Two Preputial Gland-Secreted Pheromones Evoke Sexually Dimorphic Neural Pathways in the Mouse Vomeronasal System

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Hexadecanol (16OH) and hexadecyl acetate (16Ac) are two pheromones secreted in a large quantity by mouse preputial glands and act on male and female mice differentially. Yet the underlying molecular and cellular mechanisms remain to be elucidated. In this study, we examined the activation of vomeronasal sensory neurons (VSNs) by these two pheromones and mapped the downstream neural circuits that process and relay their chemosignals. Using the calcium imaging method and immunohistochemistry, we found that a small number of VSNs were activated by 16OH, 16Ac, or both in the male and female mice, most of which were located apically in the vomeronasal epithelium, and their numbers did not increase when the concentrations of 16OH and 16Ac were raised by 10,000-fold except that of female VSNs in response to 16OH. In the accessory olfactory bulb (AOB), the two pheromones evoked more c-Fos+ neurons in the anterior AOB (aAOB) than in the posterior AOB (pAOB); and the increases in the number of c-Fos+ neurons in both aAOB and pAOB were dose-dependent; and between sexes, the female AOB responded more strongly to 16OH than to 16Ac whereas the male AOB had the opposite response pattern. This sexual dimorphism was largely retained in the downstream brain regions, including the bed nucleus of the stria terminalis (BNST), the medial amygdaloid nucleus (MeA), the posteromedial cortical amygdaloid nucleus (PMCo), the medial preoptic area (MPA), and the ventromedial hypothalamic nucleus (VmH). Taken together, our data indicate that there is one V1r receptor each for 16OH, 16Ac, or both, and that activation of these receptors evokes sexually dimorphic neural circuits, directing different behavioral outputs and possibly modulating other pheromone-induced responses.

Keywords: hexadecanol, hexadecyl acetate, vomeronasal organ, calcium imaging, c-Fos, neural circuits
INTRODUCTION

The term “pheromone” has been used to describe the chemicals used for intra-species communications (Karlson and Luscher, 1959). These chemicals can convey information about conspecific members’ social status, sexual maturation, and receptiveness, and can also trigger sexual behaviors, pregnancy block, inter-sex or infant-directed aggression, and parental caring (Leinders-Zufall et al., 2000, 2004; Brennan and Zufall, 2006; Brechbühl et al., 2011; Liberles, 2014; Isogai et al., 2018; Mohrhardt et al., 2018). Since the first pheromones were identified in insects, many more have been found across the animal kingdom (Liberles, 2014). In mammals, rodents have been the best studied model animals, which emit pheromones through feces, urine, saliva, tear, and other bodily fluids that are produced by such glands as preputial and lacrimal glands (Novotny, 2003; Zhang et al., 2008; Brechbühl et al., 2011). However, the number of molecularly identified pheromones is still limited in mammals, including proteins and peptides, e.g., the major urinary proteins (MUPs) (Hurst et al., 2001), major histocompatibility complex (MHC) class I peptides (Sturm et al., 2013), exocrine gland-secreting peptides (ESPs) (Kimoto et al., 2005; Osakada et al., 2018), as well as more volatile compounds such as bile acids and steroids, e.g., 2,5-dimethylpyrazine, 3,4-dehydro-exo-brevicomin, 2,3-dehydro-exo-brevicomin, 2-heptanone, 6-hydroxy-6-methyl-3-heptanone (HMH), 2-sec-butyl-4,5-dihydrothiazole (SBT), αβ-farnesene, and sulfated sex hormones (Novotny et al., 1990, 1999; Brennan and Zufall, 2006; Haga-Yamanaka et al., 2014; Doyle et al., 2016). Naturally produced pheromones, however, are often mixtures, of which the ratios of these different components are also important to evoking the receivers’ behavioral responses. Identifying additional semiochemicals and their blends is indeed necessary to fully understand pheromone physiology.

The pheromones are detected mainly by the sensory neurons in the vomeronasal organs (VNO) that are located at the anterior bottom of murine nasal cavity. Unlike the olfactory epithelium of the main olfactory system (MOS), the vomeronasal epithelium (VSE) can be largely divided into apical and basal layers whereas the basal layer can be further divided into two sublayers (Munger et al., 2009; Touhara and Vosshall, 2006; Dulac and Wagner, 2006; Haga et al., 2010; Szymanski and Keller, 2014). Up till now, however, very few pheromones’ neural circuits have been mapped out (Dulac and Torello, 2003; Brennan and Zufall, 2006; Ishii et al., 2017; Wei et al., 2018). And even more complicated is that the accessory olfactory system (AOS) may also be plastic and can be modulated by learning and internal physiological states (Kaur et al., 2014; Xu et al., 2016). Thus, more thorough studies are needed to elucidate the AOS circuits beyond those merely eliciting the stereotyped responses.

Hexadecanol (16OH) and hexadecyl acetate (16Ac) are two pheromones initially identified from insects, and later found to be produced in a substantial quantity by murine preputial glands (McElfresh et al., 2001; Zhang et al., 2008). Male mice, which produce more of these two pheromones than females, show some attraction to them at low concentrations but display aversion to concentrated ones while female mice are indifferent to low concentrations but attractive to high concentrations of these two pheromones; and these sex-specific responses are VNO-dependent, and probably mediated by the AOS (Zhang et al., 2008). However, the exact molecular and cellular mechanisms underlying their detection in the VNO and information processing in the central circuits remain to be revealed. In this study, we utilized calcium imaging to identify the VSNs responsible for the 16OH and 16Ac detection, and c-Fos immunohistochemistry to identify activated neurons in the central nuclei including those in the AOB, BNST, PMCo, MeA, MPA, and VmH that make up the AOS neural circuits. Our data provide new insights into the underlying molecular and
FIGURE 1 | The pheromones evoke responses of intracellular Ca\(^{2+}\) elevation from single vomeronal sensory neurons. (A,B) Fluorescence images on gray and pseudocolor scales, respectively, acquired from a VNO slice loaded with fura-2-AM. The somata of the VSNs contained most of the calcium-sensitive dye. (C) Traces of three VSNs responsive to 100 \(\mu\)M 16OH and KCl (50 mM) but not to 100 \(\mu\)M 16Ac over a period of 8 min. (D) Left: Response profiles of three representative VSNs: cell 1 responded to 16OH alone, cell 2 to 16Ac alone, and cell 3 to both; Right: the corresponding Fura-2 ratio images of the three VSNs. KCl (50 mM) was used as a positive control. Scale bar: 20 \(\mu\)m.

