

**Fatty Acid Solubilizer from the Oral Disk of the Blowfly**

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**Abstract**

**Background:** Blowflies are economic pests of the wool industry and potential vectors for epidemics. The establishment of a pesticide-free, environmentally friendly blowfly control strategy is necessary. Blowflies must feed on meat in order to initiate the cascade of events that are involved in reproduction including juvenile hormone synthesis, vitellogenesis, and mating. During feeding blowflies regurgitate salivary lipase, which may play a role in releasing fatty acids from triglycerides that are found in food. However, long-chain fatty acids show low solubility in aqueous solutions. In order to solubilize and ingest the released hydrophobic fatty acids, the blowflies must use a solubilizer.

**Methodology:** We applied native PAGE, Edman degradation, cDNA cloning, and RT-PCR to characterize a protein that accumulated in the oral disk of the blowfly, *Phormia regina*. In situ hybridization was carried out to localize the expression at the cellular level. A fluorescence competitive binding assay was used to identify potential ligands of this protein.

**Conclusion:** A protein newly identified from *P. regina* (PregOBP56a) belonged to the classic odorant-binding protein (OBP) family. This gene was expressed in a cluster of cells that was localized between pseudotracheae on the oral disk, which are not accessory cells of the taste peg chemosensory sensilla that normally synthesize OBPs. At pH 7 and pH 6, PregOBP56a bound palmitic, stearic, oleic, and linoleic acids, that are mainly found in chicken meat. The binding affinity of PregOBP56a decreased at pH 5. We propose that PregOBP56a is a protein that solubilizes fatty acids during feeding and subsequently helps to deliver the fatty acids to the midgut where they may help in the process of reproduction. As such, PregOBP56a is a potential molecular target for controlling the blowfly.

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**Introduction**

In the aftermath of the earthquake and subsequent tsunami that occurred in northern Japan on March 11, 2011, outbreaks of blowflies emerged from rotting fish, and affected public health [1]. Typically, the larvae of blowfly species produce wound myiasis. When blowflies strike sheep, the result is a reduction in the quality and quantity of wool, ewe infertility, and death, resulting in considerable economical losses in the wool industries [2,3]. Blowflies are also considered potential vectors of H5N1 avian influenza virus by mechanical transmission [4]. Additionally, there are problems with an increased incidence of insecticide-resistant influenza virus by mechanical transmission [4]. Additionally, there are problems with an increased incidence of insecticide-resistant blowflies and insecticide residues in the products of livestock [5]. Therefore, the establishment of environmentally friendly strategies for the control of blowflies is necessary especially in the wool industry and in terms of public health.

Blowflies must feed on meat in order to initiate the events involved in reproduction. In the black blowfly, *Phormia regina*, juvenile hormone (JH) biosynthesis is activated through the mevalonate pathway within 12 hours of feeding [6,7,8,9]. The increase in JH titer subsequently turns on male mating behavior [10]. In females, the increase in JH titer induces lipid and protein accumulation in the fat body from the digested meal, and increases vitellogenin synthesis. The vitellogenin and lipids released into hemolymph from the fat body accumulate into the developing oocytes under control of JH. JH is also involved in the regulation of male acceptance by females [8,11,12]. These findings imply that the newly ingested fatty acids are utilized immediately for JH synthesis in the corpora allata and continuously for vitellogenin synthesis in the fat body.

During feeding blowflies regurgitate salivary lipase, which might play a role in releasing fatty acids from triglycerides [13,14]. However, the solubility of long-chain fatty acids is extremely low in aqueous solution [15]. Although its exact function remains elusive, fatty acid-binding protein 5 has been identified in the human saliva which may help to solubilize long-chain fatty acids in human [16]. In order to effectively solubilize and ingest the hydrophobic fatty acids, the blowfly likely uses a similar solubilizer.

