

Molecular identification and expression patterns of odorant binding protein and chemosensory protein genes in *Athetis lepigone* (Lepidoptera: Noctuidae)

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The olfaction system of insects plays an important role in mediating various physiological behaviors, including locating hosts, avoiding predators, and recognizing mates and oviposition sites. Therefore, some key genes in the system present valuable opportunities as targets for developing novel green pesticides. *Athetis lepigone*, a noctuid moth can feed on more than 30 different host plants making it a serious polyphagous pest worldwide, and it has become one of the major maize pests in northern China since 2011. However, there are no reports on effective and environmentally friendly pesticides for the control of this pest. In this study, we identified 28 genes encoding putative odorant binding proteins (OBPs) and 20 chemosensory protein (CSPs) genes based on our previous *A. lepigone* transcriptomic data. A tissue expression investigation and phylogenetic analysis were conducted in an effort to postulate the functions of these genes. Our results show that nearly half (46.4%) of the *AIOBPs* exhibited antennae-biased expression while many of the *AICSPs* were highly abundant in non-antennal tissues. These results will aid in exploring the chemosensory mechanisms of *A. lepigone* and developing environmentally friendly pesticides against this pest in the future.

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Running head: OBP and CSP genes of *Athetis lepigone*

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Abstract

The olfaction system of insects plays an important role in mediating various physiological behaviors, including locating hosts, avoiding predators, and recognizing mates and oviposition sites. Therefore, some key genes in the system present valuable opportunities as targets for developing novel green pesticides. *Athetis lepigone*, a noctuid moth can feed on more than 30 different host plants making it a serious polyphagous pest worldwide, and it has become one of the major maize pests in northern China since 2011. However, there are no reports on effective and environmentally friendly pesticides for the control of this pest. In this study, we identified 28 genes encoding putative odorant binding proteins (OBPs) and 20 chemosensory protein (CSPs) genes based on our previous *A. lepigone* transcriptomic data. A tissue expression investigation and phylogenetic analysis were conducted in an effort to postulate the functions of these genes. Our results show that nearly half (46.4%) of the *AlOBPs* exhibited antennae-biased expression while many of the *AlCSPs* were highly abundant in non-antennal tissues. These results will aid in exploring the chemosensory mechanisms of *A. lepigone* and developing environmentally friendly pesticides against this pest in the future.

INTRODUCTION

The olfaction system of insects mediates a host of physiological behaviors, such as host location, predator avoidance, and mate and oviposition site recognition (Leal 2013). Many studies show that the periphery process of insect olfaction requires a set of genes, including those that encode odorant binding proteins (OBPs), chemosensory proteins (CSPs), and chemosensory receptors (Elfekih et al. 2016; Glaser et al. 2015; Larter et al. 2016; Li et al. 2015; Paula et al. 2016; Zhang et al. 2013). Generally, OBPs/CSPs located in the antennal sensillar lymph can recognize and bind external odorants that can then be transferred by OBPs/CSPs through the sensillar lymph to chemosensory receptors, odorant receptors (ORs) and ionotropic receptors (IRs). Therefore, OBPs and CSPs play key roles in helping insects recognize various odorants and regulate their behaviors (Dani et al. 2011; Zhou 2010). These functions also suggest that these protein families may present valuable opportunities as target genes for developing novel green pesticides.

Insect OBPs are a class of small, abundant, and water-soluble extracellular proteins of ~14 KDa. Most OBPs use six positionally conserved cysteines to form three interlocking disulphide bridges that stabilize the protein's three-dimensional structure (Lagarde et al. 2011; Leal et al. 1999; Pelosi & Maida 1995; Vogt & Riddiford 1981). Since the first OBP was identified in *Antheraea polyphemus* (Vogt & Riddiford 1981), many OBPs have been found in various insects based on genomic or transcriptomic methods in recent years. Based on the structural features and similarity in protein sequences, insect OBPs can be divided into three major subclasses (Li et al. 2013; Schultze et al. 2012; Spinelli et al. 2012; Zhou 2010): Classic OBPs, including pheromone binding proteins (PBPs), general odorant binding proteins (GOBPs), and two OBPs involved in the recognition of female sex pheromones and host volatiles; plus-C OBPs; and minus-C OBPs, which may also participate in the binding of host volatiles as suggested by an *in vitro* competitive binding assay.

Olfactory specific protein D (OS-D), the first insect CSP gene, was discovered in *Drosophila melanogaster* (McKenna et al. 1994). By using similar methods as for OBP identification, many CSPs have been discovered in distinct insects (Guo et al. 2011; Iovinella et al. 2013; Jacquin-

Joly et al. 2001; Liu et al. 2010; Missbach et al. 2015; Picimbon et al. 2001; Wanner et al. 2004). Unlike OBPs, CSPs are smaller and more conserved in distinct insects, which only have four conserved cysteines that form two interlocking disulphide bridges (Bohbot et al. 1998; Lartigue et al. 2002; Maleszka & Stange 1997; Pelosi et al. 2005; Zhang et al. 2014). Furthermore, OBPs are usually specifically or predominately expressed in the antennae, whereas many CSPs are expressed in the antennae and other tissues (Pelosi et al. 2005; Vogt 2005; Zhang et al. 2016a; Zhang et al. 2013), suggesting insect CSPs have both chemosensation and non-chemosensation functions as is illustrated by their association with chemosensation in moths (Jacquin-Joly et al. 2001; Sun et al. 2015; Zhang et al. 2014), limb regeneration in *Periplaneta eparata* (Nomura et al. 1992), embryo development in *Apis mellifera* (Maleszka et al. 2007), behavioral phase change in *Locusta migratoria* (Guo et al. 2011), and female moth survival and reproduction in *Spodoptera exigua* (Gong et al. 2012).