FIGURE 2 | Male VSNs respond similarly to 16OH and 16OHAc at both low and high concentrations. (A) Comparison of the percentages of male VSNs responsive to 16OH and 16Ac at a low concentration (10 nM). Out of 1175 cells examined, 14 responded to 16OH, 11 to 16Ac, 7 to both 16OH and 16Ac; and no statistically significant difference was found among them (M-L-16OH vs. M-L-16Ac, \(t = 0.383\), \(p = 1.000\); M-L-16OH vs. M-L-16OH+M-L-16Ac, \(t = 1.338\), \(p = 1.000\); M-L-16Ac vs. M-L-16OH+M-L-16Ac, \(t = 0.956\), \(p = 1.000\)). (B) Comparison of the percentages of male VSNs responsive to 16OH and 16Ac at a high concentration (100 \(\mu\)M). Out of 988 cells examined, 17 responded to 16OH, 11 to 16Ac, 8 to both 16OH and 16Ac; and no statistically significant difference was found among them (M-H-16OH vs. M-H-16Ac, \(t = 1.1\), \(p = 1.000\); M-H-16OH vs. M-H-16OH+M-H-16Ac, \(t = 1.784\), \(p = 1.000\); M-H-16Ac vs. M-H-16OH+M-H-16Ac, \(t = 0.683\), \(p = 1.000\)). (C) Comparison analysis of the male VSNs responsive to the low and high concentrations of the pheromones. No significant difference was found between the responses to the low and high concentrations of the same pheromones in male mice (M-H-16OH vs. M-L-16OH, \(t = 0.953\), \(p = 1.000\); M-H-16Ac vs. M-L-16Ac, \(t = 0.236\), \(p = 1.000\); M-H-16OH+M-H-16Ac vs. M-L-16OH+M-L-16Ac, \(t = 0.508\), \(p = 1.000\)). All values are expressed as mean ± s.e.m.
Female VSNs respond differently to 16OH and 16Ac. (A) Comparison of the percentages of female VSNs responsive to 16OH and 16Ac at 10 nM. Out of 1168 cells examined, 23 responded to L-16OH, 10 to L-16Ac, and 6 to both. Significantly more cells responded to L-16OH than to both; and no significant differences in the numbers of cells responsive to L-16OH versus to L-16Ac or to L-16Ac versus to both were found (F-L-16OH vs. F-L-16Ac, t = 2.311, p = 0.341; F-L-16OH vs. F-L-16OH + F-L-16Ac, t = 3.116, p = 0.035; F-L-16Ac vs. F-L-16OH + F-L-16Ac, t = 0.805, p = 1.000). (B) Comparison of the percentages of female VSNs responsive to 16OH and 16Ac at 100 µM. Out of 964 cells examined, 41 responded to H-16OH, 17 to H-16Ac, and 16 to both. Cells responsive to H-16OH were significantly more than those to H-16Ac or both but no significant difference was found between cells responsive to H-16Ac versus both (F-H-16OH vs. F-H-16Ac, t = 4.573, p = 0.000; F-H-16OH vs. F-H-16OH + F-H-16Ac, t = 5.196, p = 0.000; F-H-16Ac vs. F-H-16OH + F-H-16Ac, t = 0.623, p = 1.000). (C) Comparative analysis of the female VSNs responsive to the same pheromones at different concentrations. More female VSNs responded to H-16OH than to L-16OH whereas no significant differences were found between cell numbers responsive to H- versus to L-16Ac or between those to both H-16OH and H-16Ac versus both L-16OH and L-16Ac (F-H-16OH vs. F-L-16OH, t = 4.062, p = 0.001; F-H-16Ac vs. F-L-16Ac, t = 1.800, p = 1.000; F-H-16OH + F-H-16Ac vs. F-L-16OH + F-L-16Ac, t = 1.982, p = 0.751). All values are expressed as mean ± s.e.m. ∗p < 0.05, ∗∗p < 0.01, ∗∗∗p < 0.001.

The differences in responsivity between male and female mice. (A) No significant differences were found between the numbers of male versus female VSNs responsive to L-16OH, L-16Ac, or both at a low concentration (10 nM) (M-L-16OH vs. F-L-16OH, t = 1.618, p = 0.109; M-L-16Ac vs. F-L-16Ac, t = 0.310, p = 0.757; M-L-16OH + M-L-16Ac vs. F-L-16OH + F-L-16Ac, t = 0.160, p = 0.873). (B) Significantly more female VSNs responded to H-16OH than male VSNs whereas no significant differences were found between the numbers of male and female responsive cells to H-16Ac or to both (100 µM) (M-H-16OH vs. F-H-16OH, t = 4.726, p = 0.000; M-H-16Ac vs. F-H-16Ac, t = 1.254, p = 0.213; M-H-16OH + M-H-16Ac vs. F-H-16OH + F-H-16Ac, t = 1.314, p = 0.192). All values are expressed as mean ± s.e.m. ∗∗∗p < 0.001.

Cellular mechanisms of the 16OH- and 16Ac-directed dimorphic behavioral outputs.

