Identification and functional characterization of a sex pheromone receptor in the silkmoth *Bombyx mori*

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Sex pheromones released by female moths are detected with high specificity and sensitivity in the olfactory sensilla of antennae of conspecific males. Bombykol in the silkmoth *Bombyx mori* was the first sex pheromone to be identified. Here we identify a male-specific G protein-coupled olfactory receptor gene, *B. mori* olfactory receptor 1 (*BmOR-1*), that appears to encode a bombykol receptor. The *BmOR-1* gene is located on the Z sex chromosome, has an eight-exon/seven-intron structure, and exhibits male-specific expression in the pheromone receptor neurons of male moth antennae during late pupal and adult stages. Bombykol stimulation of *Xenopus laevis* oocytes expressing *BmOR-1* and *BmGq41C1* induced robust dose-dependent inward currents on two-electrode voltage clamp recordings, demonstrating that the binding of bombykol to *BmOR-1* leads to the activation of a BmGq-mediated signaling cascade. Antennae of female moths infected with *BmOR-1*-recombinant baculovirus showed electrophysiological responses to bombykol but not to bombykal. These results provide evidence that *BmOR-1* is a G protein-coupled sex pheromone receptor that recognizes bombykol.

baculovirus | bombykol | bombykal | olfactory receptor | *Xenopus laevis* oocyte

*Insects* use a unique class of chemical signals called pheromones as cues to recognize other members of the same species and as a means to induce a particular behavior in other members of the same species (1–4). The silkmoth, *Bombyx mori*, possesses the simplest pheromone system in which full sexual behavior of male moths is initiated by one achiral compound, (E,Z)-10,12-hexadecadien-1-ol (bombykol) released from the pheromone gland of female moths (2–4). Thereby, bombykol is thought to be the sole sex pheromone in *B. mori*. Bombykol, an oxidized form of bombykal, is also released by female moths, but it does not elicit orientating behavior in male moths (5). Two specialized chemosensory neurons in the long sensilla trichodea on the male antennae are fine-tuned to detect either bombykol or bombykal (5). This remarkable sensitivity and specificity is thought to be achieved by olfactory receptors (ORs) expressed in individual chemosensory neurons.

The OR gene families in insect species have been identified by comprehensive analysis of genome sequences (6–14). In *Drosophila* antennae, expression of a total of 31 conventional ORs was reported (9), and 24 of these appear to show responses to general odors with a broad and overlapping ligand-spectrum (15). Therefore, insects use a combinatorial coding strategy to discriminate various odors as has been previously addressed in mammalian species (16, 17). Receptors for insect pheromones, however, remain unidentified. Pheromone receptors should possess a fine-tuned ligand spectrum with a high discriminatory power. Further, it is reasonable to suggest that a sex pheromone receptor should be expressed specifically in male moths, but not in female moths. In this regard, male-specific OR genes have recently been identified in the genome database of tobacco budworm *Heliothis virescens* (14).

In the present study, we have undertaken a differential screening strategy to isolate the male-specific *B. mori* OR (*BmOR*) gene and cloned one gene designated *BmOR-1*. Using a *Xenopus laevis* oocyte expression system, we obtained functional evidence that *BmOR-1* encoded an OR that specifically recognized bombykol. Further, ectopic expression of *BmOR-1* in female antennae produced electrophysiological responses to bombykol. Finally, none of other male-specific ORs among 29 putative OR genes encoded in the *B. mori* genome showed any response to bombykol. Functional characterization of *BmOR-1* in both heterologous and homologous systems and the results from comprehensive genome database mining most likely identify *BmOR-1* as a single sex pheromone receptor in *B. mori*.