Athetis lepigone Möschler (Lepidoptera: Noctuidae) is a serious polyphagous pest found worldwide (Fu et al. 2014; Karsholt et al. 2013; Lindeborg 2008; Nikolaevitch & Vjatcheslavovna 2003; Zhang et al. 2009) that can feed on more than 30 different host plants species and has become one of the major maize pests in northern China since 2011 (Jiang et al. 2011; Ma et al. 2012; Shi et al. 2011). However, there are no reports on the chemosensory mechanism mediated by OBPs/CSPs between the pests and host plants. In this study, we identified 28 and 20 genes encoding putative AIOBPs (*A. lepigone* OBPs) and AICSPs (*A. lepigone* CSPs), respectively, based on our previous transcriptomic data of *A. lepigone* (Zhang et al. 2016b). Tissue expression and phylogenetic analyses were conducted in an effort to postulate the function of these genes. We found that most AIOBPs and AICSPs had high identities with those in other moths (Campanacci et al. 2001; Gu et al. 2013; Liu et al. 2015c; Zhang et al. 2015a; Zhang et al. 2013; Zhang et al. 2015b); nearly half of the AIOBPs exhibited antennae-biased expression, and many AICSPs were found in various tissues and were highly expressed in proboscises, legs, and wings, which will help us explore the chemosensory mechanism of *A. lepigone* and develop environmentally friendly pesticides against this pest in the future.

98

99 MATERIALS & METHODS

100 Insect rearing and tissue collection

101 *A. lepigone* were fed an noctuid artificial diet (Huang et al. 2002) at a temperature of 26 ± 1 °C
 102 in a 14:10 h, light:dark photoperiod. Pupae were sexed, and males and females were placed into
 103 separate enclosures. Adult moths were given a 10% honey solution after emergence. We
 104 collected 25–30 female antennae (FA), 25–30 male antennae (MA), 50–60 proboscises (Pr,
 105 ♀:♂=1:1), 10–12 abdomen (Ab, ♀:♂=1:1), 28–30 legs (Le, ♀:♂=1:1), and 28–30 wings (Wi,
 106 ♀:♂=1:1) from three-day-old virgin adults. All samples were immediately frozen in liquid
 107 nitrogen and stored at -80 °C until use.

108

109 RNA isolation and cDNA synthesis

110 Total RNA was extracted using the MiniBEST Universal RNA Extraction Kit (TaKaRa, Dalian,
 111 China) following the manufacturer's instructions, and the RNA quality was checked using a
 112 spectrophotometer (NanoDrop™ 2000, Thermo Fisher Scientific, USA). The single-stranded
 113 cDNA templates were synthesized from 1 µg total RNA from various tissue samples using the
 114 PrimeScript™ RT Master Mix (TaKaRa, Dalian, China).

115

116 Sequence analyses

117 The open reading frames (ORFs) of the putative chemosensory genes were predicted using ORF
 118 Finder (<http://www.ncbi.nlm.nih.gov/gorf/gorf.html>). The similarity searches were performed
 119 with NCBI-BLAST (<http://blast.ncbi.nlm.nih.gov/>). Putative N-terminal signal peptides for
 120 ALOBPs and AICSPs were predicted by SignalP 4.1 (<http://www.cbs.dtu.dk/services/SignalP/>)
 121 (Petersen et al. 2011).

122

123 Phylogenetic analysis

124 Phylogenetic trees were reconstructed for the analysis of *ALOBPs* and *AICSPs*, based on the gene

sequences of *A. lepigone* and those of other insects. The OBP data set contained 28 sequences from *A. lepigone* (Table S1), and 100 from other insects including *Bombyx mori* (Gong et al. 2009), *Manduca sexta* (Grosse-Wilde et al. 2011), *Sesamia inferens* (Zhang et al. 2013), and *Spodoptera littoralis* (Legeai et al. 2011). The CSP dataset contained 20 sequences from *A. lepigone* (Table S1) and 68 from other insects including *B. mori* (Gong et al. 2007), *M. sexta* (Grosse-Wilde et al. 2011), *S. inferens* (Zhang et al. 2013), and *S. littoralis* (Legeai et al. 2011). Amino acid sequences were aligned with MAFFT version 7 (<http://mafft.cbrc.jp/alignment/server/>), and phylogenetic trees were constructed using PhyML (Guindon et al., 2010) based on the LG substitution model (Le and Gascuel, 2008) with Nearest Neighbour Interchange (NNI), and branch support estimated by a Bayesian-like transformation of the aLRT (aBayes) method. Dendrograms were created and colored in FigTree (<http://tree.bio.ed.ac.uk/software/figtree/>).

Quantitative real time-PCR

Expression profiling of *AIOBPs* and *AICSPs* was performed using quantitative real time-PCR (qRT-PCR) performed in a LightCycler® 96 (Roche, Switzerland) with a mixture of 5 µL 2X SYBR® Premix Ex Taq (Tli RNaseH Plus) (TaKaRa, Dalian), 0.2 µL of each primer (10 µM), 2.5 ng of sample cDNA, and 3.6 µL of sterilized ultrapure H₂O. The reaction program was as follows: 30 s at 95°C, 40 cycles of 95°C for 5 s, and 60°C for 20 s. The results were analyzed using a LightCycler® 96 SW 1.1. The qRT-PCR primers (Table S2) were designed with Beacon Designer 7.9 (PREMIER Biosoft International, CA, USA). This was followed by the measurement of fluorescence over a 55 to 95 °C melting curve to detect a single gene-specific peak and to check the absence of primer dimer peaks, and a single and discrete peak was detected for all primers tested. Negative controls consisted of non-template reactions where the cDNA was replaced with H₂O.