In order to identify fatty acid-binding protein(s) from the black blowfly, *P. regina*, we searched here protein expression from various tissues, including the oral disk, which makes direct contacts with the meal and presumably can accumulate proteins of our interest. A protein that accumulated at relatively high levels was detected in situ hybridization was carried out to localize the expression at the cellular level. A fluorescence competitive binding assay was used to identify potential ligands of this protein.
specifically in the oral disk. cDNA cloning and phylogenetic analysis demonstrated that the protein belongs to the insect classic odorant-binding protein (OBP) family, which plays a role in solubilizing hydrophobic ligands in aqueous solution [17,18,19]. We further characterized this protein by RT-PCR, in situ hybridization, and a fluorescence competitive binding assay, and discuss its presumed function.

Materials and Methods

Protein identification

A colony of the black blowfly, *P. regina* was maintained under the conditions described by Nishimura et al. [20]. In order to identify an oral disk-specific protein, each experimental tissue at day 3 of the adult stage was collected using fine clean forceps under a microscope. The tissue was homogenized in native PAGE sample buffer using a disposable pellet pestle (Trefflab, Deger- sheim, Switzerland). The protein homogenate was separated by 15% native PAGE [e-pagel, ATTO, Tokyo, Japan], and the gel was stained with Coomassie Brilliant Blue R-250.

In order to determine the N-terminal amino acid sequence of the oral disk-specific protein, protein was extracted from 500 ml labella that were collected from the blowflies. The labella were homogenized as described above in 10 mM Tris-HCl, pH 8, and the homogenate was centrifuged at 13,000 rpm at 4°C for 10 min to remove debris. The supernatant was partially separated on a column of DEAE Sepharose Fast Flow (200 ml gel volume, GE Healthcare, Little Chalfont, United Kingdom) by stepwise elution with 10 mM Tris-HCl, pH 8, containing 50, 100, 200, 250, 300, 400, and 500 mM NaCl. Proteins in the fractions containing 150 and 200 mM NaCl were concentrated, and subsequently separated by native PAGE using Tris-glycine running buffer. Following separation, the protein was electrophoretically transferred onto a polyvinylidene fluoride membrane (Millipore, Billerica, MA, USA) using 10 mM CAPS transfer buffer containing 10% methanol, pH 11, using a Trans-Blot SD Semi-Dry Transfer Cell (Bio-Rad, Hemel Hempstead, England, UK) by stepwise elution with 10 mM Tris-HCl, pH 8, containing 50, 100, 200, 250, 300, 400, and 500 mM NaCl. Proteins in the fractions containing 150 and 200 mM NaCl were concentrated, and subsequently separated by native PAGE using Tris-glycine running buffer. Following separation, the protein was electrophoretically transferred onto a polyvinylidene fluoride membrane (Millipore, Billerica, MA, USA) using 10 mM CAPS transfer buffer containing 10% methanol, pH 11, using a Trans-Blot SD Semi-Dry Transfer Cell (Bio-Rad Laboratories, K. K. Tokyo, Japan). Protein bands were visualized by Coomassie Brilliant Blue staining. The N-terminal sequences of a newly identified oral disk-specific protein and a chemical sense-related ligand-binding protein (CRLBP) were determined by Edman degradation.

cDNA cloning and bioinformatics analysis

Total RNA was extracted from an oral disk at day 0 adult stage using TRZol Reagent (Life Technologies, Carlsbad, CA, USA). The total RNA was used as template to synthesize cDNA using a SMART RACE cDNA Amplification Kit (Clontech, Mountain View, CA, USA) and Superscript II (Life Technologies) as a reverse transcriptase. The cDNA was treated with RNase H (New England Biolabs, Ipswich, MA, USA).

