Male tarsi specific odorant-binding proteins in the diving beetle *Cybister japonicus* sharp

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Odorant binding proteins (OBPs) play critical roles in chemical communication of insects, as they recognize and transport environmental chemical signals to receptors. The diving beetle *Cybister japonicus* Sharp shows a remarkable sexual dimorphism. The foreleg tarsi of males are equipped with large suction cups, believed to help holding the female during underwater courtship and mating. Here, we identified two OBPs highly and specifically expressed in male tarsi, suggesting important functions of these structures in chemical communication. The first protein, *Cjap*OBP1, exhibits the 6 conserved cysteines motif of classic OBPs, while the second, *Cjap*OBP2, contains only four cysteines and can be assigned to the sub-class of C-minus OBPs. Both proteins were expressed in a bacterial system and the purified recombinant proteins were used for antibodies preparation. Western Blot analysis showed that *Cjap*OBP1 is predominantly expressed in male tarsi and could be also detected in antennae and palpi of both sexes, while *Cjap*OBP2, besides male tarsi, is also present in testis. Ligand-binding experiments showed a good binding affinity between *Cjap*OBP1, *Cjap*OBP2 and citral and coniferyl aldehyde, respectively. These results support a possible function of these two OBPs in the male foreleg tarsi of diving beetles in chemical communication.

Chemoreception in insects is mediated by membrane-bound receptors¹-⁵ and soluble proteins, named as odorant-binding proteins⁶-⁸ and chemosensory proteins⁹-¹². In particular, odorant-binding proteins (OBPs) are small proteins made of α-helical domains and folded into a compact structure¹³,¹⁴ further stabilized by three interlocked disulfide bridges¹⁵,¹⁶. Although six conserved cysteines are the landmark of classical OBPs, members with a lower or higher number of cysteines have been reported in all insect orders and are referred to as ‘Minus-C’, ‘Plus-C’ OBPs, respectively. In addition, atypical members with a different arrangement of the cysteines or with additional domains have also been described¹⁷-¹⁹.

OBPs are abundantly expressed in the lymph of chemosensilla, between the cuticle and the dendritic membrane of sensory neurons²⁰-²². The role of OBPs in olfactory processing is generally referred to as that of solubiliser and carrier of hydrophobic pheromones and odorants. Several recent studies have demonstrated that OBPs are also involved in detection and discrimination of semiochemicals²³-²⁹. Moreover, functional studies with receptors expressed in heterologous systems have reported that the presence of the appropriate OBPs increases the sensitivity and selectivity of the receptors to pheromones³⁰-³³. However, their specific function in sensory organs of insects and in particular the interplay between OBPs and olfactory receptors remain largely unknown.

Besides chemosensory organs, OBPs are also abundantly expressed in secretory glands and cells that often produce species specific pheromones. In such organs, OBPs are believed to help release of such semiochemicals into the environment. Several studies have been reported, in the pheromone glands of *Bombyx mori*³⁴ and *Agrotis ipsilon*³⁵, in the seminal fluid of *Aedes aegypti*³⁶ and *Helicoverpa* species³⁷, as well as in the mandibular glands of *Apis mellifera*³⁸. Some other OBPs are also involved in roles unrelated to chemoreception. For example, the OBPs6a expressed in the oral disk of the blow fly *Phormia regina* was suggested to be a fatty acid solubiliser³⁹. In *Aedes aegypti*, several OBPs were identified as components of the egg shell⁴⁰,⁴¹, suggesting their functions beyond chemoreception.

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Up to date, many OBPs have been identified in several species of Coleoptera, the largest order in the class of Insecta, including *Anomala osakana*⁴², *Papillia japonica*⁴² and *Phyllopertha diversa*⁴³, and more recently, in *Tenebrio molitor*⁴⁴, *Triobium castaneum*⁴⁵, *Anoplophora glabripennis*⁴⁶, *Dendroctonus valens*⁴⁷, *Dendroctonus ponderosae* (obtained with second generation deep sequencing) and *Lissorhoptrus oryzophilus* (Curculionoidea). However, information regarding olfactory proteins remains scarce in aquatic beetles.

