Olfactory Receptor Neurons from Antennae of Developing Male Manduca sexta Respond to Components of the Species-specific Sex Pheromone in vitro

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Male-specific olfactory receptor neurons, dissociated from developing antennae of the moth Manduca sexta and grown in long-term primary cell culture, can respond to at least one component of the female moth’s sex-pheromone blend with the opening of a nonspecific cation channel. This response does not require the coapplication of pheromone-binding protein.

Adult male sphinx moths, Manduca sexta, locate mating partners by detecting and orienting to the female’s species-specific sex-pheromone blend. The female moth releases this volatile chemical signal from a gland at the tip of her abdomen (Eaton, 1986; Itagaki and Conner, 1988). Organic-solvent extracts of this gland contain a characteristic mixture of 12 C16 and C18 aldehydes, including the predominant and most potent component, (E,Z)-10,12-hexadecadienal or bombykal (Starratt et al., 1979; Christensen et al., 1989; Kaissling et al., 1989; Tumlinson et al., 1989).

Male Manduca sexta respond to stimulation with bombykal and at least two other pheromone components via specialized olfactory receptor neurons (ORNs) innervating long, sexually dimorphic sensory hairs (sensilla trichodea) on the antennae (Sanes and Hildebrand, 1976a,b; Schweitzer et al., 1976; Kaissling et al., 1989; Keil, 1989). The pheromone components are assumed to bind to presumptive receptors in the dendritic membranes of the ORNs and thus to initiate a series of transduction processes. These processes finally lead to depolarizing receptor potentials, which initiate action potentials that encode information about the quality, quantity, and temporal pattern of the odor and carry it to the brain (Kaissling and Thorson, 1980; Kaissling, 1987). “Tip recordings” from cut trichoid sensilla on moth antennae have shown that, after pheromonal stimulation, odor-dependent potential changes occur (Kaissling, 1987). These extracellularly measured receptor potentials rise rapidly until they reach a plateau and then repolarize gradually. The rapid rise is accompanied by a decrease in resistance and by a phasic, phasic-tonic, or tonic spiking response (Schneider, 1962; Schneider and Boeckh, 1962; Zack-Strausfeld, 1979; Kaissling and Thorson, 1980; Kaissling, 1986, 1987; Zack-Strausfeld and Kaissling, 1986; Kaissling et al., 1989). In response to strong pheromonal stimuli, both the extracellularly recorded receptor potential and the impulse frequency decrease as a result of adaptation (Zack-Strausfeld, 1979; Zack-Strausfeld and Kaissling, 1986, 1987).

The transduction processes that underlie these adapting receptor potentials in moth sensilla are assumed to occur in the dendrites of the ORNs, in an unusual ionic environment. Within the hollow, hairlike cuticular sensillum, the dendrites are bathed by a receptor lymph containing circa 200 mM K+, circa 20 mM Na+, and presumably several micromolar Ca2+ (Keil, 1984; Grütert, 1985; Kaissling, 1986) as well as abundant soluble proteins. The pheromone-binding proteins (PBPs) (Vogt and Riddiford, 1981; Kaissling et al., 1985; Vogt et al., 1985; Klein, 1987; Gygöry et al., 1988; Raming et al., 1989, 1990). It has been proposed that these PBPs (in M. sexta, ca. 16 kDa; Gygöry et al., 1988) carry the lipophilic pheromone components through the aqueous receptor lymph and may be required for the recognition of the pheromone components by presumptive receptors in the dendritic membranes of the ORNs (Vogt et al., 1985; Vogt, 1987; van den Berg and Ziegelberger, 1991). Alternatively, the PBP may scavenge, and thus rapidly inactivate, the pheromone components after their interactions with receptor sites (Kaissling, 1986).

Because the ORNs are not readily accessible to intracellular or patch clamp recording techniques in situ (Zufall and Hatt, 1991), we developed a primary-cell-culture system for studies of the mechanisms underlying the responses of male-specific ORNs to pheromone (Stengl and Hildebrand, 1990). Cultures are derived from antennae of stage 3 male M. sexta pupae, within 2 d after the mitotic birth of the ORNs and their associated cells.