3’-RACE was performed using the degenerate primer, EEQKAKV-1, 5’-GA/GA/GGA/G/GA/G/GA/G/GA/G/GA/G/GC/G/C/G/TAA/G/G/GT-3’, designed on the basis of N-terminal sequence of an oral disk specific protein, EEQKAKV, which was determined by Edman degradation, and UPM supplied in the SMART RACE cDNA Amplification Kit, and Takara Ex Taq (Takara Bio, Otsu, Shiga, Japan) as a Taq DNA polymerase. The PCR cycles were as follows: 94°C for 3 min, 35 cycles of 94°C for 1 min, 55°C for 1 min, and 72°C for 1 min. The amplified PCR product was gel-purified, ligated into the Eco RV site of pBluescript II SK(+) (Agilent Technologies, Santa Clara, CA, USA), and sequenced on an ABI 3130xl sequencer (Applied Biosystems, Foster, CA, USA). 3’-RACE was performed using the gene-specific primer PregOBP56a-1 (5’-AGATTTC-GAAACCGGTAATCACAACGATC-3’) and UPM and KOD-plus (Takara, Osaka, Japan) as a Taq DNA polymerase. The PCR amplification was performed as follows: 95°C for 2 min, 35 cycles of 94°C for 15 s, 56°C for 30 s, and 68°C for 1 min. In order to further confirm the 3’-region, a second 3’-RACE was performed using the gene-specific primer PregOBP56a-2 (5’-GCCGCGTGATTTGTCACCCATTCAATTGACC-3’) and UPM. PCR was carried out as follows: 95°C for 2 min, 35 cycles of 94°C for 15 s, 67°C for 30 s, and 68°C for 1 min. The full-length cDNA sequence was determined using 9 independent clones to avoid PCR-derived sequence errors.

DNA sequence and blastp searches were performed using CLC DNA Workbench 5.7 software (CLC Bio, Aarhus N, Denmark). Phylogenetic analysis was carried out using the neighbour-joining method with 1000 bootstrap replicates using MEGA5 software [21] after sequence alignment with ClustalW.

RT-PCR

Total RNA was extracted from each experimental tissue and reverse-transcribed as described above. The PCR program was carried out using primers PregOBP56a-1 and PregOBP56a-2, or CRLBP-1 (5’-GAAGCTGGGGGATAGTCGCCACTTG-3’) and CRLBP-2 (5’-GCCATCATCACCTATCACCCTG-3’) using KOD-plus as a Taq DNA polymerase. The following cycles were used: 95°C for 2 min, 20 cycles of 94°C for 15 s, 59°C for 30 s, and 68°C for 1 min. The expression of actin was detected as an internal control [22]. The PCR-amplified products were separated on a 1.5% agarose gel electrophoresis, visualized under UV using a gel documentation system (Printgraph, ATTO, Tokyo, Japan), and cropped with Photoshop CS3 (Adobe, San Jose, CA, USA).

In situ hybridization

In situ hybridization was carried out according to the method described by Vosshall et al. [23]. The PCR amplicon generated by primers PregOBP56a-1 and PregOBP56a-2 was inserted into the Eco RV site of pBluescript II SK(+), digested with Eco RI and Hha III, and ligated into corresponding sites of pSPT18 (Roche Applied Science, Indianapolis, IN, USA). Digoxigenin (DIG)-labeled anti-sense and sense (control) probes were synthesized using a DIG-RNA labeling Kit (SP6/T7) (Roche Applied Science, Indianapolis, IN, USA). Digoxigenin (DIG)-labeled anti-sense and sense (control) probes were hybridized onto a DIG-RNA labeling Kit (SP6/T7) (Roche Applied Science) according to the instruction manual. Labella were collected from flies at day 0 adult stage, embedded in Tissue-Tek O.C.T. Compound (Sakura Finetek Japan, Tokyo, Japan), cut into 15 μm thick slices using a microtome, and dried on amino silane-coated glass slides (Matsunami Glass, Osaka, Japan). After 4% paraformaldehyde fixation for 10 min, the specimens were acetylated, prehybridized in hybridization buffer (50% formamide, 5× SSC, 5× Denhardt, 250 μg/mL yeast RNA, 500 μg/mL sonicated salmon sperm DNA, 50 μg/mL heparin, 2.5 mM EDTA, 0.1% Tween-20, and 0.25% CHAPS) at 55°C for 1 h, and hybridized overnight with 0.7 μg/mL of probe. Following probe hybridization, the specimen was washed 3 times with 5× SSC at 65°C, 0.2× SSC at 65°C for 20 min, and treated with PBS containing 0.1% Triton X-100 for 10 min. The specimen was then blocked with PBS containing 0.1% Triton X-100 and 10% normal goat serum (Wako Pure Chemical Industries, Osaka, Japan) for 1 h, and then reacted with anti-digoxigenin-Rhodamine Fab fragments (Roche Applied Science) at 1:1,000 dilution for 3 h. The slices embedded with FluorSave Reagent (Calbiochem, Darmstadt, Germany) were observed on a FLUOVIEW FV1000 confocal laser scanning microscope (Olympus, Tokyo, Japan). Other procedures were performed as reported previously [24].
Recombinant protein expression, purification, and evaluation

