,854 unigenes (mean length = 568 bp). The length distribution presented in (Supplementary Figure S1). In total, 18,408 (27.95%) of all 65,854 unigenes, 5,625 (8.54%), 7,551 (40.92%), 12,484 (18.95%), 15,070 (22.88%), 15,951 (24.22%) and 9,462 (14.36%) transcripts from *A. gifuensis* antennae were annotated using the Nr, Nt, KO, Swiss-Prot, Pfam, GO and KOG databases, respectively (Supplementary Table S2). The most abundant GO terms were biological process terms, with AgiOBP3 corresponding to the cellular process and AgiOBP15 grouped with the membrane. The cluster for cellular process was the second largest group. Most transcripts that corresponded to molecular function were related to binding and catalytic activity (Supplementary Figure S2). In the KOG classification, unigenes clustered into 26 categories (Supplementary Figure S3). Among these categories, general function prediction was the dominant category, followed by signal transduction and posttranslational modification, protein turnover and chaperon. All the unigenes annotated in the KO database were assigned to the 5 biological pathways described in the KEGG database: cellular processes, environmental information processing, genetic information processing, metabolism, and organismal systems (Supplementary Figure S4). The most common pathway was metabolism followed by genetic information processing, organismal systems and cellular processes. Signal transduction was involved in 940 genes in the environmental information processing group.

#### Statistical Analyses

For qRT-PCR analyses, the differences between means of biological replicates were tested using two-way ANOVA followed by multiple comparisons tests regardless of rows and columns using GraphPad Prism version 7.0.0 for Windows (GraphPad, Software, San Diego, California United States, www.graphpad.com). Differences between means for experiments with more than two treatments were distinguished using Tukey’s honestly significant difference (HSD) test at the \( p < 0.05 \) significance level.

### OBPs in *Aphidius gifuensis*

| No | Code | CAS        | Purity | IC50     | Ki(μM)          | IC50     | Ki(μM)          |
|----|------|------------|--------|----------|----------------|----------|----------------|
| 1  | (E)-Farnesene | 18,794–84–8 | ≥85%   | >30      | 20.30 ± 1.99   | 17.83 ± 1.69 | 4.60 ± 0.43   |
| 2  | (-)-α-Pine| 80–56–8    | ≥95%   | >30      | >30            | 24.10 ± 3.42 | 6.22 ± 0.88   |
| 3  | (E)-α-Pine  | 19,902–08–0| ≥99%   | >30      | >30            | 11.86 ± 1.27 | 3.06 ± 0.33   |
| 4  | (+)-Limonene| 138–86–3   | ≥95%   | >30      | >30            | 12.85 ± 0.25 | 3.32 ± 0.06   |
| 5  | (4aSR 7SR 7aRS)-Nepetalactone | 21,651–62–7 | ≥80%  | >30      | 16.12 ± 3.49   | 9.03 ± 0.31  | 2.33 ± 0.08   |
| 6  | 6-Methyl-5-hepten-2-one | 110–93–0 | ≥99%   | >30    | >30            | 22.32 ± 3.38 | 5.76 ± 0.87   |
| 7  | cis-3-Hexenol | 926–96–1  | ≥97%   | >30      | >30            | 17.01 ± 0.33 | 4.39 ± 0.09   |
| 8  | Methyl salicylate | 119–26–8 | ≥99%   | >30    | >30            | 13.31 ± 2.99 | 3.43 ± 0.77   |
| 9  | CAU-II-11 | -         | ≥98%   | 18.56 ± 1.73 | 8.50 ± 0.73   | 4.15 ± 0.33  | 1.07 ± 0.08   |
Figure 1 illustrates that OBP5, OBP6, OBP11, and OBP15 were highly expressed genes in both sex antennae but OBP3/14/17 showed relatively low expression levels. The phylogenetic tree of Hymenoptera OBPs was built using MEGA11 (maximum likelihood method with an LG model) OBP sequences from 5 different species (A. gifuensis, A. mellifera, M. mediator, M. pulchricornis and A. confusum). A. gifuensis OBPs are clustered together to form three homologous subgroups (lineages). Among them, OBP1, OBP5, OBP7, OBP9 and OBP17 were in one subgroup, OBP2, OBP3, OBP6, OBP8 and OBP11-OBP15 were in the other subgroup, and OBP4 fell into the third subgroup. The results showed that AgiOBPs almost spread across in clades without species specificity (Figure 2). Among these AgiOBPs, AgiOBP4 was found in the MpuOBP4 clade. AgiOBP6 exhibited a rather high similarity to other orthologs such as AmelOBP6, MmedOBP6 and MpuOBP6. AgiOBP8 also showed a high similarity to MmedOBP8 and MpuOBP8 (Figure 2).