Materials and Methods

**Synthesis of Bombykol and Bombykal.** (E,Z)-10,12-hexadecadien-1-ol (bombykol) was synthesized stereospecifically starting from 1-pentyne (Sigma-Aldrich) and 10-undecyn-1-ol (Tokyo Kasei, Tokyo) following the protocol described (18). Distillation gave pure bombykol, and the stereochemistry was confirmed by the 1H-NMR spectrum at 300 MHz. (E,Z)-10,12-hexadecadien-1-ol (bombykal) was prepared by Dess-Martin oxidation (19) of bombykol.

**Animals and Chemicals.** Silkmoths *B. mori* were kept at 25°C on a 12 h:12 h (light/dark) light cycle. Dissected tissues were frozen immediately in liquid nitrogen and stored at −80°C until use. Odorants used in this study were kindly provided by T. Hasegawa (Tokyo) and Takasago (Tokyo) or purchased from Wako and Sigma-Aldrich.

**Differential Screenings.** A *B. mori* male antennae cDNA library (20) was plated at low density (1,000–1,500 plaque-forming units per 140 × 100-mm plate), and phage DNA was transferred to nitrocellulose membranes in triplicate by using standard procedures. Each membrane was separately hybridized at high stringency with one of the following fluorescein-labeled probes: day-0 male antennae cDNA as a positive probe, day-0 male body
sis, total RNA was extracted from the egg, the head of fifth-instar larvae, and the antennae of pupae and adults. RT-PCR was performed as described above. No PCR products were produced when reverse transcriptase was excluded during reverse tran-

**BmOR-1 Genomic Sequences.** Genomic DNA of adult male moths was extracted by using GenomicPrep cells and tissue DNA isolation kits (Amersham Pharmacia). *BmOR-1* was amplified from *B. mori* genomic DNA by using LA Taq polymerase (Takara Shuzo, Kyoto) and primers 5'-GGATAGAATATCTTC-GATCTTCGCACTG-3' and 5'-TGTGGCCACCGTTGAAAGAT-CACG-3', which corresponded to nucleotides 25–47 or 1,326–1,350 in the *BmOR-1* cDNA sequence. Amplification was performed by using the following thermal program: 94°C for 1 min; 30 cycles of 94°C for 30 s, 63°C for 30 s, and 68°C for 10 min; followed by one cycle at 72°C for 10 min. A single 9.4-kb PCR fragment was produced in the PCR. The fragment was cloned into a pGEM-T vector (Promega) and sequenced with vector- or sequence-specific primers. The ends of the *BmOR-1* genome sequence that were not included in the PCR products were sequenced by using the bacterial artificial chromosome clone as described below.

**Isolation of Bacterial Artificial Chromosome Clones and Mapping Sequences of *BmOR-1*.** The *B. mori* bacterial artificial chromosome library was screened by PCR (22) by using two pairs of primers specific to two fragments of different lengths of *BmOR-1* (610-bp fragment, 5'-AACAATCTGAGAATATACCG-3' and 5'-TGAACGCGAAGCAGAAGC-3'; 602-bp fragment, 5'-TGGGAAGTGATGGATGAGAA-3' and 5'-TATATA ATTGTAGGGTGAGA-3'). One of the isolated clones contained the B17F10 OR sequence, which is known as a marker sequence of the Z chromosome of *B. mori* (information available at http://sosui.proteome.bio.tuat.ac.jp/cgi-bin/sosui.submit.html) and TMPRED (www.ch.embnet.org/software/TMPRED.form.html). Arrowheads, intron–exon boundaries. (b) Schematic representation of the genomic structure of *BmOR-1*. The position and relative size of exons and introns are drawn to scale as indicated. Structure of the *BmOR-1* gene and locations of start/stop codons are shown.