Among the diverse cells found in these cultures, one particular type with a 5 μm soma and fine processes has been correlated morphologically and immunocytochemically with ORNs in vivo (Hishinuma et al., 1988a,b; Stengl and Hildebrand, 1990). In situ morphological, immunocytochemical, and physiological studies indicate that about 35% of cells of this type are recognized by the monoclonal “male-olfactory-specific antibody”...
(MOSA) and are responsive to pheromone components (Sanes and Hildebrand, 1976a,b; Hishinuma et al., 1988a,b; Kassilling et al., 1989; Lee and Straussfeld, 1990). Patch-clamp studies show further that ORNs differentiate physiologically in vitro and acquire various types of ion channels in their soma membranes. After 3 weeks in culture, the ORNs express at least one type of Na+ channel and at least three types of K+ channels (Ziffall et al., 1991b).

Here we demonstrate that cultured ORNs can respond specifically to extracts of the female sex-pheromone gland or to the synthetic pheromone component bombykal with opening of a nonspecific cation channel. Furthermore, addition of PBP is not required for these responses.

Preliminary accounts of some of this work have been presented elsewhere (Stengl et al., 1989, 1991).

Materials and Methods

Unless otherwise specified, all culture media were purchased from Gibco (Grand Island, NY), and all chemicals and biochemicals, from Sigma Chemical Co. (St. Louis, MO).

Animals. Manduca sexta (Lepidoptera: Sphingidae) were reared from eggs on artificial diet (modified from Bell and Joachim, 1976) on a long-day photoperiod regimen (17 hr light/7 hr dark) at 25-26°C and 50-60% relative humidity. Pupae were staged as previously described (Sanes and Hildebrand, 1976a; Tolbert et al., 1983). They were usually selected for dissection between 2:00 and 5:00 AZT (Arbitrary Zeitgeber Time, with lights on at 0:00 AZT) and anesthetized by chilling on ice for 10-15 min before dissection of the antennal flagellum.

Cell cultures. A detailed description of the culture techniques has been reported elsewhere (Stengl and Hildebrand, 1990). Briefly, antennal flagella from male M. sexta pupae (stage 3 of the 18 stages of adult development) were disrupted by a combination of mechanical and enzymatic treatments. The dispersed cells were plated in concanavalin A-coated or uncoated Falcon plastic dishes, in Leibowitz L15 medium, supplemented with 5% fetal bovine serum (HyClone) and 20-hydroxyecdysone (ca. 1 µg/ml) or condition medium (supernatant fluid from cultures of a non-neural M. sexta cell line, generously provided by Drs. J. Hayashi and L. Oland, of the ARL Division of Neurobiology, University of Arizona, or extracellular fluid from antennae) (Stengl and Hildebrand, 1990). The cultures were maintained for 2-4 weeks at about 30°C at high humidity in an incubator.

Immunocytochemistry. ORNs from which recordings had been obtained were stained for immunoreactivity to the male-olfactory-specific antibody (MOSA, Hishinuma et al., 1988a,b), as described previously (Stengl and Hildebrand, 1990).

Patch-clamp technique and data analysis. Patch-clamp recording experiments closely followed the methods described by Hamill et al. (1981). Patch pipettes were made from borosilicate glass capillaries (World Precision Instruments, GC. 150 T10; or Clark Electromedical Instruments, Reading, UK) with a two-stage electrode puller (DMZ, Zeitz Instruments, Augsburg, Germany) or a Sutter Instruments micropipette puller (model P80/PC). The pipettes were coated with Sylgard (Dow Corning, Midland, MI) and then fire polished. The tip resistance was 6-12 MΩ when the electrodes were filled with physiological saline solution. The cells were viewed at 320× magnification with a Zeiss Axiovert 10 inverted microscope equipped with phase-contrast optics or at 1000× in a confocal laser-scanning microscope (TCS-4D, Leica) using a 63× oil immersion objective with phase-contrast or Hoffman modulation contrast optics. After formation of a seal between the pipette and the cell membrane, the electrode capacitance was compensated. During application of the pheromone, the membrane patches were kept at 0 mV holding potential (at the cells’ resting potential) in the cell-attached configuration.