Spatial Expression Pattern of AgiOBPs
Compared to other tissues or organs, 8 of the 14 OBPs, namely, AgiOBP3, AgiOBP5, AgiOBP6, AgiOBP7, AgiOBP8, AgiOBP11, AgiOBP12 and AgiOBP15, maintained higher expression in antennae (Figures 3, 4; p < 0.05). AgiOBP17 was highly expressed in the head. AgiOBP1/2/7/9 were expressed in legs with significantly higher expression levels (Figures 3, 4, p < 0.05). The other two OBPs, AgiOBP4 and AgiOBP13 were widely expressed among tissues and organs. Specifically, AgiOBP3/5/6/11/12/15 were specifically expressed in antennae. Among them, the expression levels of AgiOBP3/11 were even higher in male antennae. However, AgiOBP12/15 were even higher in female antennae, and AgiOBP6 showed no difference in antennae of both sexes. Moreover, AgiOBP1/2/4/5/7/9/15 showed relatively higher expression levels in legs (Figures 3, 4). Among them, AgiOBP2 was specifically expressed in female legs. And AgiOBP7/9/15 were expressed at comparatively higher levels in female legs. In contrast, AgiOBP1/4/5 were expressed at higher levels in male legs. Notably AgiOBP7 was female specific and was expressed directly in female antennae and legs. In males, AgiOBP8 was specifically expressed in antennae. In females, it was relatively highly expressed in both the antennae and abdomen. We also found that the highest level of AgiOBP17 was in the heads of both sexes. Although both AgiOBP4 and AgiOBP13 were widely expressed, AgiOBP4 showed an even higher expression level in thoraxes of both females and males. AgiOBP13 expression was significantly higher in the male abdomen. In addition, AgiOBP1 and AgiOBP9 were specifically or highly expressed in the legs of both male and female A. gifuensis. AgiOBP2 was significantly expressed in the legs of females (Figure 4, p < 0.05). In summary, AgiOBP3/5/6/11/12/15 were antennal specifically expressed OBPs. AgiOBP2/9 were specifically expressed OBPs in legs. AgiOBP17 is an OBP specifically in the head (Figure 3, p < 0.05). In addition, AgiOBP5/7 were female specific OBPs. AgiOBP8 expression was significantly higher in the antennae of males and in both the antennae and abdomen of females (Figure 4, p < 0.05).

Expression and Purification of AgiOBP7 and AgiOBP9
AgiOBP7 and AgiOBP9 were successfully expressed in the inclusion bodies using a bacterial system. After a dissolving and refolding treatment, the refolded AgiOBP7 and AgiOBP9 were purified with yields of 0.25 mg/ml as soluble proteins (Figures 5A, B). More than 15 mg of purified AgiOBP7 and AgiOBP9 was obtained using RESOURCE Q15 affinity columns, with the His-tag removed. The theoretical molecular weight values for AgiOBP7 and AgiOBP9 were very close to the measured values (AgiOBP7, 13.401 kDa; AgiOBP9, 12.498 kDa). The purified protein samples were further identified by LC–MS/MS (data not shown).

Fluorescence Competitive Binding Assays
To investigate the role of two OBPs in the odor perception of aphids, we chose alarm pheromones (Eββ; (−)-α-pinene, (−)-β-pinene, (+)-limonene, Eβ derivative (CAU-II-11), aphid sexual pheromones (4aSR 7SR 7aRS)-nepetalactone as well as volatiles of...
wheat green leaf (cis-3-hexen-1-ol) and aphid induced plant main volatiles (methyl salicylate, 6-methyl-5-hepten-2-one) as the candidate ligands for fluorescence competitive binding assays (Table 1). We first tested the binding affinities of both OBPs to the fluorescent probe N-phenyl-1-naphthylamine (1-NPN) as previously reported (Qiao et al., 2009; Fan et al., 2017). The dissociation constants of AgifOBP7/1-NPN and the AgifOBP9/1-NPN complex were 1.69 ± 0.27 µM and 0.69 ± 0.24 µM respectively (Figures 5C,D).