Expression levels of *AIOBPs* and *AICSPs* were calculated relative to the reference genes *ALGAPDH* (*A. lepigone* glyceraldehyde-3-phosphate dehydrogenase) and *AIEF* (*A. lepigone*

elongation factor-1 alpha) using the Q-Gene method in the Microsoft Excel-based software Visual Basic (Muller et al. 2002; Simon 2003). For each sample, three biological replicates were performed with three technical replicates per biological replicate.

Statistical analysis

Data (mean \pm SE) from various samples were subjected to one-way nested analysis of variance (ANOVA) followed by a least significant difference test (LSD) for mean comparisons using the SPSS Statistics 22.0 software (SPSS Inc., Chicago, IL, USA).

RESULTS

Identification of putative OBP genes in *A. lepigone*

Based on our previous antennal transcriptomic data (NCBI-SRX number: 2543665) for *A. lepigone* (Zhang et al. 2016b), we first identified 28 genes encoding putative OBPs including three *PBPs* and two *GOBPs* (Table 1). Among the 28 *AlOBPs*, 24 sequences were predicted to be full-length genes that encoded 133 to 246 amino acids; all 24 genes had a predicted signal peptide at the N-terminus. According to the number and position of conserved cysteines, insect OBPs can be divided into different subclasses: Classic OBPs, Plus-C OBPs, and Minus-C OBPs (Zhou 2010). Here, *AlOBP4* and *AlOBP9* had no conserved cysteines at the C2 and C5 positions, and, therefore, belonged to the Minus-C OBP subfamily; *AlOBP2*, *AlOBP7*, and *AlOBP14* had cysteines in addition to the six conserved cysteines; therefore, they belonged to the Plus-C OBP subfamily; the other 19 full-length *AlOBPs* belonged to the Classic OBP subfamily, which had six conserved cysteines at the corresponding positions (Fig. S1).

Identification of putative CSP genes in *A. lepigone*

Twenty putative genes encoding CSPs were identified in *A. lepigone* via antennal transcriptome analysis (Table 2). Eighteen of these had full length ORFs with 4 conserved cysteines in corresponding positions (Fig. S2), and seventeen genes (except *AlCSP14*) had a predicted signal

peptide at the N-terminus. The results of a BLASTX match showed that 80% of these CSPs (n=16) had >70% identity with other CSPs from different moths and that this was higher than the sequence identities of the OBPs (75%) (Table 2).

Phylogenetic analyses of moth OBPs and CSPs

Two phylogenetic trees, one of moth OBPs and one of moth CSPs, were constructed using protein sequences from *A. lepigone*, *B. mori*, *S. inferens*, *S. littoralis*, and *M. sexta*, (Fig. 1 and Fig. 2). Similar to other studies (He et al. 2010; Pelosi et al. 2014b; Vogt et al. 1991; Xiu & Dong 2007), the OBP tree showed that moth OBPs can be divided into PBP/GOBP, Minus-C OBP, and Plus-C OBP subfamilies. ALPBP1-3 clustered into the PBP subfamily and the ALGOBPs (1 and 2) clustered into the GOBP subfamily. Two AIOBPs (AIOBP4 and AIOBP9) clustered into the moth Minus-C OBP subfamily, and four AIOBPs (AIOBP2, AIOBP7, AIOBP10, and AIOBP14) clustered into the moth Plus-C OBP subfamily. The rest of the AIOBPs clustered with at least one orthologous moth gene. In the constructed CSP tree, our results indicated that all 20 AICSPs were distributed along various branches and each clustered with at least one other moth ortholog.

OBPs and CSPs expression patterns in *A. lepigone*

We used the qRT-PCR results to investigate the expression profiles of all AIOBPs and AICSPs. The results showed that all the OBPs and CSPs were expressed in the adult antennae of *A. lepigone*. Among the 28 AIOBPs, 13 AIOBPs (including PBPs and GOBPs) were significantly highly expressed in the antennae ($p < 0.05$, ANOVA, LSD), including 5 male-biased (ALPBP1, ALPBP2, AIOBP6, AIOBP17, and AIOBP20) and 3 female-biased (AIOBP1, AIOBP3, and AIOBP19) OBP genes. In all 28 AIOBPs, ALGOBP1 and ALPBP1 (male antennae) exhibited the highest expression levels, and AIOBP19 exhibited the lowest expression abundance (Fig. 3). In addition, eight AIOBPs (AIOBP4, 8, 11, 14, 16, 21, 22, and 23) exhibited proboscis-biased expression, AIOBP10 was expressed significantly more in the adult abdomen, and four AIOBPs (AIOBP2, 9, 13, and 18) displayed higher expression levels in adult wings than in other tissues

(Fig. 3).

Compared to *AlOBPs*, *AlCSPs* were highly expressed in adult antennae as well as in non-antennae tissues. Of the 20 identified *AlCSP* genes, only *AlCSP2*, *AlCSP6*, and *AlCSP18* had antennae-biased expression; *AlCSP2* was male-biased and *AlCSP18* was female-biased in their expression. Six *AlCSP* genes (*AlCSP1*, 9, 12, 15, 16 and 20) were highly expressed in the proboscises, and nine (*AlCSP3-5*, 7, 8, 10, 13, 14 and 19) were highly expressed in the wings; among the 20 total *AlCSPs*, *AlCSP14* and *AlCSP5* displayed the highest and lowest expression levels in the antennae, respectively (Fig. 4).

DISCUSSION

In this study, we first identified 28 and 20 genes encoding putative *AlOBPs* and *AlCSPs*, respectively, based on our previous *A. lepigone* transcriptomic data (Zhang et al. 2016b). The number of *AlOBP* and *AlCSP* genes identified for this species are similar to some reported moths, such as *C. suppressalis* (Cao et al. 2014), *H. armigera* (Liu et al. 2012), and *B. mori* (Gong et al. 2007), but there are certain different from *S. litura* (Gu et al. 2015), *S. inferens* (Zhang et al. 2013), *H. armigera* (Liu et al. 2012) and *B. mori* (Gong et al. 2009). The reasons for the differences in gene number may be due to: 1) the different chemosensory behaviors of different moths requiring distinct molecular mechanisms that have developed over evolutionary time; 2) the genomic data will help us identify more genes from *A. lepigone* as well as from other moths in the future.