Single-channel currents were measured at room temperature with an Axopatch 70 (Axon Instruments electronic, Darmstadt, Germany) or an Axopatch-1C patch-clamp amplifier (Axon Instruments Co., Burlingame, CA). The signals were recorded on videotape with a modified Sony PCM-501-ES video recorder or with an instrumentation FM tape recorder (A. R. Vetter Co., Rebersburg, PA) or acquired on line with an 80386-based microcomputer (Dell Computer Corp., Austin, TX) using pClamp software (Axon Instruments Co.). The data were analyzed with pClamp software (Axon Instruments) or with custom programs (Dudel and Franke, 1987). Single-channel currents were low-pass filtered at 2-5 kHz with an eight pole Bessel filter and digitally sampled at 25 kHz using a Hewlett-Packard P 9802 computer equipped with a Hewlett-Packard Multiprogrammer II interface. More detailed descriptions of the methods used can be found elsewhere (Dudel and Franke, 1987; Ziffall et al., 1991b).

Solutions. For recordings, cells were kept in “extracellular saline solution” containing (in mM) 156 NaCl, 4 KCl, 6 CaCl2, 5 glucose, and 10 HEPES (adjusted to pH 7.1, with NaOH). The “intracellular saline solution” used to fill pipettes for the cell-attached recordings contained (in mM) 150 KCl, 5 NaCl, 2 MgCl2, 0.1 EGTA (pCa = 7), and 10 HEPES (adjusted to 7.2 with KOH). The final K+ concentration was about 154 mM.

Extraction of pheromone glands and delivery of pheromonal stimuli. Sex-pheromone extracts were prepared by dipping female abdominal tips, isolated from adult virgins, in dimethyl sulfoxide (DMSO), dimethyl formamide (DMF), ethanol, or n-hexane. Freshly prepared for each experimental session, the “stock extract” consisted of 100 µl of solvent in which five female abdominal tips had been dipped for 7 min each (Tumlinson et al., 1989). These stock solutions were diluted 1:105 to 1:108 in 1 ml of extracellular saline solution. The hexane stock solution (which resulted in the purest pheromone extracts) was mixed with 10% DMSO, to facilitate aqueous solubility, before dilution in extracellular saline. The concentration of pheromone in dilutions of the stock extract was estimated (as “bombykal equivalents”) by means of electronicrogram (EAG) recordings, obtained with male antennae by established procedures (Christensen et al., 1989).

Detected sensory hairs were isolated from adult females (sensillum campaniforme) with a tip opening of <10 µm, driven by a Picospritzer (General Valve Corp., Fairfield, NJ). The pheromone was puffed onto the cell’s soma (≥20 µm from the pipette tip) before or after formation of a seal between the pipette and the cell membrane, respectively, to allow or prevent direct access of the pheromone to the membrane patch. The final concentration of pheromone in the gland extracts used ranged between 3 fg (10-12 mol) and 30 pg (10-11 mol) bombykal alone (control) and then tested with the pheromone solutions in the same solvent. Because the pheromone adsorbed to plastic or glass surfaces, the first application of pheromone contaminated the preparation dish. Once pheromone had been introduced into the dish, its physiologic effect (thus possibly its effective concentration) could be reduced by addition of bovine serum albumin (BSA) and extensive washing, but the physiologic responses of the cells indicated that the pheromone could not be completely removed from a contaminated dish.

Preparation of pheromone-binding protein. PBP was extracted by previous methods (Kaplow et al., 1987; Voss et al., 1987; Gjory et al., 1988). Antennal flagella (n = 44) of adult male M. sexta (1 d posteclosion) were collected over solid CO2. The frozen flagella were vortexed in pulverized solid CO2 to break off the sensory hairs. After the solid CO2 had fully sublimated away, the flagellar shafts were removed. The detached sensory hairs and scales were collected with diethyl ether. After evaporation of the ether, the fractions were taken up in 1 ml of buffer solution containing 250 mM sucrose, 2 mM EDTA, and 50 mM Tris (pH 8) and homogenized on ice. The homogenates were centrifuged at 2000 rpm for 5 min in a Sorvall RC-5B centrifuge (Du Pont Instruments, Wilmington, DE), and the supernatant fractions for 10 min at 9000 rpm. The resulting supernatant fractions were centrifuged at 27,000 rpm for 1 hr. The membrane and cytoplasmic fractions were taken up in buffer, and the protein concentrations were determined with a modified Lowry assay (Peterson, 1977; data not shown). The fractions were electrophoresed on SDS-PAGE gels or on native 12% acrylamide gels (Vogt and Riddiford, 1981) and stained with Coomassie blue. The cytoplasmatic fraction of the sensilla and scales contained two bands with a molecular mass between 14 and 21 kDa, corresponding to the published molecular mass of PBP (18 kDa) This fraction was used as PBP in physiological experiments.