Recombinant PregOBP56a was expressed under oxidative conditions of the periplasmic space of bacteria resulting in the spontaneous formation of disulfide bridges, and purified as described previously [25]. In brief, cDNA encoding mature PregOBP56a was inserted into recognition sites of McI and Eco RI of pET-22b(+) (Novagen, Madison, WI, USA) and the construct was confirmed by DNA sequencing. The BL21(DE3) transformed with the bacterial expression vector was inoculated in LB medium including Carbencillin as an antibiotic, cultured at 200 rpm, 28°C for 16 h, induced by addition of isopropyl β-D-1-thiogalactopyranoside for 3 h, and harvested by centrifugation. The recombinant protein was extracted by 3 cycles of freeze and thaw procedure, removed debris by centrifugation, and purified by a combination with ion-exchange chromatography and gel filtration as described previously [26].

The purified recombinant PregOBP56a was dissolved in water and mixed with the matrix solution, saturated sinapinic acid. The recombinant protein/matrix mixture was spotted on a target plate and dried at room temperature. The MALDI TOF mass spectrum was collected in the positive ion reflector mode with an Autoflex III mass spectrometer (Bruker Daltonics, MA, USA). The N-terminal amino acids sequence of the recombinant protein was confirmed by Edman degradation.

Mass-finger print analysis was performed as follows. Five micrograms of the recombinant PregOBP56a (5 μl of a 1 μg/μl solution) was reduced by adding 10 μl of dithiothreitol solution (10 μg/μl in 100 mM Tris-HCl pH 8.5) under a nitrogen atmosphere for 2 hours at 45°C. The reduced protein was then S-carboxymethylated by adding 10 μl of sodium iodoacetate solution (25 μg/μl in 100 mM Tris-HCl pH 8.5) for 1 hour at 45°C. The carboxymethylated recombinant PregOBP56a was purified by reverse phase HPLC (PegasilODS, 4.6×250 mm, Senshu Scientific, Tokyo, Japan), equilibrated with 20% acetonitrile containing 0.03% trifluoroacetate, and eluted with a linear gradient of 20–40% acetonitrile over 20 min at a flow rate of 1.0 ml/min. The purified protein was lyophilized and then resuspended in 1.0 ml of sodium iodoacetate solution (25 μg/μl in 100 mM Tris-HCl pH 8.5) for 20 hours at 37°C. The reduced protein was then S-carboxymethylated by adding 10 μl of sodium iodoacetate solution (25 μg/μl in 100 mM Tris-HCl pH 8.5) for 20 hours at 37°C. Subsequently, the trypsinized fragments were separated by MALDI TOF mass spectrometer and the pattern was analyzed with AutollexIII (Bruker Daltonics) using α-cyano-4-hydroxycinnamic acid as a matrix.

Circular dichroism analysis

Far UV circular dichroism spectra at a range of 187–260 nm were recorded using a J-810 spectropolarimeter (Jasco, Easton, MD, USA). The purified recombinant protein was diluted to 4 mg/ml in 20 mM ammonium acetate, pH 7 or 20 mM sodium acetate, pH 5.