**MATERIALS AND METHODS**

**Animals**

All the mice used in the experiments are 8–10 weeks old CD-1 mice. A total of 118 mice were used in this study: 5 male and 5 female mice were used for the VNO calcium imaging experiment; 24 male and 24 female mice for the pheromone-induced VNO pS6 immunofluorescence experiment, and 30 male and 30 female mice for the brain c-Fos immunohistochemistry experiment. Male mice were housed individually whereas females were housed in groups of four with food and water available ad libitum. Female mice were examined to determine their estrous stages and only estrous mice were used in this study. Before exposure to the pheromones, subjects were housed individually under a reversed 14/10 h light/dark photoperiod (lights on at 7:00 pm) with food and water available ad libitum in a clean room and the experiments were carried out in the morning hours. All experiments with
FIGURE 5 | Most responsive neurons are apical VSNs. (A,B) Images of anti-pS6 and anti-Goα double immunostaining on male and female VNO sections, respectively, following the exposure to 100 µM 16OH, 100 µM 16Ac, urine (positive control), or double distilled water containing 0.01% DCM (ddH2O, negative control). (C,D) Comparative analysis of the double immunostaining data from both the male and female VNO sections, respectively, showed that significantly more pS6-positive neurons were located in the apical VSE (aVSE) than in the basal VSE (bVSE) upon stimulation of urine, 16OH, or 16Ac (aVSE vs. bVSE: male-urine, \( t = 6.350, p = 0.000 \); male-16OH, \( t = 3.275, p = 0.010 \); male-16Ac, \( t = 2.604, p = 0.029 \); female-urine, \( t = 4.462, p = 0.002 \); female-16OH, \( t = 6.464, p = 0.000 \); female-16Ac, \( t = 4.620, p = 0.001 \)). All values are expressed as mean ± s.e.m. *\( p < 0.05 \), **\( p < 0.01 \), ***\( p < 0.001 \).

the animals were approved by the Institutional Animal Care and Use Committees of both Zhejiang University (No. 14843) and Institute of Zoology, Chinese Academy of Sciences (IOZ 2015) and followed the NIH “Guide for the Care and Use of Laboratory Animals.”

Calcium Imaging
VNO Slice Preparation
Animals were decapitated following anesthesia, and the mandible bone was cut off with a pair of scissors to remove the lower jaw. The ridged upper palate tissue was peeled off to expose the nasal cavity. The anterior and posterior ends of the nasal septum were cut to extract the VNO-containing portion, which was immediately transferred to the ice-cold oxygenated mouse artificial cerebro-spinal fluid (95% O2/5% CO2; mACSF containing 125 mM NaCl, 2.5 mM KCl, 1 mM MgCl2, 1.25 mM NaH2PO4, 25 mM NaHCO3, 2 mM CaCl2, 10 mM D-Glucose, pH 7.4) (Brechbühl et al., 2011; Ma et al., 2011). The cartilaginous capsule of the VNO was then removed to gain access to the luminal surface of the sensory epithelium. The dissected VNO
FIGURE 6 | Continued
was embedded in 3% low-melting agar and sliced coronally into 200 µm-thick sections with a vibratome at a speed of 0.5 mm/s and an amplitude of 0.7 mm (Leinders-Zufall et al., 2000).

**Calcium Imaging**

A VNO tissue slice was loaded with 10 µM calcium-sensitive dye Fura-2-AM (F1201, Life Technology, United States) for 30–60 min. Calcium imaging was performed on a Nikon microscope equipped with 20×, 40× water immersion objectives to monitor the changes in intracellular Ca²⁺ concentrations over time. Cells were illuminated sequentially at 340/380 nm with a polychromator instrument and emission at 510 nm was recorded at a rate of 5 Hz. Changes in the intracellular emission ratio at 340 and 380 nm, i.e., ratio = F340/F380 nm, were monitored with Ratio Imaging software. Pheromones were delivered at a flow rate of 1 ml/min using a peristaltic pump. Near the end of each imaging session, 50 mM KCl in mACSF was applied to the VNO slice to check the viability and responsiveness of the cells and only the responsive VSNs were included in the post hoc data analyses. The interstimulus intervals were 4 min or longer to allow the recovery of the VSNs. Solutions of 16OH and 16Ac were prepared as stock solutions of 1 M by being dissolved in dimethyl sulfoxide (DMSO) and then diluted in mACSF to two concentrations used in this study: low (10 nM) and high (100 µM). Thus, these four solutions: 10 nM 16OH, 10 nM 16Ac, 100 µM 16OH, and 100 µM 16Ac were designated as L-16OH, L-16Ac, H-16OH, and H-16Ac, respectively (Leinders-Zufall et al., 2000; Isogai et al., 2011).

**Immunohistochemistry**

### Pheromone Treatment

To stimulate mice, 16OH and 16Ac stock solutions of 1 M were prepared in dichloromethane (DCM) and diluted to 100 µM in double distilled water for VNO experiment, or diluted to two concentrations: 10 nM and 100 µM in mineral oil for AOB and brain experiment; thus, the concentration of residual DCM in the pheromone solutions was 0.01% or less. Forty microliters of the pheromones or whole urine (positive control) or 0.01% DCM in double distilled water (negative control) for VNO staining, and pheromones or 0.01% DCM in mineral oil (negative control) for AOB and brain staining were added onto a piece of filter paper placed in a mould in the bedding of the subjects that had been previously housed individually for 2 days in a clean room. The treatment lasted for 45 min for the VNO experiments or 90 min for the AOB and brain experiments. Based on sex and treatment, the animals were divided into eight groups: male mice treated with urine (Male-urine), 0.01% DCM-containing double distilled water (Male-ddH₂O), 16OH (Male-16OH), and 16Ac (Male-16Ac); and female mice treated with urine (Female-urine), 0.01% DCM-containing double distilled water (Female-ddH₂O), 16OH (Female-16OH), and 16Ac (Female-16Ac) for the VNO experiments. And for the AOB and brain experiments, the animals were divided into 10 groups: male mice treated with 0.01% DCM-containing mineral oil (designated as M-control), L-16OH (M-L-16OH), L-16Ac (M-L-16Ac), H-16OH (M-H-16OH), or H-16Ac (M-H-16Ac); and female mice treated with 0.01% DCM-containing mineral oil (F-control), L-16OH (F-L-16OH), L-16Ac (F-L-16Ac), H-16OH (F-H-16OH), or H-16Ac (F-H-16Ac).

**Tissue Preparation**

After pheromone treatment, the mice were anesthetized with sodium pentobarbital (40 mg/kg) and transcardially perfused with PBS followed by 4% paraformaldehyde (PFA) in PBS. Then, the VNO, OB, and brain were dissected out, post-fixed overnight, and cryoprotected in 30% sucrose solution in PBS overnight. Coronal sections of the VNO (12 µm thick) and brain (40 µm thick) and sagittal sections of the OB (30 µm thick) were sliced and collected. The brain sections that contained the nuclei potentially involved in the processing of 16OH and 16Ac signals, including the BNST, MPA, anterior subdivision of the MeA, ventromedial hypothalamic nucleus (Vmh), postero medial cortical amygdaloid nucleus (PMCo), were collected (Supplementary Figure 1; Paxinos and Franklin, 2004). The intervals between sections from the brain and OB were 40 and 30 µm apart, respectively.