**RT-PCR of RNAs Isolated from Tissues.** Total RNA was separately extracted from day-0 adult moth antennae, head, legs, wings, thorax and abdomen tissues by the acid guanidinium-phenol-chloroform method (23), treated with DNase I, and reprecipitated. RNA was reverse-transcribed by using oligo(dT) adaptor primer (Takara Shuzo) and avian myeloblastosis virus reverse transcriptase (Takara Shuzo) at 42°C for 35 min. cDNA of *BmOR-1*, *BmOR-2*, *PBP* and *B. mori actin 1* (24) was amplified by using Ex Taq DNA polymerase (Takara Shuzo) and the *BmOR* DNA sequence. The following primer pairs that were designed to span at least one predicted intron except for actin gene to distinguish between *BmOR-1* and *BmOR-2* transcriptase (Takara Shuzo) and avian myeloblastosis virus reverse transcriptase (Takara Shuzo) at 42°C for 35 min. cDNA of *BmOR-1*, *BmOR-2*, *PBP* and *B. mori actin 1* (24) was amplified by using Ex Taq DNA polymerase (Takara Shuzo) and the *BmOR* DNA sequence. The following primer pairs that were designed to span at least one predicted intron except for actin gene to distinguish between *BmOR-1* and *BmOR-2*.
antisense RNA. A Scale bar: 50 μm. 

The male antenna by using DIG-labeled BmOR-1 and fluorescein-labeled PBP labeling of a 2-μm plastic section of the male antenna viewed from olfactory sensilla side. (In Fig. 2.) Whole-mount in situ hybridization was performed according to Tautz and Pfeifle (25) with minor modifications. In brief, antennae of pupae on days 1–2 before eclosion were dissected from the animal, cut into pieces, fixed with 4% paraformaldehyde/PBS overnight at 4°C, treated with 50 μg/ml proteinase K/PBS for 1 h at 37°C, and then fixed for an additional 5 min in 4% paraformaldehyde/PBS. Samples were washed three times for 5 min each time in 0.1% Tween 20/PBS (PBST) at room temperature, followed by an incubation for 5 min in 1:1 PBST/hybridization buffer (50% formamide/5× SSC/50 μg/ml heparin/0.1% Tween 20/100 μg/ml herring sperm DNA). Hybridization reactions were carried out for 16 h at 60°C by using the BmOR-1 probe at a concentration of 500 ng/ml. Hybridized antennae were washed for 10 min at 60°C in hybridization buffer and 10 min in 1:1 PBST/hybridization buffer, followed by three washes in PBST for 10 min each. Hybridization was detected by using alkaline phosphatase-conjugated anti-DIG Ab (Roche; 1:5,000 in PBST) and stained with nitroblue tetrazolium/5-bromo-4-chloro-3-indolyl phosphate. Then, the antenna was embedded in Epon-82 resin (TAAB Laboratories Equipment, Reading, United Kingdom) and cut into 2-μm plastic sections.

For fluorescent in situ hybridization, day-0 adult moth antennae were fixed in 4% paraformaldehyde/PBS overnight at 4°C, dehydrated, embedded in paraffin, and cut into 10-μm sections. Tissue sections were incubated for 16 h at 60°C in 100 μl of hybridization buffer containing 500 ng/ml of both DIG-labeled BmOR-1 and fluorescein-labeled PBP antisense RNA probes. Sections were washed as described above; the hybridization signal was amplified by using the tyramide signal amplification plus fluorescein signal detection (PerkinElmer), and signal amplification was carried out according to the manufacturer’s instructions. DIG-labeled probes were visualized by using horseradish peroxidase-conjugated anti-DIG Ab (Roche; 1:100) with tetramethyl-rhodamine tyramides as a substrate, whereas fluorescein-labeled probes were visualized by using horseradish peroxidase-conjugated anti-fluorescein Ab (Roche; 1:100) with fluorescein tyramides as a substrate.

