THE VOMERONASAL ORGAN IS REQUIRED FOR THE MALE MOUSE MEDIAL AMYGDALA RESPONSE TO CHEMICAL-COMMUNICATION SIGNALS, AS ASSESSED BY IMMEDIATE EARLY GENE EXPRESSION

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Abstract—Many species use chemical signals to convey information relevant to social and reproductive status between members of the same species (conspecific), but some chemical signals may also provide information to another species (heterospecific). Both of these types of complex chemical signals may be detected by the vomeronasal organ, which sends projections to the accessory olfactory bulb and on to the medial amygdala. Previous reports in hamster and mouse suggest that the medial amygdala sorts this complex chemosensory information categorically, according to its biological relevance (salience). In the present set of experiments, male mice having undergone vomeronasal removal surgery (VNX) or a sham-operation (SHAM) were exposed to conspecific (male and female mouse urine) or heterospecific (hamster vaginal fluid and worn cat collar) chemical stimuli. Similarly to our previous report with intact male mice (Samuelsen and Meredith, 2009) Brain Res 1263:33–42, SHAM mice exhibit different immediate early gene (IEG) expression patterns in the medial amygdala dependent upon the biological relevance of the chemical stimuli. However, regardless of biological relevance, vomeronasal organ removal eliminates all responses in the medial amygdala to any of the chemical stimuli. Interestingly, VNX also disrupts the avoidance of (an unfamiliar) predator odor, worn cat collar. Here we show that the medial amygdala response to the tested chemical signals is dependent upon an intact vomeronasal organ.

Key words: olfaction, pheromone, Fos-related antigens (FRAs), predator, behaviour.

Most mammals employ chemical signals to communicate information pertaining to complex behaviors such as reproductive status, relatedness (strain), social rank (Desjardins et al., 1973) and territorial ownership (Nakamura et al., 2007). The volatile components of chemical signals appear to be detected mainly by receptors in the vomeronasal organ (VNO) with the information processed via the accessory olfactory bulb (Leinders-Zufall et al., 2000; Luo et al., 2003). Both sensory systems can potentially respond to both types of stimuli (Meredith and O’Connell, 1979; Lehman and Winans, 1982; Meredith et al., 1983; O’Connell and Meredith, 1984; Spehr et al., 2006). Inactivation of the VNO, by surgical removal or genetic manipulation, results in behavioral and physiological deficits in reproductive and other behaviors that involve chemical communication (Powers and Winans, 1975; Wysocki et al., 1984; Meredith, 1986; Fewell and Meredith, 2002; Stowers et al., 2002; Westberry and Meredith, 2003a,b; Pankevich et al., 2004, 2006; Keller et al., 2006; Kimchi et al., 2007). Recently, it has been shown that VNO receptors respond differentially to putative pheromones (Leinders-Zufall et al., 2000; Chamero et al., 2007), stimuli from animals differing in gender, strain or disease state (He et al., 2008; Riviére et al., 2009), suggesting that the information necessary to interpret and respond to conspecific biologically relevant chemical signals is present at the level of the VNO epithelium. However, interpretation of this information, which is distributed across several thousand sensory neurons, must occur more centrally.

The amygdala, particularly the medial amygdala (Lehman et al., 1980; Petrulis and Johnston, 1999; Blanchard et al., 2005), is important for the behavioral response to chemosensory information. The medial amygdala is the first site of convergence of afferent main olfactory and vomeronasal information in the brain (Shipley and Adamek, 1984; Pitkanen et al., 1997; Coolen and Wood, 1998; Meredith, 1998; Chamero, 2007). Exposure to different salient chemosensory stimuli (reproductive, territorial or predator odors) characteristically changes immediate early gene (IEG) expression in the medial amygdala (Meredith and Westberry, 2004; Samuelsen and Meredith, 2009). Chemical signal input, via the medial amygdala, is thought to be important for the proposed hypothalamic circuits involved in defensive and reproductive behavior (Canteras, 2002; Choi et al., 2005).