Fluorescence competitive binding assay

The fluorescence competitive binding assay was carried out as described previously [27,28]. In brief, N-phenyl-1-naphthylamine (1-NPN) (Tokyo Chemical Industry, Tokyo, Japan) was used as a fluorescent reporter. To measure the affinity of the recombinant PregOBP56a for 1-NPN, a 10 μg/ml solution of PregOBP56a in 20 mM phosphate-citrate buffer at pH 7, 6, or 5 was titrated with aliquots of 3.2 mM ligand solution in ethanol to final concentrations of 1.6–19.2 μM. Fluorescence was measured using a spectrophotofluorometer (F-2000, Hitach Hi-Technologies, Tokyo, Japan) after incubation for 2 min. Each sample, in the 2-ml SQ cuvette, was excited at 337 nm with emission and excitation slit widths of 10 nm and 10 nm. The emission spectra were recorded at a range of 350–500 nm. Cuvettes containing an appropriate concentration of 1-NPN were measured in parallel and the values obtained were subtracted as background fluorescent. The fluorescence intensity (440 nm) of a mixture of 1-NPN and recombinant protein without competitor was used as a reference (100%) to normalize the following measurements. The dissociation constant was calculated using Prism 6 software (GraphPad Software, La Jolla, CA, USA).

Decanoic acid, palmitic acid, stearic acid, oleic acid, linoleic acid, palmitoyl acetate, and cholesterol (Tokyo Chemical Industry) were used as competitors in the fluorescence competitive binding assay. After incubation of PregOBP56a with 1-NPN at a maximum binding concentration of 16 μM (pH 7), 14.4 μM (pH 6), or 19.2 μM (pH 5), each competitor was added to the PregOBP56a/1-NPN mixture at concentrations of 0-16 μM. The fluorescent intensity (440 nm) of the mixture of 1-NPN and recombinant protein without competitor was used as a reference (100%) to normalize the following measurements. The dissociation constant was calculated using Prism 6 software.

Results

Identification and cDNA cloning of an oral disk-specific protein

The aim of this research was to identify a potential solubilizer of long-chain fatty acids that are derived from the diet of the black blowfly, P. regina. In order to do this, protein from various tissues including the oral disk of female adult blowflies were analyzed by 15% native PAGE. Two major proteins that migrated to the middle of the native PAGE gel were detected from the oral disk. The slowly migrating protein was only observed in the oral disk (Figure 1, indicated by an arrow), but not in the rostrom. The oral disk-specific protein was identically observed in males (data not shown). It is known that a chemical sense-related lipophilic ligand-binding protein (CRLBP) is predominantly expressed in the oral disk of the blowfly. CRLBP belongs to the classic OBP family and is responsible for taste [29]. In order to characterize the newly identified oral disk-specific protein, Edman degradation was carried out. The N-terminal amino acid sequence of the protein, HLEETEEQKAKV, was different from that of CRLBP, ELITEKEAATTI. Further experiments of Edman degradation indicated that CRLBP was a protein migrating at front of the native PAGE under our experimental conditions (Tris-glycine running buffer) (Figure 1, indicated by an arrowhead). The basis of the staining intensity of Coomasie Brilliant Blue, the relative amount of our newly identified oral disk-specific protein was significantly higher than that of an OBP related to chemical reception.