In a subsequent experiment, we used a fluorescence competitive binding assay to determine the binding affinities of AgifOBP7 and AgifOBP9 to different odorants. Based on the binding curves, we calculated the median inhibitory concentration (IC50) and dissociation constant (Ki) values (Table 1). Among the tested odorants, EBF, (-)-α-pinene, (−)-β-pinene, (−)-limonene, (4aSR 7SR 7aRS)-nepetalactone, cis-3-hexen-1-ol, methyl salicylate and 6-methyl-5-hepten-2-one displayed relatively high binding affinities (Ki < 10 µM) to AgifOBP9 (Figure 5F). Interestingly, among all the tested odorants, CAU-II-11 bound most strongly (Ki = 1.07 ± 0.08 µM) to AgifOBP9 (Table 1), which is the derivative of EBF (Ki = 4.60 ± 0.43 µM). However, this was not the case with AgifOBP7, which only displayed weak binding with EBF (Ki = 20.30 ± 1.99) and (4aSR 7SR 7aRS)-nepetalactone (Ki = 16.12 ± 3.49), not much with (−)-α-pinene, (−)-β-pinene, (−)-limonene, cis-3-hexen-1-ol, methyl salicylate and 6-methyl-5-hepten-2-one. CAU-II-11, like AgifOBP9, showed the strongest binding affinity (Ki = 1.07 ± 0.08 µM) to AgifOBP7 among the examined odorants (Figure 5E; Table 1).

DISCUSSION

Odorant-binding proteins are classically defined as olfactory soluble proteins (Vogt, R. G., & Riddiford, L. M, 1981; Pelosi, 2006) and play an essential role in habitat searching and finding...
suitable mates. With the increase in insect genome projects and transcriptome sequencing projects, large numbers of OBPs have recently been identified in different insect species. In the present study, we constructed a cDNA library from the antennae of the endoparasitoid *A. gifuensis* for transcriptome sequencing and categorized the potential function of the odorant binding protein genes by bioinformatics approaches.

**FIGURE 3** The relative expression patterns of different *AgfOBP* genes in different tissues of males and females as measured by quantitative real-time polymerase chain reaction. The fold changes are relative to the transcript levels of OBP13 in the male thorax. The NADH and ACTIN genes were used as references to normalize the expression of each tested gene. The data are presented as the mean ± SD. The asterisk * and ** above the bars indicate significant differences at \( p < 0.05 \); and \( p < 0.01 \), respectively, according to two-way ANOVA.

**OBP Prediction, Cloning and Phylogenetic Analysis**

Fourteen OBPs in the *A. gifuensis* antennae transcriptome were identified in the present study. This number is similar to those in *A. mellifera* (Forêt and Maleszka, 2006), *M. mediator* (Zhang et al., 2009; Peng et al., 2017), *Meteorus pulchricornis* (Sheng et al., 2017), *Cotesia vestalis* (Zhou et al., 2021) and *Aulacocentrum*
confusum (Li et al., 2021), therefore indicating that there are similar OBP numbers in Hymenoptera insects.

The phylogenetic tree of these AgifOBPs, together with OBPs from 4 hymenopteran species, showed that the AgifOBPs segregate into the orthologous clades of the other species, rather than into A. gifuensis paralogous clades. AgifOBP4 was found in the MpuLOBP4 clade, whereas AconOBP4, MmdOBP4, and AmelOBP4 were clustered in the other one clade. AgifOBP6 and AgifOBP8 were present in the three wasps of A. gifuensis, M. mediator and M. pulchricornis, but their orthologs were rarely found in Apis mellifera (Figure 2). This also suggests that these AgifOBPs might play different roles in odor recognition or have roles other than olfaction. The comparatively conserved OBPs in hymenoptera wasps, particularly in parasitoid wasps implied that their function could be limited to the common olfactory physiology of these insects. Some study results on natural enemies of aphids support this hypothesis. For example, aphid OBP7 orthologs have been widely reported to have their affinity with the alarm pheromone EBF (Sun et al., 2012a; Zhong et al., 2012; Fan et al., 2017; Qin et al., 2020). CpalOBP10 in lacewing Chrysopa pallens, an aphid predator, belongs to the same lineage as aphid OBP7 such as in S. avenae and in A. pisum, and its affinity for EBF was also consistent with that of aphid OBP7 orthologs (Li et al., 2017; Li et al., 2019).