The diving beetle *Cybister japonicus* Sharp (Coleoptera: Dytiscidae) lives in stagnant waters⁴⁸,⁴⁹ and is part of an important group of predatory insects. The beetles of the Dytiscidae family often show a range of peculiar secondary sexual characters in both males and females. The three basal segments of the pro-tarsi in males are usually equipped with various combinations of small and large suction cups. These cups are suggested to play roles in climbing, swimming and catching preys, as well as reproduction⁵⁰–⁵². In such context, males use specialized proleg tarsi to adhere the slippery females elytra⁵⁰.

In this study, we reported the first two OBPs in the diving beetle *C. japonicas*. The structure modeling, specialized expression in male tarsi and ligand-binding properties of these OBPs suggested their potential function in chemical communication between male and female diving beetles.

**Results**

Aim of the present work is to elucidate the function of the giant male front tarsi in water beetles through a study of the proteins expressed, focusing on odorant-binding proteins. The clear sexual dimorphism and the use of these anatomical structures in courtship and mating suggested that they might be involved in chemical communication.

**Morphology of proleg tarsi of *C. japonicus***. Although the peculiar structure of male tarsi in water beetles has been reported⁴², we first undertook a morphological investigation to describe in detail such organ in the diving beetle *C. japonicus* Sharp using scanning electron microscopy (SEM). Figure 1 clearly evidences the remarkable sexual dimorphism of the tarsi. The prolegs of males are equipped with large suction cups, which are absent in females (Fig. 1A,B). The male proleg presents an expanded palette composed of protarsomeres bearing specialized adhesive setae on their ventral side (Fig. 1A). Each spatula seta connects to the palette with an off-centre stalk and its ventral surface has an oval-shaped sucker from which parallel channels extend distally (Fig. 1C,D). The structural elements of this organ are very similar to those reported for the male diving beetle *Cybister rugosus*⁵⁰ and proposed to control female movement.

**Identification of odorant-binding proteins**. We hypothesized that the exaggerated growth of male tarsi could be related to semiochemical delivery. Therefore we decided to investigate the protein composition of the tarsi in a comparative way between male and female diving beetles. Electrophoretic analysis (SDS-PAGE) of crude extracts from the tarsi of male and female *C. japonicus* showed the presence of low molecular weight bands mainly in the male sample (Fig. 2A). Given the unusual abundance of these bands, migrating with apparent molecular masses compatible with the values expected for OBPs, we decided to investigate their protein composition. Thus, we separated a crude extract of male tarsi on a native gel (Fig. 2B) and selected two fast
migrating bands for N-terminal sequencing. Previous work had shown that generally OBPs migrate on the front of native gels and can be easily separated from other proteins. The two selected bands were blotted onto PVDF membrane and subjected to N-terminal sequencing. We obtained the N-terminal 20 amino acids sequences LDDAQKAKFKAHYDLCVTET and ISPEQKEKMKKLHDECLHET for these two proteins and temporary named them as protein 1 and protein 2.

**Molecular Cloning, sequence and analysis and structural modeling of CjapOBPs.** The information on the first 20 amino acids enabled us to design degenerate primers (Supplementary Table S1) at the 5′ ends of two genes. These degenerated primers together with oligo-(dT)18V at the 3′ end were used to amplify the relative genes with PCR. 5′-RACE was then performed to complete the sequences at the 5′ end, so we obtained the full-length genes, as reported in the Materials and Methods section.

On the basis of their significant similarities with OBPs from other species in National Center for Biotechnology Information (NCBI) database, the two deduced protein sequences were named CjapOBP1 and CjapOBP2 and deposited in the GenBank database with accession numbers KX078770 and KX078771, respectively.