Results

To test the ability of cultured ORNs to respond specifically to pheromone, we obtained patch-clamp recordings from cells in antennal-cell cultures 2-4 weeks old. Except for control recordings (see Control experiments, below) on various types of an-
teneral cells, only ORN-like cells (i.e., with a 5 μm soma and fine processes) were chosen for these experiments (Stengl and Hildebrand, 1990). The number of ORNs from which recordings were obtained successfully is given as n.

To obtain responses from intact cells without dialysis of cytoplasmic components such as second messenger systems, we recorded in the cell-attached mode from the cell soma. The 

**Control experiments**

Cultured antennal cells were tested for responses to stimulation with the solvents alone. We also sought to determine whether the responses to pheromone were specific to pheromone components. Furthermore, we tested whether the cells respond at a specific time in vitro and whether the responses are specific to certain types of cells.

In more than 50 cell-attached patch-clamp experiments, cultured ORNs were stimulated first with the solvents (DMSO, DMF, ethanol, hexane–DMSO) alone, before application of the pheromone sample. In none of these control experiments were responses elicited by the solvents alone. Cells that responded to pheromone did not respond to 1 μM citral (n = 8), a potent plant-derived odorant. These findings suggested that the responses might be cell-type-specific.

None of the 15 nonreceptor cells (judged by morphological criteria) responded to pheromonal stimulation. To demonstrate that the cells we tested were male-specific ORNs, we processed male ORNs for MOSA immunocytochemistry. In one of these control experiments were responses elicited by the solvents alone. Cells that responded to pheromone did not respond to 1 μM citral (n = 8), a potent plant-derived odorant. These findings suggested that the responses might be cell-type-specific.

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Cell-attached patch-clamp recording of a pheromone-dependent cation channel in a cultured ORN. A pheromone-dependent, nonspecific cation channel opened in cell-attached patches with intracellular solution containing 150 mM K+ and 10^(-4) M Ca^2+ in the patch pipette. In A, single-channel openings of the cation channel are shown at different cell potentials. The channel opened in bursts and had amplitude substates C, closed state; O, open state. B, The I-V curve shows that the channel reversed around 0 mV. Its conductance was nonlinear, with about 20 pS estimated at 25 mV and 50 pS at -100 mV. The channel did not discriminate among K+, Na+, and Cs+. The cell's membrane potential was estimated in cell-attached recordings with symmetrical K+ concentrations from the reversal potential of the previously described delayed-rectifier channels (Zufall et al., 1991b) and confirmed in the current-clamp mode after establishing a whole-cell configuration. The Z-C curve was obtained from amplitude histograms at different pipette potentials, taking only the main amplitude state into account.

This channel opened in bursts, with rapid flickers between open and closed states, and showed no amplitude substates (Fig. 1D, squares).

After stimulation with lower concentrations of pheromone (pg-fg/ml), in 54% (76 of 48) of the pheromone-specific responses, openings of nonspecific cation channels could be noted in single-channel records (Fig. 2A). This type of channel was never observed in cell-attached experiments in the absence of applied pheromone (n > 200), whereas K+ channels resembling the delayed rectifier could be observed regularly. The current through the cation channels reversed around 0 mV cell potential (i.e., between -10 mV and 20 mV, no channel activity was observed) in the cell-attached configuration with Ca^2+-buffered intracellular saline solutions containing different principal cations (150 mM K+, Na+, or Cs+) in the patch pipette (Fig. 2B). This indicated that the channels do not discriminate among K+, Na+, and Cs+. The conductance of the channels was nonlinear: circa 50 pS at -100 mV, 36 pS at -25 mV, and 18 pS at 25 mV. Amplitude histograms at -70 mV holding potential revealed additional smaller-amplitude substates (Fig. 3). The burst length of channel openings (of the largest-amplitude state, at -70 mV holding potential) can be fitted by one exponential with a time constant of about 43 msec (Fig. 4A). The distribution of open states of the largest-amplitude state can be described by two exponentials, with time constants of about 0.6 msec and 2.3 msec at -70 mV holding potential (Fig. 4B). The cation channels opened readily and continuously after patch excision even in the absence of pheromone but were never observed in the cell-attached configuration. The conductance of the excision-activated channels was also nonlinear, but showed less inward rectification than the pheromone-dependent cation channels that were observed in the cell-attached configuration. The conductance of the excision-activated channels was also nonlinear, but showed less inward rectification than the pheromone-dependent cation channels that were observed in the cell-attached configuration.