By 5'- and 3'-RACE using a degenerate primer designed on the basis of N-terminal sequence, gene-specific primers, and UPM, a cDNA sequence of 587 bp was obtained (accession number, AB697136). The predicted mature protein was composed of 117 amino acid residues including 6 cysteine residues, which are hallmarks of insect classic OBP (Figure 2A). OBP plays a role in solubilizing hydrophobic ligands in aqueous solution in insect chemosensory organs [17,18,19]. This protein has a calculated molecular weight of 13,251 Da and isoelectric point of 6.36. Phylogenetic analysis indicated that this protein belonged to the cluster of dipteran OBPs (DmelOBP56a, CquiOBP56d, and AgamOBP[11]) but did not cluster with insect antennal binding.
To localize *PregOBP56a* expression in a female blowfly, RT-PCR analysis was carried out. We also investigated *CRLBP* expression as a reference of OBP related to chemical reception. Expression of *actin* as an internal control was detected from all experimental tissues. *PregOBP56a* was predominantly expressed in the oral disk with low level expression detected in the other chemosensory tissues including the maxillary palp and tarsus. On the other hand, *CRLBP* transcript was accumulated in the oral disk and antenna (Figure 3A). Oral disk-specific expression of *PregOBP56a* was consistent with the profile of proteins that was found by native PAGE analysis (Figure 1). The measured molecular mass of the recombinant protein was 15,245 Da (Figure S2), which was 6 Da less than the value calculated. These results suggested that the recombinant protein is a mature *PregOBP56a* that correctly formed 3 disulfide bridges as in the structure of insect classic OBPs. To analyze secondary structure of *PregOBP56a*, far-UV circular dichroism (CD) analysis was performed. The far-UV CD spectra showed one positive peak at 194 nm and two negative peaks at 209 nm and 221 nm (Figure 4), indicating that *PregOBP56a* is an α-helical-rich protein, as other insect OBPs reported previously [28,30,31,32,33,34].

Potential ligands of *PregOBP56a* expressing in the oral disk

Pseudotracheae on the oral disk of flies play a role in filtering out large particles in food [35,36,37], thus, blowflies only ingest liquid and/or tiny pieces of food. The chicken liver that is a primary component of the blowfly diet used for our colony contains hydrophobic long-chain fatty acids such as palmitic, stearic, oleic, and linoleic acids [38]. In order to address the ability of *PregOBP56a* to solubilize these fatty acids, we performed fluorescence competitive binding assay [27,28].

Addition of the fluorescent reporter, *N*-phenyl-1-naphthylamine (1-NPN) to a solution of *PregOBP56a* gave fluorescence emission spectra with a peak at 440 nm (Figure 5A). The concentrations of 1-NPN that produced maximum fluorescence were 16, 14.4, and 19.2 μM at pH 7, 6, and 5, respectively (Figure 5B). The dissociation constants of *PregOBP56a* (1-NPN) complex were 11.4, 12.3, and 17 μM at pH 7, 6, and 5, respectively.

The affinity of each ligand was evaluated using fluorescence competitive binding assay, where we measured the ability of each test compound to displace 1-NPN from the *PregOBP56a*-1-NPN complex. At pH 7, the addition of palmitic acid as a competitor to the *PregOBP56a*-1-NPN complex decreased the fluorescence intensity in a dose-dependent manner (Figure 5C). Also, at pH 7, a decrease in the fluorescence intensity (440 nm) was observed by addition of other fatty acids. Palmitic, stearic, oleic, and linoleic acids bound *PregOBP56a* with a calculated dissociation constant of 0.28, 0.35, 0.71, and 0.51 μM, respectively. However, the *PregOBP56a* did not bind to cholesterol (a major sterol in the chicken liver) and unrelated compounds such as decanoic acid (a short-chain fatty acid) or palmitoyl acetate (a long-chain ester) (Figure 5D). The binding assay results that were obtained at pH 6 were similar to those that were found at pH 7 (Figure 5E). At pH 6, palmitic, stearic, oleic, and linoleic acids bound *PregOBP56a* with calculated dissociation constants of 2.22, 1.24, 0.94,
and 2.40 μM, respectively. On the other hand, at pH 5 the binding of PregOBP56a to these fatty acids was significantly decreased (Figure 5F). Even when the tested fatty acids were added at a final concentration of 16 μM, the fluorescence intensity did not decrease below 50% (data not shown) and the dissociation constants could not be calculated.