*CjapOBP1* is a single polypeptide of 138 amino acids, with a calculated molecular weight for the mature protein of 14.1 kDa and a predicted isoelectric point of 5.49. *CjapOBP1* is similar in length, with 132 amino acids, a calculated molecular weight of 13.6 kDa and an isoelectric point of 6.74 for the mature protein. While *CjapOBP1* presents the six-cysteine signature of classic insect OBPs, *CjapOBP2* contains only four cysteines and can be assigned to the family of C-minus OBPs. The sequences of the two OBPs of *C. japonicus* were aligned with seven representative OBPs from other Coleoptera species (Fig. 3A). Phylogenetic analysis was performed based on amino acid sequences alignment of these beetle OBPs. The phylogenetic tree was generated by MEGA 6.06 with the neighbor-joining algorithm. The nine beetle OBPs separated into two groups, one group with six cysteine residues and the other having four cysteine residues (Fig. 3B). The overall structures of *CjapOBP1* and 2 (predicted by I-TASSER with highest ranking templates being 3qme and 3d77, respectively) consist of six helices (labelled α1 to α6) surrounding the hydrophobic ligand binding pocket (Fig. 3C, D). The predicted ligand binding site residues for *CjapOBP1* include seven hydrophobic amino acids (Phe67, Phe72, Ile73, Ile79, Met84, Tyr119 and Tyr122) (Fig. 3C). The putative ligand binding site residues for *CjapOBP2* consist of six hydrophobic amino acids (Met66, Phe67, Leu70, Phe72, Ile131 and Val132) (Fig. 3D).

**Specific localization of CjapOBPs in male proleg tarsi.** Using pET28a as a vector, we obtained the proteins fused with a His-tag fragment in high yields and in their soluble forms. Purification was performed by affinity chromatography on Ni-NTA His-Bind column. The electrophoresis analyses relative to expression and purification steps were reported in Supplementary Figure S1.

Western blot experiments, using a polyclonal antiserum raised against the purified protein, revealed that *CjapOBP1* is very abundant in male tarsi, with relatively low expression in palpi and antennae of both sexes (Fig. 4). *CjapOBP2*, instead was detected in testis and tarsi of males (Fig. 4). Both proteins, therefore appear to be
mainly expressed in the tarsi with a strong sex bias, suggesting their potential functions in chemical communica-
tion between male and female diving beetles.

Fluorescence binding assays. To characterize the ligand-binding properties of the two OBPs, we per-
formed fluorescence binding assays with several chemicals having different structures using N-phenyl-
1-naphthylamine (1-NPN) as a fluorescent reporter. In the absence of any information about potential
semiochemicals for this insect, as well as for related species, we selected some common plant volatiles, such as cit-
ral, coniferyl aldehyde, eugenol and 2-isobutyl-3-mthoxypyrazine, and structurally similar compounds. 1-NPN
binds $C_{jap}OBP1$ and $C_{jap}OBP2$ with dissociation constants of 3.3 and 2.1
$\mu$M respectively (Fig. 5A). Among the
ligands tested, we identified a single good ligand for each protein, citral for
$C_{jap}OBP1$ and coniferyl aldehyde
for
$C_{jap}OBP2$ (Fig. 5B,C). In particular, we can observe that several compounds structurally related to coniferyl
aldehyde, such as dihydroferulic acid, eugenol and $\alpha$-methoxycinnamaldehyde, did not show appreciable binding
to
$C_{jap}OBP2$, indicating that this protein might have a rather narrow binding structure spectrum.

Discussion
In this study, we have identified two OBPs predominantly expressed in the frontal tarsi of male $C. japonicus$. The
deduced amino acid sequences suggested that $C_{jap}OBP1$ consists of a typical framework of OBPs (six conserved
cysteines), while $C_{jap}OBP2$ contains only four cysteines and belongs to the family of C-minus OBPs (Fig. 3).
Both proteins present high sequence similarities with OBPs of other Coleoptera. The first Coleoptera insect OBPs
were identified from scarab beetles, specifically expressed in antennae, and associated with the pheromone rec-
ognition based on ligand-binding properties and single sensillum recordings evidences$^{42}$. With the advantage of
next-generation sequencing, a growing number of OBPs have also been reported in other Coleoptera species$^{54–58}$.
The diversified functions of these OBPs might be revealed in the future with future exploration.