**Double Immunostaining With Anti-pS6 and Anti-Gαo Antibodies on the VNO Sections (Jiang et al., 2015)**

The VNO sections were washed with PBS to remove the embedding medium and blocked by incubating with the blocking buffer (10% normal goat serum, 2% bovine serum albumin in 0.5% Triton X-100/PBS solution) for 1 h. The anti-Gαo primary antibody (551, Medical and Biological Laboratories, Japan) was diluted at 1:2000 in the dilution buffer (2% normal goat serum, 2% BSA in 0.5% Triton X-100/PBS) and applied to the sections for 16 h, followed by the incubation with AF568-conjugated goat-anti-rabbit secondary antibody diluted at 1:500 (ab175471, Abcam, United Kingdom) for 1 h. The anti-pS6 antibody (4856S, Cell Signaling Technology, Japan) was conjugated with AF488.
TABLE 1

| Brain region | Male mice | Female mice |
|--------------|-----------|-------------|
|              | Control (n = 6) | L-16OH (n = 6) | L-16Ac (n = 6) | H-16Ac (n = 6) | Control (n = 6) | L-16OH (n = 6) | L-16Ac (n = 6) | H-16Ac (n = 6) |
| AOB          | 100.4 ± 5.9 | 52.9 ± 6.8 | 279.0 ± 14.5 | 529.0 ± 15.6 | 492.7 ± 17.3 | 297.0 ± 14.5 | 163.9 ± 14.5 | 279.0 ± 15.6 |
| MOB          | 110.7 ± 4.5 | 57.4 ± 6.3 | 287.0 ± 14.5 | 529.0 ± 15.6 | 492.7 ± 17.3 | 297.0 ± 14.5 | 163.9 ± 14.5 | 279.0 ± 15.6 |
| POM          | 110.7 ± 4.5 | 57.4 ± 6.3 | 287.0 ± 14.5 | 529.0 ± 15.6 | 492.7 ± 17.3 | 297.0 ± 14.5 | 163.9 ± 14.5 | 279.0 ± 15.6 |
| VmH          | 100.4 ± 5.9 | 52.9 ± 6.8 | 279.0 ± 14.5 | 529.0 ± 15.6 | 492.7 ± 17.3 | 297.0 ± 14.5 | 163.9 ± 14.5 | 279.0 ± 15.6 |

Data are expressed as the mean number of c-Fos+ cells ± s.e.m. in the AOB and other brain regions of the male and female mice after exposure to 16OH and 16Ac at 10 nM (L) or 100 nM (H) for 2 h. The aAOB was compared with pAOB in the same group, and groups of the same sex were compared with control as well. AOB, accessory olfactory bulb; aAOB, anterior AOB; pAOB, posterior AOB; BNST, bed nucleus of the stria terminalis; MPA, medial preoptic area; MeA, medial amygdaloid nucleus; VmH, ventromedial hypothalamic nucleus; PMCo, posteromedial cortical amygdaloid nucleus.