**Spatial Expression Pattern**

The spatial expression profile of AgifOBPs was verified using qPCR. Our data revealed that five OBPs, namely AgifOBP3, AgifOBP5, AgifOBP6, AgifOBP11, and AgifOBP15, were expressed at a high level in the antennae (Figures 3, 4), while four OBPs, AgifOBP2, AgifOBP4, AgifOBP7 and AgifOBP8, were expressed at a medium level, and seven OBPs, AgifOBP1, AgifOBP9, AgifOBP12/13, and AgifOBP17, were expressed at a low level in the antennae (Figures 3, 4). The antennal specific OBPs (Figure 4) suggest their function of recognizing and binding odorants from the environment. Six OBPs, AgifOBP2, AgifOBP4, AgifOBP5, AgifOBP7, AgifOBP13 and AgifOBP17, showed expression patterns among sensory and nonsensory organs, indicating their possible multiple functions in olfactory perception as well as other physiological processes such as development and reproduction. Both AgifOBP1 and AgifOBP9 showed higher expression levels in the legs than the other four tissues (antennae, heads, thorax and abdomen), which could be related to the adaptation of A. gifuensis during migration as we

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**FIGURE 4 |** Relative expression of AgifOBP genes in the different tissues of male and female A. gifuensis as measured by quantitative real-time polymerase chain reaction. Relative fold changes were normalized to the transcript levels in the male thorax. The standard error is represented by the error bar (n = 3) and the different lowercase letters (a, b, c, d, e) indicate significant differences in transcript abundances (two-way ANOVA followed by Tukey’s HSD multiple comparison test, p < 0.05).
have discussed in our previous study (Xue et al., 2016), and might be involved in the procedure of taste or volatile perception or be related to olfactory sensilla on the legs (Yasukawa et al., 2010; Harada et al., 2012). A similar condition was also found for AgifOBP5, which is expressed in small amounts in the head and leg, in addition to being expressed abundantly in antennae. Apart from antennae, alternatively, these OBPs expressed in other tissues may be responsible for corresponding functions. For example, NlugOBP3 is highly expressed in the abdomen of Nilaparvata lugens and may be involved in juvenile hormone transport and play an important role in metamorphosis (He et al., 2011). Insect OBPs have been reported to act as carrier proteins in the male reproductive apparatus of mosquitoes (Li et al., 2008). After mating, the OBPs expressed by male moths are found on the surface of fertilized eggs, which helps the larvae to avoid cannibalistic behaviors (Sun et al., 2012b). For parasitic wasps, AconOBP8 was reported to be expressed predominantly in the abdomen (Li et al., 2021). Similar expression patterns of OBPs in the nonolfactory tissues were observed in Sclerodermus sp. (Zhou et al., 2015) and M. pulchricornis (Sheng et al., 2017). In our present study, qPCR analysis revealed that AgifOBP8 was also expressed in the female abdomen, and it can be speculated that OBP8 may potentially function as a pheromone-binding protein for identifying a particular signal such as the sex pheromone component in mating or oviposition behaviors, although the active component of sex pheromone in this species is still unclear.

The results obtained by qPCR are consistent with antennal transcriptome based differential expression analysis (heatmap, Figure 1). Nonetheless, any discrepancy between qPCR and differential expression analysis results illustrate the poor performance of showing local details by omics big data analysis.