To determine whether the pheromone opened the pheromone-dependent cation channels via mechanisms confined to the membrane, we stimulated outside-out patches with pheromone. Patch excision to the outside-out configuration, even in the absence of pheromone, activated cation channels. These cation channels reversed around 0 mV, did not discriminate among Na+, K+, and Cs+; and expressed conductance substates (Fig. 5). Thus, these channels closely resembled the pheromone-dependent cation channels that were observed in the cell-attached configuration.
Figure 4. The burst length and the mean open-time durations of pheromone-dependent cation channels in cultured ORNs. A, The burst length of the nonspecific cation channel (from the same recording as in Fig. 4) at -70 mV holding potential can be fitted by one exponential with a time constant of 43 msec. B, The distribution of open times of the nonspecific cation channel at the same potential (same data as in Fig. 2): events with amplitudes between -8 pA and -10 pA. The open times (τ) are fitted with the sum of two exponentials. The mean open times are 0.6 msec and 2.3 msec.

The channel was unaffected by \(10^{-6}\) m TTX, 20 mM 4-aminopyridine, 10 mM ATP, 1 mM bromo-cGMP, 1 mM bromo-cAMP, 100 μM amiloride, or Ca
+2-channel blockers including Co
+2, Sr
+2, and Ni
+2 (applied outside). The channels were blocked in most cases by 20 mM tetraethylammonium chloride (applied to outside-out patches; data not shown). Replacement of Cl
- by aspartate, glutamate, or acetate did not affect the currents through the cation channels, but no currents through cation channels were observed if the cations were replaced by choline.

In at least 17% (8/48) of the pheromone-specific responses, the previously described cGMP-blockable delayed-rectifier K
+ channels (Zufall et al., 1991 b) exhibited increased probability of opening after stimulation with pheromone (Fig. 6) and closed after prolonged exposure to pheromone. Because the records also contained responses of several cation channels or were too short, we did not analyze them further. In at least 6% (3 of 48) of all pheromone-dependent responses, channels opened (Fig. 1B, squares) that resembled the previously described Ca
+2-dependent K
+ channels (Zufall et al., 1991 b) in kinetics and amplitude (data not shown).

Coapplication of PBP (Fig. 7A) was not required to obtain specific pheromone-dependent responses. After 10-60 min of incubation with high concentrations of pheromone (ca. 30 pg/ml), the pheromone-dependent cation channels and the delayed-rectifier channels closed. If PBP or BSA was then applied (at a concentration of about 10 mM), channels with a reversal potential of about 0 mV opened (Fig. 7B). Channel activity that reversed around 0 mV could be suppressed again with application of more pheromone, or thereafter elicited again with application of more PBP (n = 3).

Figure 5. An outside-out patch-clamp recording of a cation channel activated by patch excision from a cultured ORN. Excision of a membrane patch to the outside-out configuration with 156 mM NaCl outside and intracellular solution with 156 mM KCl inside opened a nonspecific cation channel. Single-channel openings at different holding potentials (A) show that the channel opened in bursts with amplitude substates (c, closed state; o, open state of the channel). B, The channel reversed around 0 mV, with nonlinear conductance of about 37 pS at +30 mV and about 53 pS at -100 mV. The channel discriminated little between Na
+, K
+, and Cs
+. Thus, this excision-activated cation channel resembles the pheromone-dependent cation channel (Fig. 2).
membranes at specific times in their development in vitro (Zufall 1991b), we assumed that pheromone-dependent ion channels would also be expressed in their somata, before becoming localized to the dendritic sites of their final deployment. This assumption was supported by findings in other neuronal culture systems in which Ca$$^{2+}$$ or Na$$^{+}$$ channels are first expressed in the soma membranes and then become localized to the dendritic sites of their final deployment.