Statistical Analysis

Calcium imaging data were expressed as the mean ± s.e.m. The data distribution and homogeneity of variances were checked. The differences between groups with different sexes, pheromones, and concentrations were assessed by two-way ANOVA with sex as between-group factor and treatment as within-group factor. If significant interaction was found, a post hoc analysis with Bonferroni’s multiple comparison tests was performed to identify significant differences with PASW Statistics 18.0 software. c-Fos immunohistochemistry data using Zenon Rabbit IgG Labeling Kit (Z25302, Molecular Probes, United States) and applied to the sections for 1–2 h. Finally, the sections were washed with 1 × PBS and covered with the mounting medium containing DAPI.
FIGURE 7 | Quantitative analysis of c-Fos+ AOB neurons of the male and female mice after exposure to 16OH and 16Ac. (A) Significantly more activated neurons were found in both male and female aAOB and pAOB following stimulation by all these four stimuli: low or high 16OH or 16Ac in comparison with the control samples except the L-16Ac in female mice (M-L-16OH vs. M-control, \( t = 6.780, p = 0.000; M-H-16OH vs. M-control, t = 18.717, p = 0.000; M-L-16Ac vs. M-control, t = 10.148, p = 0.000; M-H-16Ac vs. M-control, t = 24.412, p = 0.000; F-L-16OH vs. F-control, t = 11.996, p = 0.000; F-H-16OH vs. F-control, \( t = 34.776, p = 0.000; F-L-16Ac vs. F-control, t = 2.834, p = 0.022; F-H-16Ac vs. F-control, t = 6.600, p = 0.000). H-16OH or H-16Ac evoked more c-Fos+ AOB neurons than the corresponding L-16OH and L-16Ac, respectively, in the mice of the same sex (M-H-16OH vs. M-L-16OH, \( t = 4.266, p = 0.000; M-L-16Ac vs. M-L-16OH, \( t = 7.529, p = 0.000; F-H-16Ac vs. F-L-16Ac, t = 4.030, p = 0.002). (B) Comparative analysis of the c-Fos+ cells in the aAOB of the same sex exposed to the same pheromone at either the low or high concentration. Significantly more activated neurons were found in both male and female aAOB following stimulation by all these four stimuli: low or high 16OH or 16Ac in comparison with the control samples whereas H-16OH or H-16Ac evoked more c-Fos+ aAOB neurons than the corresponding L-16OH and L-16Ac, respectively, in the mice of the same sex (M-L-16OH vs. M-control, \( t = 8.302, p = 0.000; M-H-16OH vs. M-control, t = 20.212, p = 0.000; M-H-16OH vs. M-L-16OH, t = 11.910, p = 0.000; M-L-16Ac vs. M-control, t = 12.126, p = 0.000; M-H-16Ac vs. M-control, t = 27.836, p = 0.000; M-H-16Ac vs. M-L-16Ac, t = 10.078, p = 0.000; F-L-16OH vs. F-control, \( t = 14.637, p = 0.000; F-H-16OH vs. F-control, \( t = 37.370, p = 0.000; F-H-16OH vs. F-L-16OH, t = 22.733, p = 0.000; F-L-16Ac vs. F-control, \( t = 3.499, p = 0.010; F-H-16Ac vs. F-control, \( t = 7.529, p = 0.000; F-H-16Ac vs. F-L-16Ac, t = 4.030, p = 0.002). (C) Comparative analysis of the c-Fos+ cells in the pAOB of the same sex mice exposed to the same pheromone at the either low or high concentration (M-L-16OH vs. M-control, t = 4.266, p = 0.000; M-L-16OH vs. M-control, t = 14.180, p = 0.000; M-H-16OH vs. M-L-16OH, t = 8.915, p = 0.000; M-L-16Ac vs. M-control, t = 6.651, p = 0.000; M-H-16Ac vs. M-control, t = 16.729, p = 0.000; M-L-16Ac vs. M-L-16Ac, t = 10.078, p = 0.000; F-L-16OH vs. F-control, \( t = 7.596, p = 0.000; F-H-16OH vs. F-control, \( t = 26.515, p = 0.000; F-H-16OH vs. F-L-16OH, t = 18.919, p = 0.000; F-L-16Ac vs. F-control, \( t = 0.950, p = 1.000; F-H-16Ac vs. F-control, \( t = 3.639, p = 0.002; F-H-16Ac vs. F-L-16Ac, t = 3.007, p = 0.041). (D) Comparative analysis of c-Fos+ neurons in the AOB of males versus females stimulated by 16OH or 16Ac at the low or high concentration (F-L-16OH vs. M-L-16OH, t = 7.119, p = 0.000; F-H-16OH vs. M-H-16OH, t = 17.963, p = 0.000; M-L-16Ac vs. F-L-16Ac, t = 5.881, p = 0.000; M-H-16Ac vs. F-H-16Ac, \( t = 16.039, p = 0.000; M-L-16Ac vs. F-L-16Ac, t = 5.426, p = 0.000; F-L-16OH vs. F-L-16Ac, t = 9.632, p = 0.000; F-H-16OH vs. F-H-16Ac, \( t = 18.335, p = 0.000; F-L-16Ac vs. F-L-16Ac, t = 4.187, p = 0.000; F-H-16Ac vs. F-H-16Ac, \( t = 11.258, p = 0.000). (E–I) Comparison of the efficacy of 16OH versus 16Ac on evoking the c-Fos response in the AOB. 16OH seemed to be more effective on the female aAOB, aAOB, or pAOB than on the male counterparts while 16Ac was more effective on male AOB, aAOB, or pAOB than on the female counterparts (AOB: M-L-16Ac vs. M-L-16OH, \( t = 3.388, p = 0.015; M-L-16Ac vs. M-H-16Ac, \( t = 5.426, p = 0.000; F-L-16OH vs. F-L-16Ac, \( t = 9.632, p = 0.000; F-H-16OH vs. F-H-16Ac, \( t = 28.576, p = 0.000; aAOB: M-L-16Ac vs. M-L-16OH, \( t = 3.825, p = 0.004; M-L-16Ac vs. M-H-16OH, \( t = 7.624, p = 0.000; F-L-16OH vs. F-L-16Ac, t = 11.138, p = 0.000; F-H-16OH vs. F-H-16Ac, \( t = 29.841, p = 0.000; pAOB: M-L-16Ac vs. M-L-16OH, \( t = 2.838, p = 0.029; M-H-16Ac vs. M-H-16OH, \( t = 2.548, p = 0.139; F-L-16OH vs. F-L-16Ac, \( t = 6.646, p = 0.000; F-H-16OH vs. F-H-16Ac, \( t = 22.559, p = 0.000).
FIGURE 8 | Continued
were also expressed as the mean ± s.e.m, and checked for distribution and homogeneity of variances. Two-way ANOVA was performed with both sex and treatment as between-group factors. After the significant interaction was found, similar post hoc analyses with Bonferroni’s multiple comparison tests were carried out as with the calcium imaging data using PASW Statistics 18.0 software.

RESULTS

Hexadecanol and Hexadecyl Acetate Can Activate the Vomeronasal Sensory Neurons of Male and Female Mice

To determine at the cellular level whether hexadecanol (16OH) and hexadecyl acetate (16Ac) can indeed function as pheromones on mice, we investigated VSN responses to these two compounds using calcium imaging on the vomeronasal sensory epithelium slices. The VSNs were loaded with the calcium-sensitive dye Fura 2-AM and stimulated sequentially with the two pheromones (Figures 1A,B). The results showed that increases in the intracellular calcium concentrations were observed in some VSNs in response to 100 µM 16OH and 100 µM 16Ac. Based on their responsiveness profiles, these VSNs could be classified into four types: those responsive to 16OH alone, to 16Ac alone, to both, or neither (Figures 1C,D).

From 5 male mice, out of 1175 VSNs examined, 14 (1.19%), 11 (0.93%), and 7 (0.6%) of VSNs responded to L-16OH, L-16Ac, or both, respectively (Figure 2A) whereas out of 988 VSNs examined, 17 (1.7%), 11 (1.1%), and 8 (0.81%) responded to H-16OH, H-16Ac, or both, respectively (Figure 2B). On the other hand, from 5 female mice, out of 1168 VSNs examined, 23 (1.97%), 10 (0.86%), and 6 (0.51%) responded to L-16OH, L-16Ac, or both, respectively (Figure 3A) whereas out of 964 VSNs examined, 41 (4.25%) responded to H-16OH, 17 (1.76%) to H-16Ac, and 16 (1.66%) to both, respectively (Figure 3B). Two-way ANOVA analysis with sex as a between-group factor and pheromone treatment as a within-group factor found that the effects of interaction (F(5,90) = 3.3, p = 0.0088), sex (F(1,18) = 11.8, p = 0.0029), and pheromone treatment (F(5,90) = 10.56, p < 0.0001) were all significant. A post hoc analysis with Bonferroni’s correction showed that no statistically significant difference was found among the numbers of cells responsive to L-16OH, L-16Ac, or both (Figure 2A), or among the numbers of cells responsive to H-16OH, H-16Ac, or both (Figure 2B), neither was found between the numbers of cells responsive to L- versus H-16OH, or L- versus H-16Ac in the male mice (Figure 2C).