Previously we have shown that patterns of IEG expression in posterior medial amygdala depend on the biological relevance of the chemical stimuli. A range of conspecific (same species) chemical signals from males and females, relevant for reproductive or agonistic behavior, all activate both anterior and posterior medial amygdala. A range of non-related stimuli from other species (heterospecific) activate anterior medial amygdala, but fail to activate pos-
terior medial amygdala. This categorical difference is seen in mice (Samuelsen and Meredith, 2009) and hamsters (Meredith and Westberry, 2004; Meredith et al., 2008). Thus, the pattern of IEG expression in medial amygdala differs categorically according to the biological relevance of the chemical stimuli. Given the importance of the VNO to normal behavioral and physiological function, we reasoned that surgical removal of the VNO would result in a disruption of IEG expression patterns in the medial amygdala to biologically relevant chemosensory stimuli.

Here we show that the VNO is necessary for significant medial amygdala response to critical chemical signals in sexually inexperienced (naive) male mice. Surgical removal of the vomeronasal organ (VNX) actually eliminates all significant medial amygdala IEG expression above control levels to all tested chemical signals, regardless of biological relevance. The tested stimuli include male and female mouse urine (mMU, fMU), known to convey reproductive and agonistic information to male mice (Hurst and Beynon, 2004; Pankevich et al., 2004, 2006; Nakamura et al., 2007), and hamster vaginal fluid (HVF), a heterospecific stimulus known to convey reproductive information from female to male hamsters (Johnston, 1977), but presumably irrelevant for mice. In a separate experiment, collars worn by a cat, which mice and rats normally avoid (Dielenberg et al., 1999, 2001; Takahashi et al., 2005, 2007; Samuelsen and Meredith, 2009), were used as a biologically relevant heterospecific stimulus.

**EXPERIMENTAL PROCEDURES**

**Animals**

Forty eight (Exp. 1) and 24 (Exp. 2) sexually naive 3–4 month old male C57 BL/6 mice (Jackson Laboratory) were maintained on a reverse 12/12 h light/dark cycle with food and water ad libitum. All animal procedures were approved by the Florida State University Institutional Animal Care and Use Committee. Animals had no contact with any heterospecific stimuli before the experimental session, no contact with females or female stimuli since weaning and no contact with the male or female conspecific stimulus donors.

**Vomeronasal organ removal surgery (VNX)**

In both sham-operation (SHAM) and VNX surgery, a midline incision was made along the palate exposing the VNO capsule. For SHAM surgery, the exposed VNO capsule was examined and then the palatal incision was closed with two to three sutures and cyanoacrylate adhesive. For VNX, a dental drill was used to separate the VNO capsule from the rostral palatal bones. Using forceps, the medial palatine process of the maxillary bones was broken, separating the VNO capsule from the hard palate at its caudal end. The two VNO capsules were detached with forceps, with care to ensure that the capsule contents were removed with the capsules, and the palatal incision was closed as above. Experiments were conducted 7 days post surgery. To confirm VNX, decalcified noses were cut coronally into 20 μm sections, stained with Gills hematoxylin and examined microscopically for intact vomeronasal epithelium (Fig. 1). Fifteen animals with residual VNO tissue or excess nasal debris were excluded.

**Stimuli**

Female mouse urine was collected from three to four adult mice placed in a metabolic cage over a 5 day period. Five days of collection was used in order to collect urine from all estrus stages of normally cycling female mice (Champlin et al., 1973). Male mouse urine was collected in a similar manner. Hamster vaginal fluid was collected from two to four adult female hamsters in behavioral estrus using a spatula, mixed together and placed in centrifuge tubes. As in our previous experiments (Meredith and Westberry, 2004; Samuelsen and Meredith, 2009), all liquid stimuli were diluted 1:10 by weight with distilled water (purified by reverse osmosis and polishing with activated carbon) and centrifuged for 30 min (Fisher clinical centrifuge at medium speed). The supernatant was decanted and held at −20 °C until presentation. Heterospecific stimuli are diluted to avoid any neophobia in test animals; conspecific stimuli are diluted equally to match, where possible. Soft nylon cat-collars (CC) (PETCO Single Ply Nylon Collar) were unworn (control) or were worn for 2 weeks by a neutered male house cat. The collars were removed, placed in zip-lock plastic bags and held at −20 °C until use.