Many studies have shown that insect *OBPs* are mainly expressed in the antennae of both sexes and that they may play key roles in the process of host location, mating, and oviposition by allowing the insect to accurately recognize environmental odorants (Larter et al. 2016; Leal 2013; Qiao et al. 2009; Zhou et al. 2009). The phylogenetic tree of moth *OBPs* showed that *AlOBPs* were divided into different subfamilies, including the PBP/GOBP, Minus-C OBP, and Plus-C OBP proteins suggesting that the structural diversity of *AlOBPs* may be involved in chemosensation and/or in other physiological processes. Based on the qRT-PCR analyses, we

found that 46% of the 28 *AlOBPs* were highly expressed in the antennae indicating that these *AlOBP* proteins have putative roles in the odorant reorganization of *A. lepigone*. Similar to our previous work and to other studies (Gu et al. 2015; McKenzie et al. 2014; Zhang et al. 2016a; Zhang et al. 2013), we found that there were five *AlOBP* genes highly expressed in non-antennal tissues (legs and wings), including one abdomen-biased *AlOBP*-encoding gene and four wing-biased *AlOBP* genes, indicating that these OBPs may have other non-chemosensory functions.

Five *AlPBP*/*GOBPs* displayed higher expression in the adult antennae (especially *AlGOBPI* and *AlPBP1*), which is consistent with that reported for *PBP*/*GOBPs* in other moths (Liu et al. 2013; Liu et al. 2015b; Zhang et al. 2013). According to recent functional studies of moth *PBP*/*GOBPs* (Jin et al. 2014; Liu et al. 2015a; Liu et al. 2013; Sun et al. 2013; Zhu et al. 2016) and *D. melanogaster* LUSH protein (*OBP76a*) (Ha & Smith 2006; Stowers & Logan 2008; Zhou et al. 2004), we hypothesize that the *AlPBP*/*GOBPs* may also play important roles in recognizing the sex pheromones of female moths and some host plant volatiles. Additionally, there are three male-biased and three female-biased *AlOBP* genes, indicating that these sex-biased OBPs may participate in the reorganization of female or male sex pheromones, plant volatiles from oviposition sites, or other sex-related odorants, and need further analysis to explore their exact roles such as through fluorescence competitive binding assays (Liu et al. 2015b), CRISPR/Cas9 mediated genome editing (Zhu et al. 2016), and gene mutations (Stowers & Logan 2008).

Studies on *CSP* genes in certain insects have shown that they are smaller and more conserved than *OBP* genes and that they are widely expressed in different parts of the insect body (Calvello et al. 2005; Gong et al. 2007; Pelosi et al. 2014a; Zhang et al. 2013). Our BLASTX results showed that the *AlCSPs* had relatively high identities with other moth *CSPs* indicating high conservation of *CSPs* among moths. Our results agreed with those from studies using ligand-binding assays that found that some *CSPs* in other Lepidopterans have chemosensory roles including in *Mamestra brassicae* (Jacquin-Joly et al. 2001), *B. mori* (Qiao et al. 2013), and *S. inferens* (Zhang et al. 2014). Compared to the *AlOBP* genes highly expressed in the antennae, only three *AlCSPs* had antennae-biased expression, indicating that these three genes may be

involved in the recognition and transmission of sex pheromones, host volatiles, and other odorants. On the other hand, many insect CSPs are broadly expressed in non-chemosensory tissues and have non-chemosensory functions, such as SexiCSP3, which has been shown to have effects on the survival and reproduction of *S. exigua* (Gong et al. 2012), and AmelCSP5, which is involved in embryonic integument formation in *A. mellifera* (Foret et al. 2007). In this study, many AICSPs were found in various tissues and were highly expressed in non-chemosensory tissues suggesting that these AICSPs (especially AICSP14, which had the highest expression) may be involved in other physiological functions apart from chemosensory ones.

Furthermore, we found that there were eight AIOBPs (28.5% of all AIOBPs) and six AICSPs (30.0% of all AICSPs) that displayed proboscis-biased expression. OBP and CSP gene expression in the proboscis has been observed in other insects including *Apolygus lucorum* (Hua et al. 2012), *Lygus lineolaris* (Hull et al. 2014), *S. podoptera* (Liu et al. 2015c), and *A. dissimilis* (Sun et al. 2016). Further functional studies have also confirmed the gustation function of some genes: OBP49a in *D. melanogaster* is involved in the suppression of sweet taste by bitter chemicals (Jeong et al. 2013); some OBPs in *D. melanogaster* can modulate sucrose intake in response to a panel of nine bitter compounds by RNAi-mediated methods (Swarup et al. 2014); and CSP4, which is exclusively presented in the proboscis of two sibling species — *H. armigera* and *H. assulta* — an act as a wetting agent to reduce the surface tension of aqueous solutions (Liu et al. 2014). Therefore, the 14 AIOBPs and AICSPs with proboscis-biased expression may play similar gustation functions in *A. lepigone*.

In the future, these AIOBPs and AICSPs can help us develop environmentally friendly pesticides against *A. lepigone* based on reverse chemical ecology (Dominguez et al. 2016; Zhou 2010). We can explore the functions of candidate OBPs/CSPs *in vitro* to screen compounds with high binding affinities (e.g., host plant volatiles or sex pheromones) to target the OBPs/CSPs. These compounds could then be investigated as potential pesticides or sexual attractants. Further, with genetic modification by the CRISPR/Cas9 editing system (Hsu et al. 2014; Li et al. 2016; Zhu et al. 2016), we can knock out the candidate OBPs and CSPs to construct various mutant

strains and then release the effective strains into the field to disrupt the chemical communication behaviors of the pest.