$C. japonicus$ has marked sexual dimorphism of the tarsi with SEM investigation (Fig. 1). It has been known that
males use their giant front tarsi to hold on the back of females during their courtship$^{50–52}$. The high expression
of OBPs in the specialized front tarsi of males may suggest significant roles interfering with sex communication.
Because during the underwater courtship the female $C. japonicus$ could release sex pheromones used by males
for partner localization that was observed in the diving beetle Rhantus suturalis$^{59}$. Therefore it is reasonable to
hypothesize that these OBPs specific expressed in male foreleg tarsi *C. japonicus* may be involved in the sex pheromone reception. Work is in progress in our lab to determine the functions of *Cjap*OBP1 and *Cjap*OBP2 through functional RNA interference and behavior studies. In swallowtail butterfly *Papilio xuthus*, several chemosensory proteins and OBPs were identified in the female foreleg tarsi, which is served as a chemosensory organ in this species, suggesting an important function of these proteins in oviposition behavior of *P. xuthus*. Most recently, two putative OBPs show abundant expression in palps and foreleg tarsi of *Amblyomma americanum*, which contain the olfactory Haller's organ. These *A. americanum* OBPs may also play roles in sex pheromones reception. Future functional studies of these foreleg tarsi abundant OBPs will shed new lights on the understanding of molecular basis and evolution of chemoreception.

We can speculate that such proteins, rather than being involved in chemosensing, act as solubilisers and carriers of pheromones, and OBPs are transferred to females during such process together with their load of pheromones, as reported in several other species.

Investigating the binding affinity and distribution of expression of OBPs is useful for understanding the physiological function and mechanism of olfactory recognition. The fluorescent binding experiments showed that each protein is coupled with a single good ligand, citral for *Cjap*OBP1 and coniferyl for *Cjap*OBP2. The components of the sex pheromones in these beetles are unknown, but we can hypothesize that molecules structurally similar to citral and coniferyl aldehyde might represent good candidates, on the basis of our preliminary binding experiments.
Materials and Methods

Ethics statement. The use of rabbits in our experiments has been approved by the Institutional Animal Care and Use Committee of China Agricultural University (permit number: SYXK 2007–0023). All experimental protocols and procedures were carried out according to relevant regulations and guidelines established by this committee. All efforts were made to minimize the suffering of the rabbits.

Insect collection and rearing. Diving beetle *Cybister japonicus* Sharp were obtained from Guangdong Province, China, and raised in the Grassland Science Department (China Agricultural University, Beijing, China) at 28–30 °C, with photoperiod of 18 h:6 h (light: dark). The aquariums were filled with tap water, and the water was renewed twice a week. The aquariums were covered with fly screen to prevent beetles from flying away.

Scanning electron microscopy. For scanning electron microscopy (SEM), the samples of male prolegs were fixed in 70% ethanol for 2 h, and then cleaned in ultrasonic bath (250 W) for 1 min in the same solution. After treatment with 100% ethanol for 30 min, the samples were dried in air. Prolegs of male were mounted on holders, and after gold-coating, the samples were examined in a FEI Quanta 200 SEM (FEI Company, the Netherlands).

Figure 5. Affinity of *Cjap*OBPs to candidate chemical ligands. (A) Both *Cjap*OBP1 and *Cjap*OBP2 bind the fluorescent probe 1-NPN, with dissociation constants of 3.3 µM and 2.1 µM, respectively. (B) Competitive binding curves of selected ligands to the two *Cjap*OBPs. Mixtures of proteins (5 µM) and 1-NPN (2 µM) were titrated with 1mM solutions of the ligands in methanol. Among the ligands tested, *Cjap*OBP1 shows good affinity only to citral, and *Cjap*OBP2 bind coniferyl aldehyde strongly, but not dihydroferulic acid, eugenol and α-methoxy cinnamaldehyde, which are structurally related to coniferyl aldehyde. (C) The chemical structures of the chemicals applied.
Identification of *C. japonicus* OBPs. The different parts of the *C. japonicus* including antennae (30 pairs), tarsi (10 for male, and 30 for female), palpi (30 pairs), and sex organs (3) of both sexes were collected on ice and ground under liquid nitrogen. The samples were extracted with 0.3–1.0 mL 50 mM PBS (pH: 7.4), containing 1% PMSF and centrifuged at 13000 rpm for 20 min. Supernatants were collected and analyzed by 15% SDS-PAGE. For sequencing analysis, a sample of male tarsi was separated on a 15% native gel and blotted onto PVDF membrane. Fast migrating bands were subjected to automated Edman degradation, using a Procise 492 protein sequencer (Applied Biosystems, Foster City, Calif.), equipped with a 140C microgradient apparatus and a 785A UV detector (Applied Biosystems, Foster City, Calif.) for the automated identification of phenylthiohydantoin-amino acids.