**Testing procedure and stimulus presentation**

All mice were single housed for 7 days post-surgery and before testing, minimizing exposure to other male odors. On the day of the experiment, mice were placed in a clean cage with clean corn cob bedding and allowed 2 min to acclimate to the surroundings.
Experiment 1: Polyester swab-tips (Puritan Medical Products Company) were used to present the stimuli. Clean swabs (control) or swabs containing ~200 µl of liquid stimulus were presented in the middle of the cage and replaced every 3 min for a total of 15 min; a total of five scented swabs or five control swabs per animal. Experiment 2: A 2.5 × 1.27 cm² piece of CC worn for 2 weeks or a piece of clean CC was presented in the middle of the clean cage and left for the entire 15 min trial. For both experiments, behavior was recorded using a computer program and numbered key pad with each key corresponding to a different behavior. The computer records the latency, the number of presses and total elapsed time each key is depressed. Behaviors recorded were the number of times the mouse contacted or very closely investigated the stimulus, the duration of investigation of the stimulus, number of rears, time spent rearing and general investigation of the cage. All animals were tested in the first 6 h of the dark phase of the light cycle in a room lit by red light.

Immunocytochemistry

Forty-five minutes after the initial stimulus exposure, mice were anesthetized with Nembutal and perfused with cold 0.1 M PBS followed by 4% paraformaldehyde (PFA). Brains were removed and post-fixed overnight in 4% PFA. The next morning brains were placed in 30% sucrose overnight for cyroprotection. Using a freezing microtome, brains were sliced into 40 µm sections. Alternate free-floating coronal sections were washed in 0.1 M PBS, blocked in a solution of 5% normal goat serum (30 min) and incubated in rabbit anti-fos-related antigens (FRAs) primary antibody solution (SC253—detects c-Fos, Fos B, Fra-1 and Fra-2; 1:10,000; Santa Cruz Bio/technology, Santa Cruz, CA, USA) for 20–24 h at room temperature. The next day, sections were washed in 0.1 M PBS and incubated in biotinylated goat anti-rabbit secondary antibody solution (1:400; Vector Laboratories, Burlingame, CA, USA) for 2 h. Sections were processed in ABC reagent (Vector Laboratories) for 1 h and stained with diamino benzidine (DAB) (Vector Laboratories). FRAs expression was assessed by averaging numbers of densely labeled cell nuclei within areas of interest on both sides of the brain in three adjacent sections per anatomical area. Areas of interest included: (1) Anterior medial amygdala (MeA), which was divided into ventral anterior medial amygdala (MeAv) and dorsal anterior medial amygdala (MeAd); (2) Posterior medial amygdala (MeP), which was divided into ventral posterior medial amygdala (MePv) and dorsal posterior medial amygdala (MePd); both as indicated in the mouse brain atlas (Paxinos and Franklin, 2003; Fig. 3). Image analysis software (ImagePro Plus, Media Cybernetics, Inc., Bethesda, MD, USA) was used to count all densely labeled cell nuclei within the borders of the neuroanatomical nucleus of interest. The numbers are presented as means and standard errors. Data are presented separately for MeP as a whole and MeA as a whole, in addition to the data for individual subdivisions, in order to facilitate comparisons with earlier reports that did not provide data for subdivisions.

Statistics

IEG expression comparisons were analyzed for each experiment by two-way analysis of variance (ANOVA) with factors surgery (SHAM or VNX) and stimulus (Experiment 1: CON, mMU, fMU and HVF; Experiment 2: control-CC and worn CC). Post-hoc comparisons were made using the Holm-Sidak test. Behaviors were analyzed using Two-way ANOVAs, with post-hoc comparisons using the Holm-Sidak test. Behavioral responses were analyzed using factors of surgery (SHAM and VNX) and stimulus (Experiment 1: CON, mMU, fMU and HVF; Experiment 2: control-CC and worn CC). Reported behaviors include number and cumulative duration of close investigation or contact with the stimulus swab/collar, number of rears, and time spent rearing.