CONCLUSION

In conclusion, we identified an extensive set of putative OBP- and CSP-encoding genes in *A. lepigone* based on our previous antennal transcriptomic data. As the first step towards understanding the functions of these genes, we conducted comprehensive and comparative phylogenetic analyses and developed gene expression profiles for OBPs and CSPs and found that most of the AIOBPs and AICSPs had high identities with other moth odorant genes. Nearly half of the AIOBPs displayed antennae-biased expression, but many AICSPs were detected in all tissues tested and were highly expressed in non-antennal tissues. Understanding the tissue and sex-biased expression patterns will help identify the functions of AIOBPs and AICSPs, which in turn will aid in elucidating the chemosensory mechanism of *A. lepigone* and developing environmentally friendly pesticides against this pest in future.

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ADDITIONAL INFORMATION AND DECLARATIONS

Data Availability

The following information was supplied regarding data availability: The sequences of AIOBPs and AICSPs have been supplied as Supplemental Information.

Supplemental Information

Supplemental information for this article can be found online.

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Tables Legends:

Table 1. The BLASTX match of OBP genes in *A. lepigone*.

Table 2. The BLASTX match of CSP genes in *A. lepigone*.

Figure Legends:

Fig. 1. Phylogenetic tree of moth OBPs. The *A. lepigone* translated genes are shown in blue. This tree was constructed using phyML based on alignment results of MAFFT. Al: *A. lepigone*, Bm: *B. mori*, Si: *S. inferens*, Sl: *S. littorali*, Ms: *M. sexta*.

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Fig. 3. Expression patterns of OBP genes in *A. lepigone*. The relative expression level is indicated as mean \pm SE (N = 3). Different capital letters mean significant difference between tissues ($p < 0.05$, ANOVA, LSD). FA, female antennae; MA, male antennae; Pr, proboscises; Ab, abdomen; Le, legs; Wi, wings.

Fig. 4. Expression patterns of CSP genes in *A. lepigone*. The relative expression level is indicated as mean \pm SE (N = 3). Different capital letters mean significant difference between tissues ($p < 0.05$, ANOVA, LSD). FA, female antennae; MA, male antennae; Pr, proboscises; Ab, abdomen; Le, legs; Wi, wings.

Supplemental Information

Table S1. Amino acid sequences of AIOBPs and AICSPs obtained in the study.

Table S2. Primers used for qRT-PCR.

Fig. S1. Multiple alignment of AIOBPs.

Fig. S2. Multiple alignment of AICSPs.

Table 1 (on next page)

Table 1. The BLASTX match of OBP genes in *A. lepigone*

1 **Table 1. The BLASTX match of OBP genes in *A. lepigone*.**

2

Gene	ORF	Signal	Complete	Best Blastx Match				
Name	(aa)	Peptide	ORF	Name	Acc. No.	Species	E value	Identity (%)
GOBP1	163	1-18	Y	general odorant binding protein 1	ABI24160.1	<i>Agrotis ipsilon</i>	8.00E-83	95
GOBP2	162	1-21	Y	general odorant binding protein 2	AHC72380.1	<i>Sesamia inferens</i>	2.00E-92	91
PBP1	167	1-23	Y	pheromone binding protein 1 precursor	AAC05702.2	<i>Mamestra brassicae</i>	3.00E-88	90
PBP2	170	1-24	Y	pheromone binding protein 2 precursor	AAC05701.2	<i>Mamestra brassicae</i>	5.00E-58	90
PBP3	164	1-22	Y	pheromone-binding protein 3	AFM36758.1	<i>Agrotis ipsilon</i>	2.00E-85	90
OBP1	116	N	N	SexiOBP14	AGP03460.1	<i>Spodoptera exigua</i>	7.00E-54	88
OBP2	146	1-17	Y	odorant binding protein 6	AGR39569.1	<i>Agrotis ipsilon</i>	2.00E-84	88
OBP3	120	N	N	odorant binding protein 8	AKI87969.1	<i>Spodoptera litura</i>	5.00E-79	85
OBP4	138	1-16	Y	odorant-binding protein 18	AFI57167.1	<i>Helicoverpa armigera</i>	2.00E-52	85
OBP5	147	1-21	Y	pheromone binding protein 4	AAL66739.1	<i>Mamestra brassicae</i>	1.00E-81	84
OBP6	134	1-17	Y	ABPX	AGS36754.1	<i>Sesamia inferens</i>	2.00E-54	83
OBP7	203	1-20	Y	odorant-binding protein 19	AGC92793.1	<i>Helicoverpa assulta</i>	2.00E-69	83
OBP8	147	1-20	Y	oderant binding protein 6	AFM77984.1	<i>Spodoptera exigua</i>	4.00E-56	82
OBP9	133	1-16	Y	odorant binding protein 9	AGH70105.1	<i>Spodoptera exigua</i>	5.00E-84	80
OBP10	96	N	N	odorant binding protein 1	AGR39564.1	<i>Agrotis ipsilon</i>	2.00E-58	79
OBP5	147	1-21	Y	pheromone binding protein 4	AAL66739.1	<i>Mamestra brassicae</i>	1.00E-81	84
OBP11	152	1-21	Y	pheromone binding protein 4	AAL66739.1	<i>Mamestra brassicae</i>	1.00E-30	78
OBP12	141	1-26	Y	odorant binding protein 8	AGH70104.1	<i>Spodoptera exigua</i>	9.00E-78	77
OBP13	184	1-20	Y	odorant binding protein	AII00978.1	<i>Dendrolimus houi</i>	1.00E-106	75
OBP14	186	1-17	Y	odorant binding protein 1	AGR39564.1	<i>Agrotis ipsilon</i>	8.00E-97	75
OBP15	155	1-24	Y	SexiOBP11	AGP03457.1	<i>Spodoptera exigua</i>	2.00E-82	73
OBP16	148	1-21	Y	OBP7	AEB54591.1	<i>Helicoverpa armigera</i>	7.00E-54	70
OBP17	246	1-19	Y	odorant binding protein	AII00994.1	<i>Dendrolimus kikuchii</i>	2.00E-74	67
OBP18	149	1-22	Y	OBP5	AEB54581.1	<i>Helicoverpa armigera</i>	8.00E-58	65
OBP19	71	1-22	N	OBP6	AGS36748.1	<i>Sesamia inferens</i>	2.00E-25	65