Gene Expression in X. laevis Oocytes and Electrophysiological Recording. Stage V–VII oocytes were treated with 2 mg/ml collagenase type I (Sigma-Aldrich) in Ca2+-free saline solution (82.5 mM NaCl/2 mM KCl/1 mM MgCl2/5 mM Hepes, pH 7.5) for 1–2 h at room temperature. Oocytes were then microinjected with 50 ng of BmOR-1 cRNA and 25 pg of BmGqα cRNA synthesized from pSPUTK-BmOR-1 and pSPUTK-BmGqα, respectively (26). Oocytes injected with 50 ng of BmOR-1 cRNA or 25 pg of BmGqα cRNA were used as negative controls. Injected oocytes were incubated for 5–7 days at 18°C in Barth’s solution (88 mM NaCl/1 mM KCl/0.3 mM Ca(NO3)2/0.4 mM CaCl2/0.8 mM MgSO4/2.4 mM NaHCO3/15 mM Hepes, pH 7.6) supplemented with 10 μg/ml penicillin and streptomycin.

Whole-cell currents were recorded with a two-electrode voltage clamp (OC-725, Warner) (27). Data acquisition and analysis were carried out by using Digidata1322A and pCLAMP software (Axon Instruments, Foster City, CA). Ca2+-dependent Cl− current was monitored by applying 200-msec depolarizing pulses every 2 sec of +60 mV from the holding potential of −80 mV.
responses were averaged for each concentration: 30 bombykol responses in 20 bombykol was monitored by an increase in Ca\textsuperscript{2+} current. (Upper) Current traces of oocytes injected with cRNAs of BmOR-1 and BmG\textsubscript{q} (BmOR-1 + BmG\textsubscript{q}) or noninjected control oocytes (no injection) before and after application of 100 \mu M bombykol. (Lower) Time course of the response plotted as the current amplitude at the end of each depolarizing pulse. Bombykol (100 \mu M) was applied at the time indicated by arrowheads. The data are representative of bombykol responses in 20 X. laevis oocytes from five different animals. (c) Time course of current recordings of oocytes expressing BmOR-1 + BmG\textsubscript{q}, or BmOR-1 and BmG\textsubscript{q} separately. Bombykol (Left) or bombykal (Right) (100 \mu M) was applied at the time indicated by arrowheads. Data ± SE (n = 3–4). (d) Dose-dependent increases in amplitudes of bombykol-induced currents. Current recordings at different doses were measured on different oocytes. Currents of oocytes showing responses were averaged for each concentration: 30 \mu M, n = 3; 50 \mu M, n = 3; 100 \mu M, n = 9.

At the fifth pulse, 40 \mu l of concentrated stock-solution of bombykol or bombykal was applied to the recording bath (160-\mu l volume) to give the indicated concentrations.

**Generation of BmOR-1 Recombinant Baculovirus.** The entire protein-coding sequence of BmOR-1 was subcloned into the transfer vector pVL1393 to create pVL-BmOR-1, which was designed to express the native BmOR-1 protein. Spodoptera frugiperda IPLB-SF21-AE (SF21) cells were cotransfected with 2 \mu g of resultant transfer vector DNA and 1 \mu g of the linearized hybrid nuclear polyhedrosis virus (HyNPV) (28) genomic DNA, which is a host range-expanded Autographa californica NPV by using a lipofection reagent (Invitrogen). The recombinant baculovirus was plaque-purified and amplified to titers of 1 x 10\textsuperscript{7} plaque-forming units/ml. The recombinant baculovirus resulting from pVL-BmOR-1 was named HyBmOR-1.

**Baculovirus-Infected Female Moths.** At 4 days before eclosion, 50 \mu l of virus solution containing 5 x 10\textsuperscript{5} plaque-forming units of the recombinant or WT baculovirus was percutaneously injected into the fifth abdominal segment of female pupae by using a 1-ml plastic syringe with a 26-gauge needle. Inoculated pupae were reared at 25°C until eclosion.