Female pheromone-gland extracts were chosen as the main olfactory stimuli to maximize the stimulation of all pheromone-sensitive ORNs. From in vivo studies, it is known that there are at least three types of pheromone-sensitive antenna ORNs, which respond to different components of the pheromone-gland extract (Kaissling et al., 1989). To assess the specificity of the responses to the gland extract, we compared different solvent extracts, and in some of our experiments the synthetic pheromone component bombykal was employed. The similarities in the responses to bombykal and to the pheromone-gland extracts suggest that the responses could have been due mainly or entirely to bombykal in the gland extracts. In support of this interpretation, solvents alone did not elicit any response, the pheromone concentrations used were in the physiological range (see Materials and Methods), and cells that responded to pheromone did not respond to the plant odorant citral. Moreover, the responses were specific to the ORN cell type and could be elicited by the odor-sensitive neurons (recognized on the basis of their 5 1/2 mm somata and their fine bipolar processes in vitro; Stengl and Hildebrand, 1990). Finally, we showed that cells that responded with opening of cation channels were MOSA immunoreactive. Because MOSA immunoreactivity is characteristic of male-spe-
specific ORNs in situ (Hishinuma et al., 1988a,b), it provided further confirmation of the specificity of the responses to pheromone. Although dose–response curves usually provide further indication of response specificity, they were not determined in our experiments because we could not ascertain accurately the concentration and arrival time of the pheromonal stimulus at the cell. In a more general way, however, concentration dependence was exhibited: at higher concentrations of pheromone (picogram of bombykal), large inward currents obscured single-channel events. Single-channel events could be resolved only after reduction of the pheromone concentration by extensive washing or addition of BSA to the medium. Even before application of the pulse of pheromone solution, diffusion of the pheromone out of the puffer pipette appeared to be sufficient to open the cation channel in cell-attached recordings. This extraordinary sensitivity is consistent with possible involvement of second-messenger cascades in the opening of the pheromone-dependent cation channels. The relatively long response delays (from <100 msec up to several seconds) might also be accounted for by interposed second-messenger actions. Because the cultured ORNs responded to the pheromone, however, even when formation of the seal preceded delivery of the pheromonal stimulus, second-messenger control of the cation channel appears to be likely.

Considerable evidence for involvement of second-messenger systems in olfactory transduction, in vertebrates as well as in invertebrates, is accumulating (Pace et al., 1985; Nakamura and Gold, 1987; Breer et al., 1988, 1990; Firestein and Shepherd, 1989; Firestein and Werblin, 1989; Boeckhoff et al., 1990; Zufall and Hatt, 1991). After exposure to pheromone, insect ORNs exhibit relatively long-lasting increases in cGMP (Ziegelberger et al., 1990). These increases in cGMP occur in the somata rather than the dendritic regions of the ORNs. G Protein-dependent increases of inositol 1,4,5-triphosphate (IP₃) occur in insect ORNs within milliseconds after pheromonal stimulation, but in contrast to the situation in vertebrates, no increases in Ca²⁺ could be detected (Breer et al., 1988, 1990; Boeckhoff et al., 1990; Ziegelberger et al., 1990). Thus, a G-protein–dependent phospholipase C (Nishizuka, 1984; Berridge, 1987), which might cause an IP₃-dependent rise in internal Ca²⁺, may play a role in the production of odor-dependent potential changes in insect ORNs. Furthermore, a guanylate cyclase or possibly a phosphodiesterase might be involved in the olfactory transduction mechanisms. The involvement of diffusible second messengers in olfactory transduction processes thus makes it likely that channels not only in the outer dendritic segment but also in the inner dendritic segment and the soma may underlie odor-dependent potential changes that influence the firing of action potentials in ORNs. Therefore, more than one type of second-messenger–mediated channel might be involved in the generation of the receptor potentials in ORNs.