**RNA extraction, cDNA synthesis, and cloning.** Total RNA was extracted using the TRizol® Reagent (Sigma, USA), following the manufacturer’s procedure. cDNA was synthesized with total RNA and Reverse Transcript PCR kit (Qiagen, Germany), along with the protocol provided. The PCR products were amplified using degenerated primers designed based on the amino acid residues of the N-terminal sequencing and oligo-(dT)αV primer (Supplementary Table S1). After a step at 50°C for 30 min and 95°C for 15 min, the reaction was performed for 35 cycles (95°C for 0.5 min, 47°C for 0.5 min, 72°C for 1 min), followed by a final step of 7 min at 72°C.

Crude PCR products (both about 500 bp) were then ligated into a pGEM-T (Promega, USA) vector. After transformation of *Escherichia coli* XL-1 Blue competent cells with the ligation product, positive colonies were selected by PCR using the plasmid’s primers SP6 and T7, then extracted and sequenced. Analysis of five colonies for each sample gave the same sequences.

**5’ RACE-PCR of putative *Cjap* OBP gene fragments.** Reverse transcription and 5’ rapid amplification of cDNA ends (RACE) were carried out with the SMART-RACE cDNA Amplification Kit (Clontech, USA). Total RNA from the front tarsi of male beetle was used as a template for first-strand synthesis with a 5’-CDs primer A and SMARTer II A oligo according to manufacturer’s directions. Second-strand synthesis was carried out with the PCR kit (Tiangen, China) with the 5’ RACE outer primer UPM (Clontech, USA) and specific forward primer. The primers were shown in Supplementary Table S1. The 20μl reaction mixture was initially incubated at 5 cycles of 94°C for 30 s, 72°C for 3 min, followed by 5 cycles of 94°C for 30 s, 70°C for 30 s, and 72°C for 3 min, and end with 27 cycles of 94°C for 30 s, 68°C for 30 s, and 72°C for 3 min. The single reaction product was gel isolated and cloned in pGEM-T (Promega, USA). Eight positive clones for each sample were sequenced.

**In silico structural analysis.** Amino acid sequences of *Cjap*OBPs and other beetle OBPs, including: *Lory*OBP2 (KF383281.1) and *Lory*OBP14 (KF383275.1) from *Lissorhoptrus oryzophilus*; *Tmo*OBP4 (KP071918.1) from *Tenebrio molitor*; *Tcas*OBP10 (XM_970591.2) from *Tribolium castaneum*; *Acor*OBP10 (KM251650.1) from *Anomala corpulenta*; *Hpar*OBP10 (KR733556.1) from *Holotrichia parallela*; *Dpon*OBP13 (KP736119.1) from *Dendroctonus ponderosae* were alignment using MUSCLE (http://www.ebi.ac.uk/Tools/muscle/). The secondary and tertiary structures of *Cjap*OBP1 and *Cjap*OBP2 were predicted by an online protein structure homology modeling server I-TASSER (http://zhanglab.ccmb.med.umich.edu/I-TASSER/). Then the Protein Data Bank (PDB) coordinate file of the highest ranking model was loaded into Chimera (http://www.cgl.ucsf.edu/chimera) for molecular visualizing and modification.

**Phylogenetic tree construction.** The insect OBP amino acid sequences were aligned using MUSCLE alignment default settings through MEGA version 6.06 (http://www.megasoftware.net/). The phylogenic tree was then generated by the neighbor-joining algorithm since the p-distance was <0.8 in overall distance. A total of 2,000 bootstrap replications were used to test of phylogeny. Finally, the selected tree was created with the cut-off value of 50%.