OBP20	170	1-23	Y	odorant binding protein 4	AKI87965.1	<i>Spodoptera litura</i>	2.00E-76	61
OBP21	153	1-21	Y	SexiOBP9	AGP03455.1	<i>Spodoptera exigua</i>	2.00E-77	59
OBP22	146	1-25	Y	SexiOBP12	AGP03458.1	<i>Spodoptera exigua</i>	1.00E-72	58
OBP23	145	1-17	Y	odorant binding protein	ADY17886.1	<i>Spodoptera exigua</i>	1.00E-85	40

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Table 2 (on next page)

Table 2. The BLASTX match of CSP genes in *A. lepigone*

1 **Table 2. The BLASTX match of CSP genes in *A. lepigone*.**

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Gene	ORF	Signal	Complete	Best Blastx Match				
Name	(aa)	Peptide	ORF	Name	Acc. No.	Species	E value	Identity (%)
CSP1	124	1-16	Y	chemosensory protein 15	AGH20053.1	<i>Helicoverpa armigera</i>	5.00E-62	83
CSP2	124	1-15	Y	chemosensory protein precursor	NP_001037066.1	<i>Bombyx mori</i>	2.00E-38	61
CSP3	122	1-18	Y	ejaculatory bulb-specific protein 3-like	XP_012549936.1	<i>Bombyx mori</i>	2.00E-45	74
CSP4	294	1-16	Y	chemosensory protein	AIW65104.1	<i>Helicoverpa armigera</i>	2.00E-130	78
CSP5	56	N	N	chemosensory protein	AII01011.1	<i>Dendrolimus houi</i>	3.00E-17	62
CSP6	150	1-19	Y	putative chemosensory protein	AGY49270.1	<i>Sesamia inferens</i>	6.00E-72	78
CSP7	114	1-19	Y	sensory appendage protein-like protein	AAK14793.1	<i>Mamestra brassicae</i>	1.00E-28	61
CSP8	127	1-18	Y	chemosensory protein 6	AGR39576.1	<i>Agrotis ipsilon</i>	5.00E-63	91
CSP9	127	1-16	Y	chemosensory protein	AAF71289.1	<i>Mamestra brassicae</i>	3.00E-59	83
CSP10	123	1-18	Y	chemosensory protein 8	AGR39578.1	<i>Agrotis ipsilon</i>	4.00E-68	85
CSP11	123	1-16	Y	chemosensory protein	AIW65100.1	<i>Helicoverpa armigera</i>	2.00E-65	76
CSP12	128	1-18	Y	chemosensory protein CSP2	ABM67689.1	<i>Spodoptera exigua</i>	4.00E-70	81
CSP13	123	1-19	Y	chemosensory protein	AIX97829.1	<i>Cnaphalocrocis medinalis</i>	1.00E-56	81
CSP14	46	N	N	putative chemosensory protein	AGY49260.1	<i>Sesamia inferens</i>	3.00E-25	100
CSP15	122	1-16	Y	chemosensory protein 10	AFR92094.1	<i>Helicoverpa armigera</i>	1.00E-73	89
CSP16	130	N	Y	chemosensory protein 15	NP_001091781.1	<i>Bombyx mori</i>	3.00E-42	59
CSP17	127	1-18	Y	putative chemosensory protein	AGY49267.1	<i>Sesamia inferens</i>	1.00E-70	81
CSP18	123	1-18	Y	chemosensory protein 8	AFR92092.1	<i>Helicoverpa armigera</i>	8.00E-43	74
CSP19	120	1-16	Y	chemosensory protein 4	AGR39574.1	<i>Agrotis ipsilon</i>	1.00E-60	79
CSP20	107	1-18	Y	chemosensory protein 5	AGR39575.1	<i>Agrotis ipsilon</i>	4.00E-53	97

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Figure 1

Fig. 1. Phylogenetic tree of moth OBPs

The *A. lepigone* translated genes are shown in blue. This tree was constructed using phyML based on alignment results of MAFFT. Al: *A. lepigone*, Bm: *B. mori*, Si: *S. inferens*, Sl: *S. littorali*, Ms: *M. sexta*.