The pheromone-dependent cation channel described here is a good candidate to be one of the second-messenger–gated channels involved in the mechanisms that underlie the depolarizing receptor potentials in pheromone-sensitive ORNs. Because this type of channel is not voltage gated and is observed only in cell-attached recordings after pheromonal stimulation, its activation apparently depends upon pheromone. The opening of this channel leads to inward currents carried by K⁺, Na⁺, and possibly also Ca²⁺ at negative membrane potentials. Thus, these cation channels depolarize the cell if opened at the cell's resting potential. The suggestion that the opening of cation channels might be one of the events that underlie the pheromone-dependent generator currents is consistent with previous observations that the depolarizing receptor potentials are accompanied by a decrease in membrane resistance (Kaisiling and Thorson, 1980; Kaissling, 1987). This suggestion is further supported by recent findings of pheromone-dependent, second-messenger–mediated (diacylglycerol, cGMP) cation channels on extruded dendrites of moth ORNs in situ and second-messenger–dependent (Ca²⁺, cGMP) cation channels on dendritic segments of ORNs in Antherea polyphemus (Zufall and Hatt, 1991; Zufall et al., 1991a). The pheromone-dependent cation channel of cultured ORNs in M. sexta has a conductance different from both cation channels observed in A. polyphemus. Current experiments examine whether the observed pheromone-dependent cation channel in M. sexta, which otherwise shares reversal potential, ion selectivity, and the property of expressing multiple conductance states with both of the cation channels observed in A. polyphemus, also depends on second messengers (Stengl et al., 1991, 1992).

The M. sexta cation channels activated during excision of membrane patches appear not to be blockable by amiloride or cGMP, both of which block the Ca²⁺-dependent cation channels activated by patch excision in A. polyphemus. It seems likely, however, that increased Ca²⁺ concentrations (via influx from the extracellular medium or damage to internal stores) during the process of patch excision activated these channels in both species. Ion channels activated by patch excision have also been found in other systems, but their functions and mechanisms of activation are unknown, although it is also assumed that Ca²⁺ and ATP may play a role in their activation (Yazejian and Byerly, 1989; McClintock and Aehe, 1990).

Whether the cation channels activated via patch excision belong to the same type of cation channel as the pheromone-dependent cation channels measured cell-attached remains to be demonstrated. Experiments in progress on M. sexta show that cultured ORNs are equipped with Ca²⁺–dependent cation channels as well as with protein kinase C–dependent cation channels that are affected differently by cGMP (Stengl and Hildebrand, 1991; Stengl et al., 1991, in press). These data will be presented in a separate paper (M. Stengl and J. G. Hildebrand, unpublished observations). Further studies should ascertain whether the apparently different cation channels in both species of moths are truly different types of cation channels or whether they belong to the same channel type, which changes its properties via phosphorylation by protein kinases during prolonged pheromone exposure.

The opening of a channel that resembles in its kinetics and amplitude the previously described Ca²⁺–dependent K⁺ channels (Zufall et al., 1991b) implies a possible increase of at least 10-fold in internal Ca²⁺ after application of pheromone. The observed changes in the probability of opening of the nucleotide-sensitive delayed-rectifier K⁺ channels could possibly be accounted for by changes in the levels of ATP or cGMP or by changes in the cell's potential (Zufall et al., 1991b). Because increases in cGMP after pheromonal stimulation have been found in insect ORNs (Ziegelberger et al., 1990), the closing of the delayed-rectifier K⁺ channel after prolonged exposure to pheromone might be caused by cGMP. Because the current amplitude of the delayed rectifier did not decrease in all recordings, the pheromone-dependent increase in activity is probably not caused by a depolarization of the ORN.

In view of the fact that application of PBP was not necessary to obtain pheromone-dependent responses in cultured ORNs...
as well as in recent in situ experiments (Zufall and Hatz, 1991), it seems that the pheromone need not be presented in the form of a pheromone-PBP complex to a presumptive receptor site. We cannot exclude the possibility, however, that PBP might have been produced in vitro and therefore present in cultured ORNs in a membrane-bound form in which it was not washed away by superfusion. The PBP could be used interchangeably with BSA (which unspecifically binds lipophilic substances), apparently reducing pheromone concentrations and thus possibly preventing channel closure. Thus, our preliminary results are consistent with the possibility that PBP may also serve to scavenge the pheromone and hence to prevent adaptation due to overstimulation by pheromone. However, this study did not attempt to analyze the function of the PBP. The influence of the PBP on the time course and adaptation of the response to pheromone remains to be investigated further.

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