In the female mice, however, significantly more VSNs responded to L-16OH than to both, whereas no significant differences in the numbers of the VSNs responsive to L-16OH versus to L-16Ac or to L-16OH and L-16Ac were found (Figure 3A). When the high concentration was applied, significantly more VSNs responded to H-16OH than to H-16Ac or to both, whereas cells responsive to H-16Ac were not significantly more than those to both (Figure 3B). Further analysis showed that H-16OH activated more female VSNs than 16OH, but the numbers of the cells responsive to 16Ac alone or to both were unaffected by their concentrations (Figure 3C).

Comparative analysis of the VSN responsiveness between sexes indicated that no significant differences were found between male and female VSNs in response to L-16OH alone, L- or H-16Ac alone, both L-16OH and L-16Ac, or both H-16OH and H-16Ac except that significantly more female VSNs than male VSNs responded to H-16OH (Figure 4).

The Activated VSNs Are Mostly Located in the Apical Vomeronasal Sensory Epithelium

To determine which type of the VSNs responsive to the two pheromones, we used an antibody to the phosphorylated small ribosomal protein 6 (pS6) to identify activated VSNs and another antibody to G protein-coupled receptor 37 (GPR37) to identify both aVSE and bVSE of the control mice exposed to double distilled water (ddH2O) were not significantly different (Figures 5C,D).
FIGURE 9 | Continued
Projection of 16OH and 16Ac Signals to the Accessory Olfactory Bulb

To determine the projection of 16OH and 16Ac signals from the VNO to AOB, we used c-Fos and Gα2 immunostaining to identify and locate the activated neurons to the anterior or posterior half of the AOB (Figure 6). The numbers of c-Fos+ neurons in the AOB were significantly greater in both male and female mice exposed to both low and high 16OH and 16Ac than that in the control mice exposed to 0.01% DCM-containing mineral oil (Figure 6 and Table 1). In comparison of the c-Fos expression between the two AOB halves, we found that the aAOB had more c-Fos+ neurons than the pAOB in both male and female mice in response to low and high 16OH or 16Ac (Figure 6D and Table 1). Furthermore, H-16OH or H-16Ac evoked more c-Fos-positive neurons than L-16OH or L-16Ac, respectively, in the AOB of the mice of the same sex (Figure 7A).

When the data from the aAOB and pAOB were processed separately, we found that 16OH and 16Ac induced stronger c-Fos expression at both low and high concentrations than control, except that in the female pAOB, L-16Ac failed to do so. Further, H-16OH and H-16Ac were more effective in eliciting c-Fos responses than their low counterparts in both male and female, aAOB or pAOB (Figures 7B,C).

As for sexual differences, the female AOB had more c-Fos+ cells than the male AOB in response to both L- and H-16OH whereas the male AOB had more c-Fos+ cells than the female AOB in response to both L- and H-16Ac. The pattern still held even when the data from the two AOB halves were analyzed separately: the female aAOB and pAOB responded significantly stronger than male aAOB and pAOB to both L- and H-16OH whereas in response to L- and H-16Ac, male aAOB and pAOB had more c-Fos+ neurons than their female counterparts (Figures 7D–F).

Comparative analysis of the efficacy of the two pheromones indicated that on the female AOB, L- or H-16OH was more effective than L- or H-16Ac whereas on the male AOB, L- or H-16Ac was more effective than L- or H-16OH. And the pattern was still true if the c-Fos immunostaining data were analyzed for aAOB and pAOB separately except the male pAOB (Figures 7G–I).

Following the initial processing in the AOB, pheromonal signals are known to be sent to the next orders of the neurons in several brain regions, including the BNST, MeA, PMCo, and then from both BNST and amygdaloid nuclei further downstream to the MPA and VMH (Supplementary Figure 1; Dulac and Wagner, 2006). To determine how the 16OH and 16Ac signals are processed in the higher brain regions, we carried out c-Fos immunohistochemistry on the brain sections containing the aforementioned nuclei to identify the neurons constituting the neural circuitry for these two pheromones.

16OH and 16Ac Induce c-Fos Immunoreactivity in the BNST

After exposure to low and high 16OH and 16Ac, the male and female mice were examined for the c-Fos-immunoreactivity in the BNST (Figure 8). The results showed that in comparison with the 0.01% DCM-containing mineral oil-treated control samples, either L-16OH or L-16Ac could evoke significantly more c-Fos signals from male or female BNST where H-16OH and H-16Ac were able to do so on both male and female BNST except H-16Ac on the female BNST. And H-16OH and H-16Ac induced significantly more c-Fos+ cells than their counterparts of the low concentration on both male and female BNST except H-16Ac on the female BNST (Figure 8K and Table 1).

Comparison between sexes indicates that the female BNST responded more strongly to H-16OH than the male BNST whereas the male BNST did to H-16Ac than the female (Figure 8L). Comparison between the two pheromones indicates that H-16OH was more effective than H-16Ac on the female BNST whereas H-16Ac was more so than H-16OH on the male BNST (Figure 8M).

16OH and 16Ac Activate Neurons in the Medial Amygdaloid Nucleus

A major portion of the AOB output is known to be relayed to the MeA. To determine how the neurons in the nucleus are activated by 16OH and 16Ac, c-Fos immunoreactivity was also examined on the anterior subdivision of this nucleus (Figure 9). The results showed that the c-Fos immunoreactivity pattern mirrored, to some extent, that found in the BNST, e.g., in the male MeA, both L- and H-16Ac and H-16OH were able to elicit more c-Fos immunoreactivity than control, but only H-16Ac was more effective than L-16Ac in the male, whereas, in the female MeA, only H-16OH was able to evoke significantly more c-Fos immunoreactivity.
FIGURE 10 | Continued
immunoreactivity than control. However, both L-16OH and L- and H-16Ac failed to augment c-Fos response in the female MeA. Comparison between sexes or between the two pheromones lead to the similar results as found in the BNST, that is, the female MeA more robustly responded to H-16OH whereas the male MeA did to H-16Ac; and H-16OH was more effective than H-16Ac on the female MeA whereas the opposite was true on the male MeA (Figure 9 and Table 1).