Figure 2. Nucleotide and deduced amino acid sequences and phylogenetic analysis of oral disk-specific protein (PregOBP56a). A, cDNA and deduced amino acid sequences of an oral disk-specific protein. A signal peptide is shown in italic. N-terminal amino acid sequence determined by Edman degradation is underlined. Arrowheads show 6 cysteine residues, which are hallmarks of insect classic OBPs. The three arrows indicate annealing sites of a degenerate (EEQKAV-1) and two gene-specific (PregOBP56a-1 and PregOBP56a-2) primers. B, Phylogenetic tree of odorant-binding protein 56a like-protein sequences. Phormia regina odorant-binding protein 56a is shown in bold. Accession numbers of each protein are given in the parenthesis. Bootstrap values were determined from 1,000 replications. Bar indicates 5% divergence. doi:10.1371/journal.pone.0051779.g002
Discussion

In blowflies the absorption of fatty acids from the meat diet is essential for normal progress of reproductive processes including JH synthesis, vitellogenesis, and mating behavior [8,12,39]. Although long-chain fatty acids are released from the triglyceride in meat by digestion of salivary lipase, these compounds are insoluble in an aqueous solution [13,15]. In addition, since the pseudotrachea on the oral disk filter out particles, the blowflies only ingest a liquefied meal [35]. Therefore, some solubilizer is likely indispensable for the absorption of fatty acids.

PregOBP56a belongs to an insect classic OBP family. OBPs are known to accumulate in the sensillum cavity in chemosensory tissues and to encapsulate hydrophobic semiochemicals such as pheromones and odorants, and transfer then to the odorant receptor complex in the dendrite of the receptor neuron [17,18,19]. Pheromone-binding protein from the silkworm, Bombyx mori (BmorPBP) belonging to the same family is composed of 6 $\alpha$-helices knitted with 3 disulfide bridges [40]. Because of the structural properties, this protein appears to be highly stable to endopeptidase digestion, heat inactivation, and low pH [26,32]. Although crystallographic assignment is required, based on our results obtained by MALDI TOF mass spectrometry and circular dichroism analysis, PregOBP56a might have similar stability as BmorPBP.

A slight difference in the far-UV CD spectra of PregOBP56a was observed between pH 7 and pH 5 (Figure 4). A similar difference was previously observed in dipteran OBP [25]. These results suggested that the $\alpha$-helical content of PregOBP56a might be unwound slightly, presumably causing dissociation of ligands at pH 5. It is known that mosquito OBPs form a dimer and that ligands bind in the long tunnel-binding cavity that is formed by dimer [41,42,43]. PregOBP56a possesses short peptides at the N- and C-termini, the structure of which is observed among dipteran OBPs but not in lepidopteran OBPs. Therefore PregOBP56a might bind fatty acids in a similar manner. Further analyses based on NMR spectroscopy and crystallography will clarify the molecular mechanism related to ligand binding and release.
Figure 5. Binding of PregOBP56a to \( N \)-phenyl-1-naphthylamine (1-NPN) and potential ligands derived from chicken meat. A, A typical emission spectra following the addition of 1-NPN (3.2–16 \( \mu \)M final concentration) to recombinant PregOBP56a (10 \( \mu \)g/ml of 20 mM phosphate-citrate buffer, pH 7). The emission spectra were generated following excitation at a wavelength of 337 nm. B, Binding of 1-NPN to PregOBP56a at pH 7, 6, and 5. In these experiments PregOBP56a (10 \( \mu \)g/ml) was incubated with various concentrations (1.6–19.2 \( \mu \)M) of 1-NPN and emission at 440 nm was plotted. Values are means ± standard deviation, n = 3. C, Typical emission spectra following the addition of a competitor to PregOBP56a bound to 1-NPN. PregOBP56a (10 \( \mu \)g/ml) in 20 mM phosphate-citrate buffer at pH 7 was pre-incubated with 16 \( \mu \)M of 1-NPN (final concentration). Then, palmitic acid (0–3.2 \( \mu \)M final concentration) was added to the reaction, and the emission spectra (excitation a wavelength of 337 nm) were recorded. D, Binding of PregOBP56a to various ligands at pH 7. For each set of data, fluorescence values at a wavelength of 440 nm were plotted as percent of that obtained in the absence of the competitor. Values are means ± standard deviation, n = 3. The competitors are shown.
Conclusions