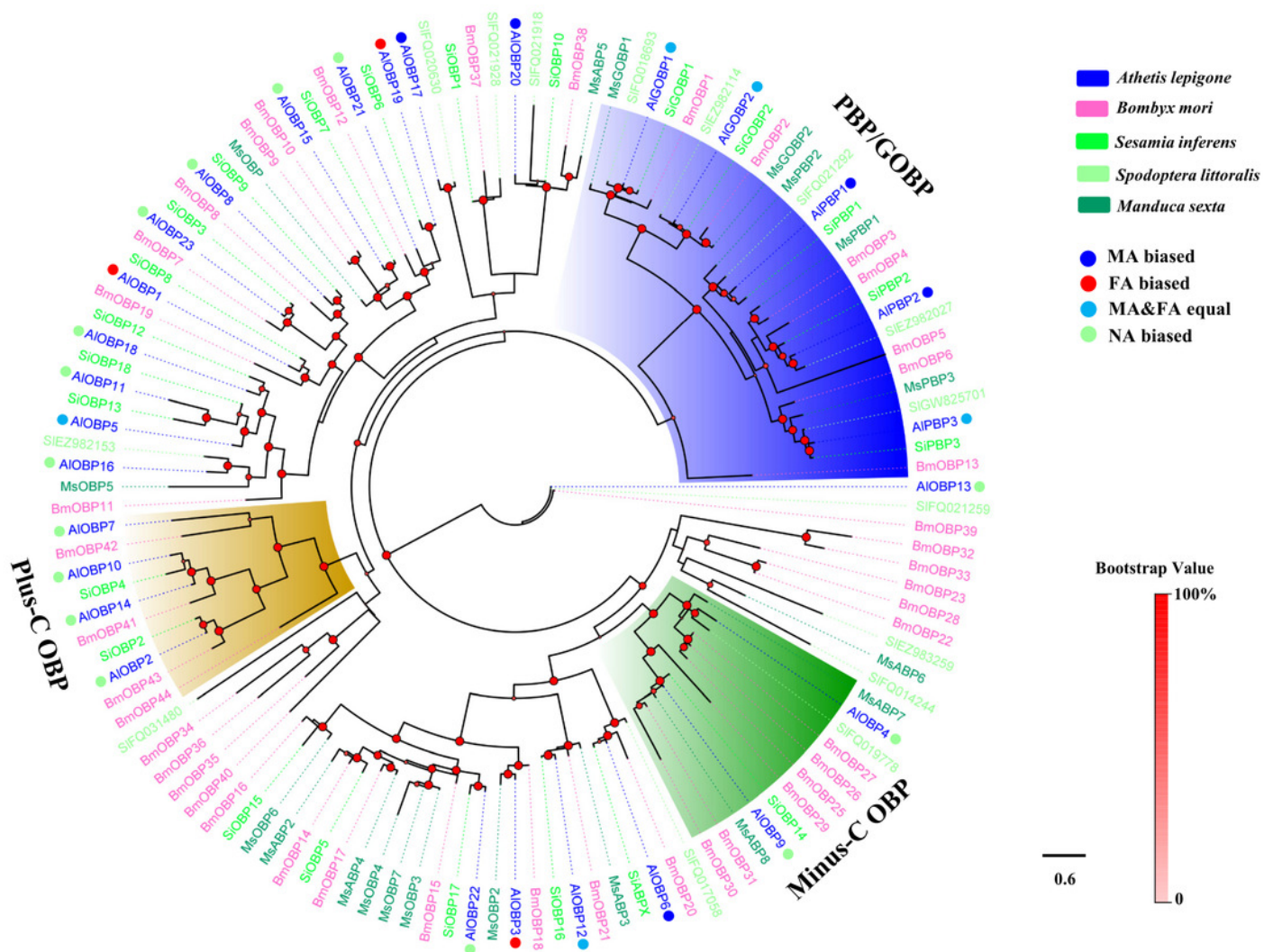


Figure 2

Fig. 2. Phylogenetic tree of moth CSPs

The *A. lepigone* translated genes are shown in blue. This tree was constructed using phyML based on alignment results of MAFFT. Al: *A. lepigone*, Bm: *B. mori*, Si: *S. inferens*, Sl: *S. littoralis*, Ms: *M. sexta*.