**16OH and 16Ac Activate Neurons in the Posteromedial Cortical Amygdaloid Nucleus**

The PMCo is another nucleus that directly receives signals from the AOB. Comparative analyses showed that H-16OH but not L-16OH was able to produce significantly more c-Fos immunoreactivity in both male and female PMCo than control whereas only H-16Ac produced significantly more c-Fos signals than control in the male but not female PMCo whereas L-16Ac were ineffective on either male or female PMCo (M-L-16OH vs. M-control, t = 1.384, p = 0.100; M-H-16OH vs. M-control, t = 3.570, p = 0.008; M-L-16Ac vs. M-control, t = 2.122, p = 0.388; M-H-16Ac vs. M-control, t = 8.317, p = 0.000; F-L-16OH vs. F-control, t = 1.193, p = 1.000; F-H-16OH vs. F-control, t = 11.845, p = 0.000; F-L-16Ac vs. F-control, t = 0.913, p = 1.000; F-H-16Ac vs. F-control, t = 1.841, p = 0.716; M-H-16OH vs. M-L-16OH, t = 2.187, p = 0.335; F-H-16OH vs. F-L-16OH, t = 10.651, p = 0.000; M-H-16Ac vs. M-L-16Ac, t = 6.194, p = 0.000; F-H-16Ac vs. F-L-16Ac, t = 0.267, p = 1.000). The female PMCo was more sensitive than the male to H-16OH whereas the male PMCo was more so to H-16Ac than the female, and H-16OH was more effective than H-16Ac on the female MeA whereas on the male MeA, the opposite was true (F-H-16OH vs. M-H-16OH, t = 8.447, p = 0.000; M-H-16Ac vs. F-H-16Ac, t = 6.303, p = 0.000; F-L-16OH vs. M-L-16OH, t = 0.0175, p = 0.988; M-L-16Ac vs. F-L-16Ac, t = 1.037, p = 0.305; F-H-16OH vs. F-H-16Ac, t = 10.004, p = 0.000; M-H-16Ac vs. M-H-16OH, t = 4.746, p = 0.000; F-L-16OH vs. F-L-16Ac, t = 2.187, p = 0.716; F-L-16Ac vs. M-L-16Ac, t = 0.739, p = 1.000). N = 6 for each group. All values are expressed as mean ± s.e.m. **p < 0.01, ***p < 0.001.

**16OH and 16Ac Activate Neurons in the Ventromedial Hypothalamic Nuclei**

The signals from the MPA are known to be relayed to the VMH. c-Fos immunoreactivity analyses showed that the response patterns were nearly identical to those found in the MPA, that is, all stimulations, L- or H-16OH, L- or H-16Ac generated more c-Fos signals than control on both male and female VMH (Figure 12 and Table 1). But the concentration-dependent increases in c-Fos signals were found for 16OH on both male and female VMH only. The female VMH was more sensitive than the male VMH to H-16OH whereas the opposite was true to H-16Ac. H-16OH was more effective than H-16Ac on the female VMH (Figure 12).

**DISCUSSION**

The mammalian olfactory system is comprised of two subsystems: the MOS and the AOS. While the former is mainly responsible for detecting volatile odorants, the latter is largely for detecting less or non-volatile pheromones, although some cross-detections between these two have been reported as well (Holy, 2018; Mohrhardt et al., 2018). Unlike a relatively large number of odorants and their corresponding olfactory receptors, very few pheromones or their receptors have been structurally identified or deorphanized. Hexadecanol (16OH) and hexadecyl acetate (16Ac) are two semi-volatile and rare chemically pure pheromones that exert differential effects on male and female mice (Zhang et al., 2008). In-depth studies on these two pheromones can facilitate our understanding of the AOS mechanisms.

Calcium imaging of the VNO slices indicated that a small percentage (0.6–1.9%) of male and female VSNs responded to 16OH only, 16Ac only, or both 16OH and 16Ac (Figures 1–4), which is very similar to the percentages of VSNs activated by several other known pheromones, including the small molecules: 2,5-dimethypyrazine, SBT, 2,3-dehydro-exo-brevicomin, α- and β-farnesenes, 2-heptanone, and HMH,
FIGURE 11 | Continued
and the large molecules such as peptides ESPs and major urine proteins (Leinders-Zufall et al., 2000; Kimoto et al., 2007; Ferrero et al., 2013; Dey et al., 2015). Notably, only one vomeronasal receptor has been identified for each of these pheromones: Vmnr1g9 for 2-heptanone, Vmnr2r116 for ESP1, and V2Rp4 for ESP22 (Boschat et al., 2002; Haga et al., 2010; Osakada et al., 2018). Thus, given the similar percentages of the activated VSNs, we conclude that it is likely only one vomeronasal receptor each for detecting 16OH, 16Ac, or both.

Further, our data also showed that the VSNs responsive to 16OH, 16Ac, or both were preferentially located in the apical VSE, and that the numbers of responsive neurons did not increase in most cases even when the pheromone concentrations were raised by 10,000-fold, from 10 nM to 100 μM, with one exception of the female VSNs that significantly increased by twofold upon stimulation of 100 μM 16OH, i.e., H-16OH (Figures 2–5). These results indicate that probably there is only one V1r in the male and female VSE sensing 16OH, 16Ac, or both, respectively, whereas the female VSNs may express an additional V1r for detecting the high concentration of 16OH. On the other hand, however, the basal VSNs also responded, to a lesser extent, to these chemicals. Thus, it is also possible that there is another V2r responsive to each of 16OH and 16Ac or both, respectively.

The pheromonal signals detected by VSNs are then projected to the AOB. Our c-Fos immunostaining data indicate that significantly more c-Fos+ neurons were found in the aAOB than in the pAOB (Figure 6), which is in congruency with the calcium imaging data showing the apical location of most responsive VSNs (Figure 5). However, the male pAOB responded to all four stimuli: L- and H-16OH, L- and H-16Ac whereas the female pAOB did to three of the four: L- and H-16OH, and H-16Ac, but not to L-16Ac (Figure 7), which is also consistent with the VNO results showing calcium responses also from the basal VSNs to 16OH and 16Ac, or both, supporting the idea of V2r responsive to these two pheromones.