In this study we demonstrated that PregOBP56a is expressed in a cluster of cells that is found between pseudotracheae. To date, OBPs are expressed not only in chemosensory tissues, but also in various parts of an insect body [28,44,45,46,47,48,49,50]. It is reported that odorant-binding protein 19d from Drosophila melanogaster (DmeOBP19d) [previous name, PBPRP2] uniquely accumulates in the subcuticular space of the pseudotracheal region of the oral disk [47]. This localization resembles that of PregOBP56a. Dipteran insects might generally use this space to store OBPs. In the blowfly, it has been shown that the sensillar lymph containing CRPBP is exuded through a top pore of the taste sensillum on the oral disk [29,51]. Similarly, PregOBP56a might be released through an unidentified pore(s) on the cuticle of oral disk or pseudotracheae. Further immunohistochemistry and electron microscopic study will clarify this secretion mechanism.

PregOBP56a showed specific binding to palmitic, stearic, oleic, and linoleic acids that are primary compounds found in chicken meat. This binding was highest, at pH 7 and pH 6 but decreased dramatically at pH 5. At 6 h postmortem the pH of the meat decreases to 6.1 by lactate generation during glycolysis [52]. PregOBP56a might be capable of binding palmitic, oleic, and linoleic acids in the meat up to 6 h postmortem. In addition, the pH of intestinal tract of the blowfly decreases from 7 [crop] to 3.5–6.2 (anterior midgut), and to 2–3.3 (coiled midgut) [53]. Thus, all fatty acids delivered by PregOBP56a will likely be dissociated in the midgut lumen. Subsequently, they can be absorbed and converted diacylglycerols in the enterocytes as is done in other insects [54]. Once in the hemolymph, lipophorin may take them to the target organs such as, corpus allata, fat bodies, and ovaries, respectively. We propose that PregOBP56a is a protein that solubilizes fatty acids during feeding and help in their delivery to the midgut for continuation of normal reproductive processes.

Conclusions

Blowflies are economic pests of the wool industry and also insects of medical importance. Blowflies must feed on meat for normal progress of reproductive processes including juvenile hormone synthesis, vitellogenesis, and mating. The long-chain fatty acid components of meat are essential for this process. The newly identified PregOBP56a was specifically expressed in a cluster of the cells that are localized between pseudotracheae on the oral disk. These cells come into direct contact with the meat meal during feeding. PregOBP56a bound fatty acids at pH 7 and pH 6, but not at pH 5. These results suggested that PregOBP56a solubilizes fatty acids from fresh meat and delivers them to the midgut where low pH environment facilitates release and subsequent absorption. Since blowflies require long-chain fatty acids for normal reproductive processes, PregOBP56a might be a potential molecular target for controlling the reproduction of blowflies.

Supporting Information

Figure S1 Mass-finger printing of recombinant PregOBP56a. A, Mass-fingerprinting of recombinant PregOBP56a. Molecular weight and corresponding positions were represented. Peaks with asterisks were [M+Na]+ ions. B, Tryptic fragments of PregOBP56a. Identified fragments were represented in bold letters.

(PPT)

Figure S2 MALDI TOF mass spectrometry of recombinant PregOBP56a.

(PPT)

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Author Contributions

Conceived and designed the experiments: YI JI WSL. Performed the experiments: YI JI. Analyzed the data: YI JI WSL.Contributed reagents/materials/analysis tools: YI JI WSL. Wrote the paper: YI JI WSL.

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