**Heterologous expression and purification of recombinant *Cjap*OBP proteins.** pGEM plasmids containing sequences of the mature *Cjap*OBPs were digested with NdeI and BamHI restriction enzymes. Then the products were ligated into a pET28a vector (Novagen, Germany) linearised with the same enzymes. The recombinant pET-*Cjap*OBP expression plasmid was transformed into *E. coli* BL21 (DE3) competent cells. Single colonies were grown overnight in 4 ml of Luria-Bertani (LB) broth supplemented with 50μg/ml kanamycin at 37°C with shaking at 200 rpm. The culture was diluted 1:100 with fresh LB broth (supplemented with 50μg/ml kanamycin) and grown at 37°C with shaking at 200 rpm until the OD₆₀₀ reached approximately 0.5. Isopropyl β-D-1-thiogalactopyranoside (IPTG) (Merck, Germany) was then added to the culture to a final concentration of 0.4 mM to induce expression of the target products for approximately 3 h at 37°C. After sonication of the bacterial pellet and centrifugation, all proteins were present in the supernatant, and the recombinant *Cjap*OBP protein were purified using a Ni-NTA His-Bind column following the provided protocol (Novagen, Germany).

**Preparation of antisera.** Polyclonal antisera against *Cjap*OBP1 and *Cjap*OBP2 were obtained by injecting rabbits subcutaneously and intramuscularly with 500μg of purified recombinant protein, followed by three additional injections of 300μg after 10 days each time. The protein was emulsified with an equal volume of Freund’s complete adjuvant (Sigma, St. Louis, MO, USA) for the first injection and incomplete adjuvant (Sigma, St. Louis, MO, USA) for further injections. The blood of rabbits was taken 7 days after the last injection, and centrifuged at 6000 rpm for 30 min. The supernatant of the blood was used without further purification. Rabbits were individually housed in large cages, at constant temperature, and all operations were performed according to ethical guidelines in order to minimize pain and discomfort to animals. The immunizations were performed according to the protocol approved by the animal care committee.
Western blot analysis. Extracts were prepared from antennae, mouth parts, tarsi, sex organs of adults of both sexes, as described in the Section of “Identification of C. japonicus OBPs” in “Materials and Methods”. After electrophoretic separation, gels were electroblotted onto nitrocellulose (NC) membrane (Millipore, USA) using a semi-dry protocol. Following overnight treatment with 2% powdered skimmed milk, the membrane was incubated with the crude antiserum at a dilution of 1:500 (2 h) and then with Goat Anti-Rabbit IgG (H+L), Horseradish Peroxidase Conjugate (Invitrogen, USA) (a dilution of 1:1000 for 1 h). Immunoreacting bands were detected by treatment with 4-chloro-1-naphthol and hydrogen peroxide.

Binding assays. To measure the affinity of the fluorescent ligand N-phenyl-1-naphthylamine (1-NPN), a 2μM solution of the protein in 50 mM Tris-HCl (pH 7.4) was titrated with aliquots of 1 mM 1-NPN in methanol to final concentrations of 1–16μM. The probe was excited at 337 nm, and emission spectra were recorded between 380 and 450 nm. The affinity of other ligands was measured in competitive binding assays, using the protein and the fluorescent reporter 1-NPN at the concentration of 5μM and 2μM respectively, while increasing the final concentration of each competitor up to 16μM. Dissociation constants for 1-NPN were evaluated from Scatchard plots of the binding data, and other ligands were calculated from the corresponding IC50 values, using the equation: 

\[ K_D = \frac{[I_{IC50}]/[1+1-NPN]}{K_{1-NPN}} \]  

being the free concentration of 1-NPN and K1-NPN being the dissociation constant of the complex protein/1-NPN.

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Acknowledgements

This work was supported by Natural Science Foundation of China (31372364), and the China Agriculture Research System (CARS-35-07), P. R. China. We thank Prof. Paolo Pelosi at Institute of Plant Protection, Chinese Academy of Agricultural Sciences for polishing the manuscript.
Author Contributions
L.-P.B. and L.-M.S. performed the experiments; X.J. provided the insect; J.-D.L. and X.-B.T. reared the insects; L.-P.B., X.-M.W. and Z.-H.Z. designed the work, and L.-P.B. and F.Z. analyzed the data and wrote the manuscript. All authors read and approved the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Song, L.-M. et al. Male tarsi specific odorant-binding proteins in the diving beetle Cybister japonicus sharp. Sci. Rep. 6, 31848; doi: 10.1038/srep31848 (2016).

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