The numbers of male and female c-Fos+ neurons in the AOB, aAOB, or pAOB were 16OH and 16Ac dose-dependent (Figure 7), which is different from the observation on the VSE, of which only the number of the female 16OH-responding VSNs was 16OH concentration-dependent. This discrepancy may be explained by the fact that VSN firing rates vary from 2.5 to 80 Hz (Stowers et al., 2002; Noradi et al., 2008; Kim et al., 2012), and that individual VSNs innervate multiple AOB glomeruli; conversely, some individual glomeruli receive projections from multiple VSNs expressing different vomeronasal receptors (Belluscio et al., 1999; Rodriguez et al., 1999; Wagner et al., 2006), although a mitral cell tends to receive inputs from different glomeruli that are innervated by sensory neurons expressing the same type of V1r or V2r receptor (Del Punta et al., 2002). Thus, high concentrations of 16OH and 16Ac may trigger more spikes from VSNs, increasing firing rate, which cannot be detected by the VSN calcium imaging, but activate more glomeruli, generating more c-Fos+ AOB neurons. Further studies, however, are needed to validate this notion.

Mitral cells in the AOB send their axons to other AOS circuit hubs in the brain. Interestingly, the BNST receives inputs only from the aAOB, not from pAOB (Martinez-Marcos and Halpern, 1999; Mohedano-Moriano et al., 2007). The c-Fos immunoreactivity pattern in the BNST was closely similar to that in the AOB: the male BNST exhibited significantly more c-Fos+ neurons in response to H-16OH and H-16Ac whereas the female BNST showed significant responses to H-16OH (Figure 8). These results indicate that the information is relayed largely intact from the AOB to BNST in response to the high concentration of 16OH or 16Ac.

Mitral cells in the AOB can also send axons to the MeA and PMCo, followed by the projection of BNST and MeA neurons to the MPA, and finally to the VmH. The overall profiles of c-Fos immunoreactivity in these nuclei were similar to that found in the BNST, and can be summarized as following: (1) dose-dependency: stronger c-Fos responses were induced by higher concentrations of 16OH and 16Ac than lower ones, respectively; (2) sexual dimorphism: the female nuclei responded more strongly to 16OH whereas male nuclei did so to 16Ac; and (3) the relative effectiveness between the two pheromones: significant differences in evoking c-Fos responses in the nuclei were found between the two pheromones only at their high concentrations but not at the low concentrations. There are, however, some subtle variations. For example, the female MeA exhibited...
FIGURE 12 | Continued
insignificant increases in the number of c-Fos+ neurons in response to not only L- or H-16Ac but also L-16OH, as in the BNST and PMCo (Figures 9, 10), whereas the male BNST and PMCo nuclei did not respond to L-16OH or L-16Ac. In the MPA and VmH, males responded equally to H-16OH and H-16Ac.

Neurons in the nuclei examined so far in this study are involved in various pheromone-induced activities. For example, activation of some BNST neurons can suppress female sexual behavior (Osakada et al., 2018), and even the PMCo, which does not have much direct connections to the hypothalamic nuclei, process and relay most information from the AOB to the MeA and posterolateral amygdaloid cortex (Gutierrez-Castellanos et al., 2014). The MeA, which receives excitatory signals from the AOB and the PMCo, sends axons to different amygdala and hypothalamic areas, directing sex-specific behavioral responses as well as endocrine and autonomic responses (Brennan and Zufall, 2006; Ishii et al., 2017). The MPA is known to be critical to control inter-male and pup-directed aggression, sexual behaviors, and parental care (Wu et al., 2014; Wei et al., 2018). Finally, some neurons in the VmH are involved in initiating aggression, parental caring, and mating as well (Lin et al., 2011; Kohl et al., 2018). However, the only documented behavioral responses induced by 16OH and 16Ac include attraction and aversion, with no evident aggression, mating, or parental caring observed yet (Zhang et al., 2008). Our c-Fos data indicate that 16OH- and 16Ac-activated neurons in the brain regions can be parts of the AOS circuit hubs, contributing to the modulation or fine-tuning of the behavioral outputs determined by these ensembles of neurons (Li et al., 2017; Li and Dulac, 2018). Further investigations are needed to unravel the exact roles played by the 16OH- and 16Ac-activated neurons in the population coding of the pheromonal signals.

Like many other pheromones, 16OH and 16Ac also evoke sexually dimorphic responses. In the VNO, similar numbers of male VSNs responded to both 16OH and 16Ac; in contrast, many more female VSNs did to 16OH than to 16Ac (Figures 1–3); and upon stimulation of H-16OH, far more VSNs were activated in females than in males (Figures 4, 5). In the AOB and most other brain regions, it seems largely consistent that males responded more strongly to 16Ac, females more strongly to 16OH, especially at the high concentration, except in the male MPA and VmH (Figures 6–12). It is known that AOS brain regions between sexes differ in neuron numbers and sex hormone receptor expression (Li and Dulac, 2018). The AOS can be modulated by the inputs from the MOS, by physiological states and experiences. For example, the enzyme aromatase can convert testosterone into estrogen, which acts on the neurons expressing the estrogen receptors, modulating their activity, while mating experience makes a subset of male MeA neurons more responsive (Dey et al., 2015; Xu et al., 2016; Lischinsky et al., 2017). Additional studies are needed to reveal the behavioral and physiological consequences resulted from or modulated by 16OH and 16Ac sexual dimorphic effects.

CONCLUSION

In conclusion, our study demonstrates the activation of VSNs by 16OH and 16Ac, and the activation of neurons in the circuit hubs of the AOS in a sexually dimorphic manner. Our data indicate that there is one V1r each for 16OH, 16Ac, or both, and possibly another V1r for the concentrated 16OH. And activation of the neurons in the brain regions strongly suggests that 16OH and 16Ac can modulate the AOS outputs such as aggression, mating, and parental caring. Further studies on these two abundantly produced and chemically identified natural pheromones can facilitate our understanding of this ancient and vital system in many mammals.

DATA AVAILABILITY STATEMENT

All datasets generated for this study are included in the manuscript/Supplementary Files.

ETHICS STATEMENT

The animal study was reviewed and approved by the Institutional Animal Care and Use Committees of both the Zhejiang University and the Institute of Zoology, Chinese Academy of Sciences.
AUTHOR CONTRIBUTIONS

QL, J-XZ, and LH designed the experiments, analyzed and interpreted the data, and wrote the manuscript. QL, XG, PW, YZ, and YW performed the experiments.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fncel.2019.00455/full#supplementary-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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