Results

Exposure to mMU, fMU or CC stimuli, all representatives of behaviorally salient stimuli, increased IEG expression in both MeA and MeP in sham-operated (SHAM) male mice. HVF, selected as a representative non-salient stimulus, increased IEG expression in MeA, but not MeP of SHAM mice. These expression patterns confirm our previous findings, using intact mice, of the characteristic and categorical difference between medial amygdala response to biologically relevant and to non-relevant stimuli (Samuelsen and Meredith, 2009) and conceptually identical results in hamsters (Meredith and Westberry, 2004). Data are presented for MeA and MeP as a whole, in addition to data for subdivisions in order to facilitate comparison with these reports. After VNX surgery, none of the stimuli significantly increased IEG expression in medial amygdala of VNX male mice.

Experiment 1

Response to fMU. SHAM male mice exposed to fMU had significantly greater FRAs expression in MeA (F(1,39)=50.7, P<0.001) and MeP (F(1,39)=35.4, P<0.001), compared to control-SHAM mice exposed to clean-swabs (Supplemental Fig. 1). They also had a significant increase in FRAs expression compared to control in all four medial amygdala subdivisions (ventral anterior medial amygdala (MeAv): F(1,39)=43.2, P<0.001; dorsal anterior medial amygdala (MeAd): F(1,39)=31.2, P<0.001; ventral posterior medial amygdala (MePv): F(1,39)=35.0, P<0.001; dorsal posterior medial amygdala (MePd): F(1,39)=20.9, P<0.001) (Fig. 2). SHAM mice also had significantly greater IEG expression in response to fMU compared to SHAM males exposed to clean-swabs (P<0.001).
to fMU than VNX mice, in MeA and MeP (MeA: $F(1,39)=22.6$, $P<0.001$; MeP: $F(1,39)=58.4$, $P<0.001$), and in each subdivision (MeAv: $F(1,39)=65.6$, $P<0.001$; MeAd: $F(1,39)=50.0$, $P<0.001$; MePv: $F(1,39)=62.0$, $P<0.001$; MePd: $F(1,39)=30.4$, $P<0.001$). Representative brain sections of the FRAs expression in both MeA and MeP to fMU in SHAM and VNX animals are shown in Fig. 3.

There was no significant difference in FRAs expression between fMU-exposed VNX mice and SHAM control mice exposed to clean-swabs in any region or subregion of medial amygdala.

Response to mMU. SHAM mice exposed to mMU had significantly greater FRAs expression in MeA ($F(1,39)=22.6$, $P<0.001$; MeP: $F(1,39)=98.4$, $P<0.001$), and in each subdivision (MeAv: $F(1,39)=71.6$, $P<0.001$; MeAd: $F(1,39)=60.0$, $P<0.001$; MePv: $F(1,39)=82.0$, $P<0.001$; MePd: $F(1,39)=40.4$, $P<0.001$). Representative brain sections of the FRAs expression in both MeA and MeP to mMU in SHAM and VNX animals are shown in Fig. 3.
36.2, P<0.001) and MeP (F(1,39)=16.5, P<0.001), compared to control SHAM male mice exposed to a clean swab (Supplemental Fig. 1). They also had significant FRAs expression compared to control in all four medial amygdala subdivisions (MeAv: F(1,39)=41.7, P<0.001; MeAd: F(1,39)=13.7, P<0.001; MePv: F(1,39)=15.4, P<0.001; MePd: F(1,39)=10.0, P<0.001) (Fig. 2). SHAM mice also had significantly greater IEG expression in response to mMU than VNX mice, in MeA and MeP (MeA: F(1,39)=79.2, P<0.001; MeP: F(1,39)=14.3, P<0.001), and in each subdivision except MePd (MeAv: F(1,39)=29.4, P<0.001; MeAd: F(1,39)=6.75, P<0.013).