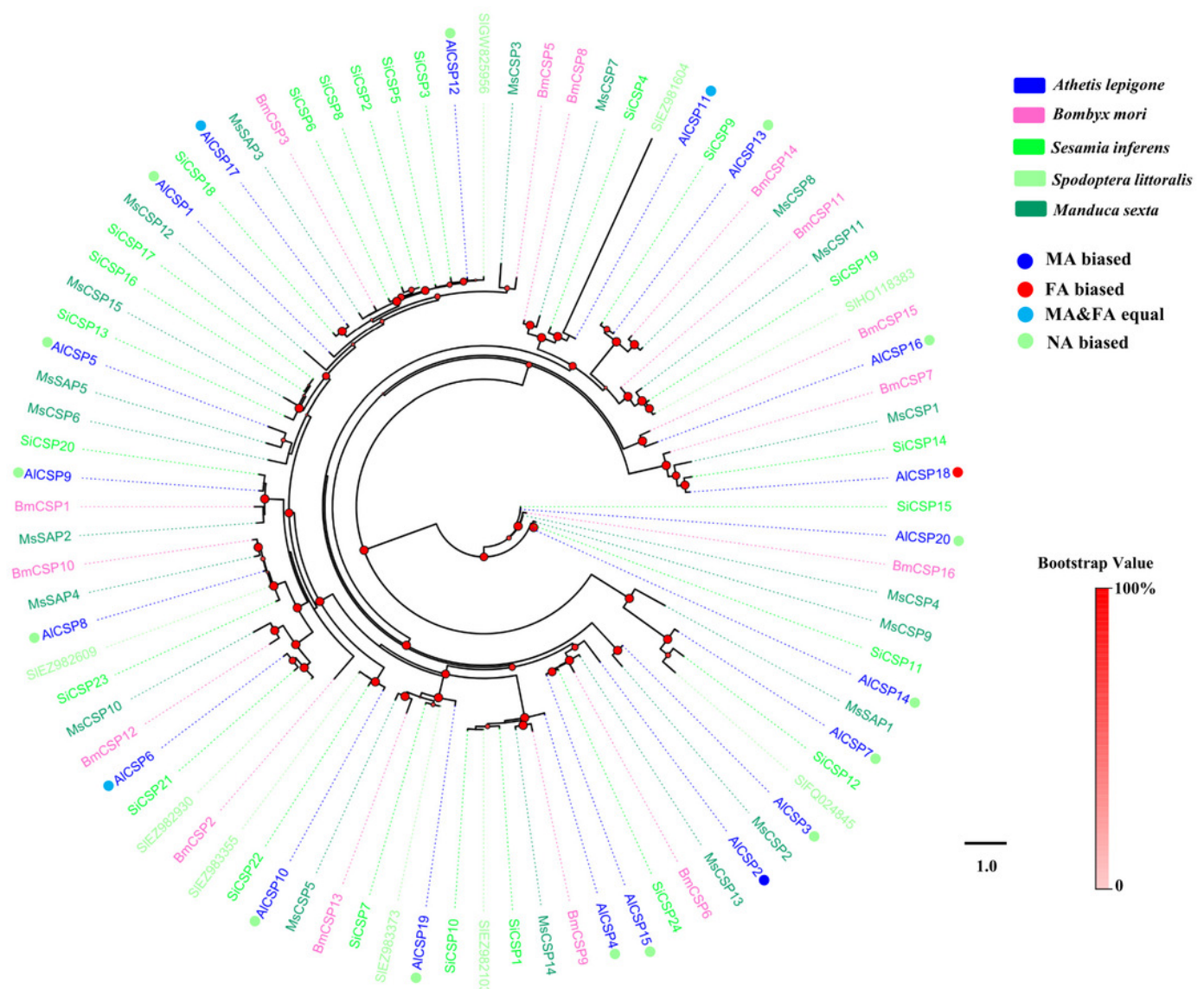


Figure 3

Fig. 3. Expression patterns of OBP genes in *A. lepigone*

The relative expression level is indicated as mean \pm SE (N = 3). Different capital letters mean significant difference between tissues ($p < 0.05$, ANOVA, LSD). FA, female antennae; MA, male antennae; Pr, proboscises; Ab, abdomen; Le, legs; Wi, wings.

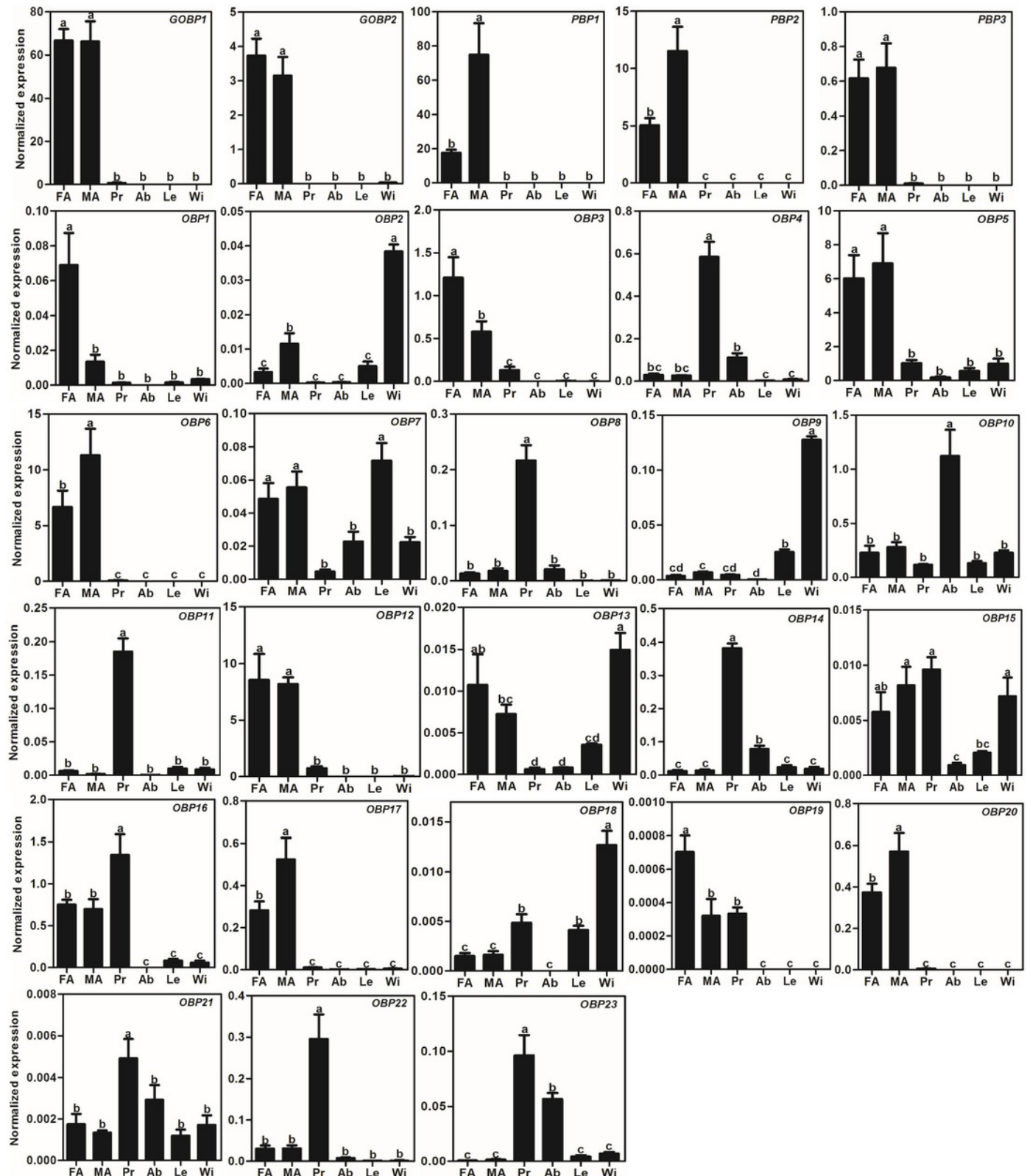


Figure 4

Fig. 4. Expression patterns of CSP genes in *A. lepigone*

The relative expression level is indicated as mean \pm SE (N = 3). Different capital letters mean significant difference between tissues ($p < 0.05$, ANOVA, LSD). FA, female antennae; MA, male antennae; Pr, proboscises; Ab, abdomen; Le, legs; Wi, wings.