**Ligand-Binding Properties**

Table 1 indicates that the proteins AgifOBP7 and AgifOBP9 have broad binding activities across the aphid alarm pheromone components, aphid sex pheromone, green leaf volatiles, aphid-induced plant volatiles and EBF derivatives. AgifOBP9 showed higher binding activities than AgifOBP7 with all the five types of compounds. Similar results have been found in its prey aphid A. pisum, in which ApisOBP9 also exhibited higher affinities with all the compounds than ApisOBP7 (Qin et al., 2020), although there is no evolutionary homology between the two species. For AgifOBP7 and AgifOBP9, EBF derivatives had higher binding properties than the lead EBF and other compounds. These results are consistent with studies on the characterized OBPs of ApisOBP1, ApisOBP3, and ApisOBP6-OBP10 in A. pisum (Sun et al., 2012a; Qin et al., 2020). Both proteins show preferential binding to several related compounds. AgifOBP7 bound the above five types of compounds from strong to...
As the natural enemy of aphids, *A. gifuensis* locates aphids using the cues of aphid pheromones and plant volatiles (Powell et al., 1998; Kang et al., 2017). Our results indicate that AgiOBP7 is specific to aphid pheromone components and EBF derivatives, and that AgiOBP9 has a broad spectrum of binding to compounds. Other OBPs in aphid natural enemies also bind to aphid pheromone components and plant volatiles. For example, OBP3, OBP4, OBP6, OBP7, OBP9, and OBP10 in *Chrysopa pallens* bind plant volatiles and aphid alarm pheromone EBF, and, OBP10 specifically binds EBF (Li, et al., 2017; Li et al., 2019). The new OBPs from the aphid natural enemy *Eupeodes corollae*, OBP12, OBP15 and OBP16 also bound with EBF and plant volatiles, among which OBP12 and OBP15 strongly bound EBF (Wang et al., 2022).

Many natural enemies such as *Aphidius ervi*, *Aphidius uzbekistanicus*, and *Adalia bipunctata* are attracted to EBF (Buitenhus et al., 2004). To confirm the functions suggested by the phylogenetic tree and tissue expression profiles, AgiOBP7 and AgiOBP9 were selected to perform a potential functional study. Overall, the odorants exhibited relatively high binding affinities (Ki < 10 μM) to AgiOBP9 (Figure 5F; Table 1). Interestingly, among all the tested odorants, CAU-II-11 had the strongest binding affinity (Ki = 1.07 ± 0.08 μM) to AgiOBP9 (Table 1), which is the derivative of EBF (Ki = 4.60 ± 0.43 μM). This finding is in line with prior research on ApisOBP3/7/9 using CAU-II-11 (Qin et al., 2020). This result further supports that both aphid-induced volatiles as well as EBF are used by *A. gifuensis* in aphid location and that AgiOBP9 may be involved in this process.

In summary, we first predicted 15 OBPs based on the antennal transcriptome of both male and female *A. gifuensis*. Fourteen of these OBPs were verified by gene cloning. Furthermore, their detailed spatial expression pattern showed that most OBPs are mainly expressed in the sensory organs, but some are widely expressed in various tissues or organs such as the thorax and abdomen. Finally, at least one female particularly expressing OBP (AgiOBP9) showed affinity to EBF in a fluorescence competition experiment, which further indicated the likely molecular basement of sensing the aphid alarm pheromone at the molecular level in *A. gifuensis*. In addition, what cannot be ignored is the presence of OBPs expressed in other nonsensory organs such as the abdomen, which supports the existence of carrier transport functions other than for foreign chemicals and therefore broader ligand ranges of wasp OBPs. Our findings may shed insight into parasitic wasps’ olfactory sensitivity to host hints, as olfactory organs recognize pheromones and odorant substances that influence both host hunting and oviposition activities and will help us better understand parasitic wasp host foraging and mating behaviors, which will aid in the strengthening and better utilization of *A. gifuensis* as a powerful and natural biocontrol strategy. As a result, we anticipate that additional research into the aforementioned topics will improve the efficacy of parasitoid-based biological control approaches against aphid pests.

**DATA AVAILABILITY STATEMENT**

The data presented in the study was deposited in the Sequence Read Archive (SRA) repository, the accession number are SRR18251541 and SRR18251542.

**AUTHOR CONTRIBUTIONS**

XJ and JF collected the insect organs or tissues, analyzed the transcriptome data, performed the molecular experiments such as RNA extraction, gene cloning and qPCR, and wrote the draft. JF and YQ assembled the RNA seq data. XJ and YX established the insect laboratory population. FF discussed the data. JF conceived and designed the study, and performed the personalized bioinformatic analysis. JC organized and directed the project. All authors contributed to the manuscript revision.

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**SUPPLEMENTARY MATERIAL**

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