There was no significant difference in FRAs expression between mMU-exposed VNX mice and SHAM control mice exposed to clean-swabs in any region or subregion of medial amygdala.

**Response to HVF.** SHAM male mice exposed to the heterospecific odor, HVF, exhibited significantly higher FRAs expression in MeAv, MeAd (Fig. 2) and overall MeA (Supplemental Fig. 1) as compared to both SHAM control males exposed to clean-swabs (Fig. 2) (MeA: F(1,39)=17.9, P<0.001; MeAv: F(1,39)=20.6, P<0.001; MeAd: F(1,39)=7.37, P<0.01) and to VNX mice exposed to HVF (MeA: F(1,39)=22.9, P<0.001; MeAv: F(1,39)=21.4, P<0.001; MeAd: F(1,39)=13.1, P<0.001). There was no significant FRAs expression in SHAM mice or VNX mice, after exposure to HVF, in the measured areas of MeP or its subdivisions.

There was no significant difference in FRAs expression between HVF-exposed VNX mice and SHAM control mice exposed to clean-swabs in any region or subregion of medial amygdala.

**Behavioral response to stimuli.** In Experiment 1, there was a significant main effect of surgery (F(1,39)=7.9, P<0.01), but no significant main effect of stimulation, and no significant interaction. VNX mice exposed to IMU spent significantly more time investigating the stimulus swab than their SHAM-operated counterparts (F(1,39)=10.4, P<0.005). Although not significant, VNX mice spent more time than SHAM mice investigating HVF (F(1,39)=3.5, P=0.07). There were no significant differences in any other of the measured behaviors (Fig. 4).

**Experiment 2**

**Responses to CC.** SHAM mice exposed to a 2.5 cm piece of CC (worn for 2 weeks) had significantly greater FRAs expression in MeA (F(1,23)=20.0, P<0.001) and MeP (F(1,23)=22.1, P<0.001) compared to SHAM mice exposed to a piece of clean collar (Supplemental Fig. 2). They also have increased FRAs expression in the subregions MeAv (F(1,23)=17.2, P<0.001), MeAd (F(1,23)=17.3, P<0.001) and MePv (F(1,23)=44.9, P<0.001) (Fig. 5). There was no significant difference in the MePd subregion with CC exposure. SHAM mice also had greater FRAs expression in response to CC than VNX mice in MeA (F(1,23)=38.6, P<0.001) and MeP (F(1,23)=40.2, P<0.001), and in each subregion except MePd (MeAv: F(1,23)=31.8, P<0.001; MeAd: F(1,23)=33.8, P<0.001; MePv: F(1,23)=69.1, P<0.001). In MePd, only the difference between SHAM and VNX mice exposed to CC was significant (F(1,23)=7.81, P<0.011).

There was no significant difference in FRAs expression between CC-exposed VNX mice and SHAM control mice exposed to clean-swabs in any region or subregion of medial amygdala.
Behavioral response to CC stimuli. There were no main effects of surgery or stimulus in Experiment 2, but there was a significant interaction ($P<0.01$), indicating a difference in response to the stimulus between SHAM and VNX mice. Although SHAM mice spent less time investigating worn CC than control CC, the difference was not significant. VNX mice exposed to CC spent significantly more time investigating the stimulus collar than either VNX clean-collar controls ($F(1,23)=5.3, P<0.035$) or SHAM CC exposed mice ($F(1,23)=11.7, P<0.005$). Although not significant, SHAM mice spent less time investigating CC than those which were exposed to control CC ($F(1,23)=4.0, P=0.06$). There was no difference in any of the other measured behaviors (Fig. 6).