**Electroantennogram (EAG) Recording.** Antennae of the day-0 adult moths were excised at the base, and a few segments at the tip were clipped off. Antennae were immediately used for EAG by attaching an indifferent electrode to the tip of the antenna and a recording electrode to the base of the antenna. Electrodes were filled with 0.1 M NaCl, and signals from a DC preamplifier (MEZ-7101, Nihon Kohden) were plotted on a chart recorder (WX1200, Graphtech). Bombykol or bombykal (40 \mu l) in 1% dimethyl sulfoxide (DMSO) at 100 ng/\mu l were loaded on 1.2 cm\textsuperscript{2}-filter paper (Advantec no. 2, Toyoroshi, Tokyo) and inserted into a plastic pipette tip. A charcoal-purified airstream was passed through the pipette tip, through Tygon tubing (inner diameter, 4 mm), and directed on the antenna from a distance of ∼5 mm. The airflow rate was adjusted at 350 ml/min, unless otherwise mentioned, and the stimulus was applied by manually opening or closing valves.

**Results and Discussion**

We hypothesized that the bombykol receptor is a male antennae-specific OR because bombykol elicits electrophysiological signals only in the male moth antennae. Thus, we used the differential cloning strategy to identify the bombykol receptor. A CDNA clone (designated BmOR-1) encoding a 430-aa protein with seven putative transmembrane domains that are characteristic of the G protein-coupled receptor superfamily (Fig. 1a) was isolated through differential screening of a CDNA library of a B. mori male antennae. The amino acid sequence deduced from the BmOR-1 clone showed significant sequence similarity (up to 42%) with ORs in insect species such as Drosophila melanogaster (6–9), Anopheles gambiae (10–12), H. virescens (13, 14), and D. simulans (29). BmOR-1 exhibited the closest proximity to a putative OR in H. virescens, HR13 (14), according to phylogenetic analysis (Fig. 1a Lower). The BmOR-1 gene consists of eight exons and seven introns in 9,870 bp of genomic DNA (Fig. 1b). The genomic DNA sequence was screened against a bacterial artificial chromosome library from B. mori (22) and mapped to the Z chromosome, one of the two sex chromosomes (Z and W). B. mori males are the homogametic sex, ZZ, whereas the females are the heterogametic sex, ZW (30). Southern blot analysis showed that BmOR-1 is a single copy gene in the genome (data not shown). In addition to BmOR-1, we cloned the BmOR-2 gene by using the known insect ORs with a homology-based method. BmOR-2 shows 65–87% amino acid identity with ORs that belong to the insect Or83b gene family (31) (Fig. 1a).

RT-PCR experiments demonstrated that BmOR-1 is expressed only in the antennae of male B. mori (Fig. 2a); the BmOR-1 gene transcript was first detected 4 days before eclosion, increased during the late pupal stages, and was continu-
localized to the specific cells in the epithelium of the male antennae on the side carrying chemosensory hairs (Fig. 2c, Left), and no hybridization signal was detected in the female antennae (Fig. 2c, Right). Distribution patterns and density of the labeled cells in the male antennae (Fig. 2d) were similar to those in pheromone-sensitive long sensillum trichodea (32). Furthermore, when viewed in 2-μm plastic sections, BmOR-1-reactive cells were slightly ellipsoidal and were located at 5–10 μm beneath the antennal cuticle (Fig. 2e) where OR neurons reside. No labeling was observed with sense RNA probes (data not shown).

The developmental expression profile of PBP, which is thought to be involved in pheromone detection (33, 34), was similar to that of BmOR-1 (Fig. 2b, Middle), except that PBP was also detected, although at lower levels, in the antennae of female adults and pupae (data not shown). Double labeling of the male adult antennae with BmOR-1 and PBP antisense RNAs demonstrated that BmOR-1-labeled cells (red) were localized on the sensory epithelium side and surrounded by the PBP-labeled cells (green) in both longitudinal and cross sections (Fig. 1f). These results are consistent with the previous observation that PBP mRNA was expressed in supporting cells that surround the pheromone receptor neurons (32). These results suggest that BmOR-1 is expressed exclusively in the pheromone receptor neurons in the antennae of male moths.