DISCUSSION

In this report, we provide evidence that IEG response to biologically relevant chemical signals in the medial amygdala depends on an intact VNO. We have previously reported that the pattern of IEG expression in medial amygdala to biologically relevant (mainly conspecific) chemical signals is categorically different from the pattern of most heterospecific stimuli. These non-relevant heterospecific chemosensory stimuli do elicit an amygdala response, but have little or no effect on behavior and may be unimportant to the responding animals. In particular, biologically relevant stimuli increased IEG expression in both MeA and MeP, but stimuli with no apparent relevance increased expression only in MeA. This is the case for both mice (Samuelsen and Meredith, 2009) and hamsters (Meredith and Westberry, 2004). As mouse VNO neurons selectively respond to biologically relevant features of conspecific urine stimuli (Leinders-Zufall et al., 2000; Chamero et al., 2007; He et al., 2008), we expected the responses to conspecific signals in the medial amygdala to be disrupted by VNX. Although, we did not test a wide range of stimuli here, our evidence suggests that surgical removal of the VNO eliminates significant IEG response in the medial amygdala in male mice, regardless of the biological relevance (or species of origin) of the chemical stimulus.

Immediate early gene expression in the medial amygdala in response to chemical signals was essentially unaffected by SHAM surgery. The only stimulus/area that exhibited a different pattern of IEG expression than in our previous report on intact animals (Samuelsen and Meredith, 2009) was a significant increase in IEG expression in the MePd upon exposure to mMU. In our previous report, the response in MePd was slightly higher, but not significantly different from control. This variation may be due to small individual differences in male mice and the larger number of mice in the previous experimental group. MePd is thought to be an important contributor to reproductive circuits in the hypothalamus (Choi et al., 2005), so an increase in FRAs expression to mMU is unexpected; although the phenotype and projections of the responding cells in each case are not yet known and may be different.

Similar to our demonstration here, Pankevitch et al. (2006) reported a decrease of medial-amygdala IEG-expression following VNX in male mice, but in response to the volatile components of fMU. In male hamsters, we have repeatedly found that VNX eliminates medial-amygdala IEG-response to HVF, an important chemosignal for hamsters (Fernandez-Fewell and Meredith, 1994; Westberry and Meredith, 2003b), as well as responses to other chemo-sensory stimuli (J.M. Westberry and M. Meredith, unpublished observations).

The behavioral responses of SHAM mice were essentially similar to those previously reported in intact male mice (Samuelsen and Meredith, 2009). However, when analyzed by two-way ANOVA, the higher levels of investigation of clean control swabs, compared to scented swabs, was not significant for SHAM mice. All tested mice appear to treat clean swabs as potential bedding material and spend time shredding the tip in a characteristic behavior. This behavior was not seen with scented swabs, in either SHAM or VNX mice. Thus, the increased time VNX mice spent investigating scented swabs was not due to time spent shredding. VNX mice clearly can detect/discover male and female mouse urine (Pankevitch et al., 2004, 2006) and probably all the stimuli used here, by olfaction. The increased investigation times after VNX may indicate an attempt to stimulate the VNO, whether to obtain olfactory information or for the proposed reinforcing properties of VNO input (Martinez-Ricós et al., 2007). These animals had no prior contact with any heterospecific stimuli or any of the donors of conspecific stimuli, but they had experience with urine of mature males and also with
urine of a mature female (prior to weaning). The increased investigation of the conspecific stimuli after VNX can be interpreted as a response to the novelty of an olfactory “signature” in the absence of the corresponding VNO input, that is a novel sensory experience for a stimulus otherwise belonging to a familiar class. This explanation does not immediately appear to be likely for the heterospecific stimuli. However, these are complex stimuli and it is possible that a heterospecific stimulus may contain some component in common with a salient chemosensory stimulus, which cannot be easily distinguished without vomeronasal input, thus eliciting additional investigation. Manipulation of a complex perceptual signature, by adding rather than subtracting components, may underlie the increase of aggression in male mice when androstenone, a pheromonal component of boar saliva, is added to a mouse urine stimulus. A similar increase of aggression does not occur when androstenone is presented in water (Ingersoll and Launay, 1986).

Under many circumstances, in sexually naïve hamsters and mice, previous experiments have shown that a functional VNO is necessary for normal social (Wekesa et al., 1994; Stowers et al., 2002; Kobayakawa et al., 2007) and sexual behaviors (Meredith, 1986; Keller et al., 2006; Kimchi et al., 2007) as well as preferences for social odors (Johnston and Peng, 2000; Pankevich et al., 2004, 2006; Woodley et al., 2004). However, a recent report suggests that female mating behavior is not affected by VNX (Martel and Baum, 2009).

In this experiment, we report that the behavioral response to a predator odor is disrupted by VNX. VNX mice spent significantly more time investigating a piece of worn CC as compared to both VNX mice exposed to a clean collar control and SHAM mice exposed to worn CC. In a previous experiment (Samuelsen and Meredith, 2009), intact males, as sham-operated males here, appeared to avoid worn CC pieces. Both intact and SHAM mice use the “stretch-attend” posture (Yang et al., 2004) in investigating CC stimuli. The intact animals spent significantly less time investigating CC stimuli than clean collar control stimuli, although, as with sham-operated animals here, there was no difference in the number of investigatory contacts. Although the SHAM mice had much shorter investigation times for worn CC, when analyzed in a two-way ANOVA with the VNX data, they were not significantly different from investigation of clean collars. This difference was significant if the SHAM data were analyzed separately by one way ANOVA, as in the previous experiment with intact animals. The two-way analysis does confirm a significant difference between investigation time for SHAM and VNX animals. The behavioral response of VNX mice suggests that they do not recognize the biological relevance of these chemical signals.

An overall increase in IEG expression in MeP, as well as in MeA, is characteristic of all biologically relevant stimuli tested, in both mice (Samuelsen and Meredith, 2009) and hamsters (Meredith and Westberry, 2004). While each biologically relevant chemosensory stimulus carries a potentially different message, the medial amygdala response is “categorical” because the presence or absence of an overall MeP response separates clearly relevant stimuli from those with no obvious biological relevance. Stimuli with no clear biological relevance, which evoke IEG expression only in MeA, include, in hamsters; artificial activation of amygdala input (Meredith and Westberry, 2004; Nolte and Meredith, 2005) as well as male and female mouse signals (Meredith and Westberry, 2004). In mice, they include steer urine and hamster stimuli (Samuelsen and Meredith, 2009). All the stimuli used here are unlearned in that the animals had no previous experience with the individual stimulus donors, but they may be familiar with male and female mouse odors as categories of stimuli (see above). An interesting next step will be to examine how the medial amygdala response is modified by experience with particular individual signals.

Removal of vomeronasal sensory input to medial amygdala is one obvious consequence of VNX, but a loss of subsequent hormonal responses could also contribute to the results observed here. For example, intact, but not VNX, male mice increase LH secretion in response to female urine (Coquelin et al., 1984), presumably a result of increased Gonadotropin-releasing hormone (GnRH) release into the pituitary portal vessels. There is a subsequent increase in circulating testosterone over the next half hour (Wysocki et al., 1983). This 30 min delay means testosterone changes are not major factors in the behavioral responses and probably not in IEG responses reported here. However, GnRH is also released into the brain and intraventricular infusion of GnRH can alter behavior (Fernandez-Fewell and Meredith, 1995) as well as IEG responses in the amygdala (Westberry and Meredith, 2003a). Endogenous GnRH release directly at the site of action in the brain should be faster-acting than exogenous GnRH infused in the cerebral ventricles, and could be fast enough to feed-back onto amygdala circuits and influence results here. There is also evidence for estrogen effects on conspecific-signal recognition and behavioral response to predator odors (Pierman et al., 2006; Kavaliers et al., 2008), so a rapid change in aromatization of testosterone to estrogen could also influence brain circuits. Changes in steroid receptor function within the time course of these experiments are also possible (Blake and Meredith, unpublished observation).

The categorization of biologically relevant conspecific and heterospecific chemical signals in the mouse medial amygdala, whether learned or unlearned, depends upon a functional VNO. Even though there are direct projections from the main olfactory bulb to the medial amygdala (Prosistaiga et al., 2007; Kang et al., 2009), our results suggest that main olfactory input alone is not sufficient to activate the characteristic categorical IEG expression pattern of the medial amygdala to biologically relevant chemical signals.

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REFERENCES


APPENDIX

Supplementary data

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