We hypothesized that BmOR-1 is involved in the detection of the sex pheromone in the male antennae and tested the ability of BmOR-1 to recognize bombykol by using two-electrode voltage clamp recordings of Xenopus oocytes with depolarizing step pulses from a holding potential of −80 mV. Robust Ca2+-dependent Cl− currents were observed upon stimulation with bombykol, but not with bombykal, when BmOR-1 was coexpressed with BmGq (G protein α-subunits in B. mori belonging to the Gαq family) (Fig. 3). Oocytes injected with cRNA of BmOR-1 or BmGq alone did not respond to bombykol (Fig. 3c). Additionally, we tested ligand specificity of BmOR-1 by using 41 odorants that have previously been reported to elicit electronic responses in B. mori adult antennae (35). However, none of the odorants tested at 100 μM produced any response to the oocytes coexpressing BmOR-1 with BmGq and showing response to bombykol (data not shown). These results demonstrate that BmOR-1 specifically recognizes bombykol, and subsequently couples to BmGq, resulting in activation of Ca2+-dependent Cl− channels in oocytes. Dose-dependent response curves (Fig. 3d) show that relatively high concentrations of bombykol are necessary to fully activate BmOR-1 expressed in Xenopus oocytes, probably because of the low level of BmOR-1 expression and inefficient signal transduction mediated by BmGq in oocytes. The EC50 value of bombykol for BmOR-1 of 34 μM was ≈14-fold lower than that of a ligand for Drosophila Or43a, a conventional odorant receptor (36). Responses were observed in 10–15% of the injected oocytes, but preliminary results suggest that coexpression with BmOR-2 improved the yield of oocytes responding to bombykol by greatly enhancing membrane translocation of BmOR-1 (T. Nakagawa and K.T., unpublished results). Nonetheless, these results provide evidence that BmOR-1 functions as a bombykol receptor in a heterologous cell system.

A search of the Silkworm Genome Research Program database (http://kaikoblast dna affrc go.jp) for sequences similar to BmOR-1 and other known insect OR sequences identified an additional 29 putative OR gene sequences. RT-PCR experiments revealed that 23 of these sequences were expressed in the antennae. Two of these were expressed specifically in the male antennae but not in the female antennae (data not shown). None of these receptors showed a response to bombykol in the oocyte expression system (data not shown), suggesting that BmOR-1 is most likely a single receptor for bombykol.
We next examined whether the bombykol response observed in the male antennae could be reconstituted in the female antennae by ectopically expressing BmOR-1 by using recombinant baculovirus carrying BmOR-1-encoding sequence (HyBmOR-1). When B. mori female pupae were infected with HyBmOR-1 4 days before eclosion, BmOR-1 transcripts were detected in the antennae of the infected day-0 female adult moths (Fig. 4a). Next, we measured electrophysiological responses of the antennae of female moths ectopically expressing BmOR-1 when exposed to the sex pheromone. We recorded EAG responses of the antennae under an air stream containing bombykol or bombykal (4 μg of each on a filter paper) at a flow rate of 350 ml/min. HyBmOR-1-infected female antennae responded to bombykol, but not to bombykal (P < 0.05 by Scheffe’s test, Fig. 4b and c), indicating the strong specificity for bombykol. None of WT host range-expanded baculovirus (HyNPV)-infected female antennae responded to bombykol, but not to bombykal, indicating that BmOR-1 functions as a highly specific receptor for bombykol in the silkmoth antennae.

The discovery of BmOR-1 as an insect sex pheromone receptor sheds light on mechanisms underlying highly sensitive and specific detection of sex pheromones and subsequent signal integration by the brain. Identification and functional analyses of sex pheromone receptors from other insect species will provide insight into the molecular evolution of species-specific pheromones and their receptors, as well as contribute to the development of unique methodologies for controlling plant and after harvest pests.


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