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Dual processing of sulfated steroids in the olfactory system of an anuran amphibian

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Chemical communication is widespread in amphibians, but if compared to later diverging tetrapods the available functional data is limited. The existing information on the vomeronasal system of anurans is particularly sparse. Amphibians represent a transitional stage in the evolution of the olfactory system. Most species have anatomically separated main and vomeronasal systems, but recent studies have shown that in anurans their molecular separation is still underway. Sulfated steroids function as migratory pheromones in lamprey and have recently been identified as natural vomeronasal stimuli in rodents. Here we identified sulfated steroids as the first known class of vomeronasal stimuli in the amphibian *Xenopus laevis*. We show that sulfated steroids are detected and concurrently processed by the two distinct olfactory subsystems of larval *Xenopus laevis*, the main olfactory system and the vomeronasal system. Our data revealed a similar but partially different processing of steroid-induced responses in the two systems. Differences of detection thresholds suggest that the two information channels are not just redundant, but rather signal different information. Furthermore, we found that larval and adult animals excrete multiple sulfated compounds with physical properties consistent with sulfated steroids. Breeding tadpole and frog water including these compounds activated a large subset of sensory neurons that also responded to synthetic steroids, showing that sulfated steroids are likely to convey intraspecific information. Our findings indicate that sulfated steroids are conserved vomeronasal stimuli functioning in phylogenetically distant classes of tetrapods living in aquatic and terrestrial habitats.

**Keywords:** intraspecific chemical communication, *Xenopus laevis*, pipidae, main olfactory system, vomeronasal system

**Introduction**

The mammalian olfactory system is typically defined by at least two anatomically and molecularly segregated subsystems, the most prominent being the main and vomeronasal system. The main olfactory system generally senses volatile chemicals primarily signaling food and environmental cues, whereas the vomeronasal system mainly detects pheromones, which signal intraspecific social
cues (Liberles, 2014). Several recent studies, however, showed that this separation may not be as strict as originally thought (Zufall and Leinders-Zufall, 2007). Fishes generally have a single olfactory system. Characteristic morphological sensory neuron types, as well as genetic components of both the mammalian main olfactory epithelium (MOE) and vomeronasal organ (VNO) coexist in their unique olfactory epithelium. Accordingly, it mediates all of the behavioral responses known to be associated with the sense of smell (Hamdani and Doving, 2007).

Amphibians are transitional species, typically having aquatic larvae and terrestrial or secondarily aquatic adults. Most amphibians, including *Xenopus laevis* (see Dittrich et al., 2015), have anatomically segregated main and vomeronasal systems (Eisthen and Polese, 2006). However, at a cellular and molecular level the systems are not yet fully separated. The MOE of larval *Xenopus* still closely resembles the single olfactory epithelium of fishes. It is made of ciliated as well as microvillous sensory neurons expressing olfactory receptor genes of all main families (Hansen et al., 1998; Hagino-Yamagishi et al., 2004; Date-Ito et al., 2008; Gliem et al., 2013; Syed et al., 2013). Also, at least two different transduction pathways are present (Gliem et al., 2013; Sansone et al., 2014a,b). Conversely, the larval VNO is already very similar to that of rodents, comprising microvillous neurons (Hansen et al., 1998), which express solely vomeronasal type-2 receptor genes (Hagino-Yamagishi et al., 2004; Syed et al., 2013; Sansone et al., 2014b).

Many general mammalian odorants, mostly volatile chemicals, have been identified. The number of known mammalian pheromones, many of them urinary constituents, is much lower (Liberles, 2014). Sulfated steroids have been reported to be the predominant vomeronasal ligands in female mouse urine (Nodari et al., 2008). The olfactory system of fishes detects general odorants and substances with pheromonal activity (Bazáes et al., 2013). Some sulfated steroids function as migratory pheromones in lamprey (Sorensen et al., 2005). In aquatic amphibians (larval and adult) the main olfactory system is generally sensitive to the same odorant classes as in fishes. Terrestrial species typically sense the same odorants as mammals (Eisthen and Polese, 2006). Pheromonal communication is relatively widespread in caudate amphibians, however, much less is known in anurans, where acoustic communication is thought to be the main mode of intraspecific communication (Woodley, 2014).

Here we report that the same synthetic sulfated steroids active in rodents, elicit strong olfactory responses in both the main and vomeronasal system of larval *Xenopus laevis*. Sulfated steroids thus appear to be conserved vomeronasal stimuli in phylogenetically distant classes of tetrapods. Analysis of neuronal responses revealed similarities and differences in the processing of these stimuli in the two subsystems. Furthermore, mass spectrometry analysis revealed multiple sulfated compounds in *Xenopus* excretions with mass-to-charge ratios consistent with sulfated steroids. The fact that these excretion products activated a substantial subset of sensory neurons responsive to the synthetic compounds indicates that sulfated metabolites are involved in amphibian intraspecific communication.

## Materials and Methods

### Confocal Ca$^{2+}$ Imaging in Acute Slices of the Olfactory Organ

Slices (130–150 µm) of the olfactory organ of larval *Xenopus laevis* (either sex, stages 50–54; see ref. Nieuwkoop and Faber, 1994) were prepared and loaded with Fluo-4 AM (Life Technologies) as described in earlier work of our lab (Sansone et al., 2014a). All procedures for animal handling were approved by the governmental animal care and use office (Niedersächsisches Landesamt für Verbraucherschutz und Lebensmittelsicherheit, Oldenburg, Germany, Protocol No. T24.07) and were in accordance with the German Animal Welfare Act as well as with the guidelines of the Göttingen University Committee for Ethics in Animal Experimentation. Reproducibility of stimulus-induced responses was verified by regularly repeating the application at least twice. As a negative control we regularly applied standard bath solution containing the same DMSO concentration as the stimuli. The minimum interstimulus interval was at least 2 min. Changes of intracellular calcium concentrations were monitored at 1 Hz using an inverted or upright laser-scanning confocal microscope (LSM 510/Axiovert 100 M or LSM 780/Axio Examiner, Carl Zeiss).

### Multi-photon Ca$^{2+}$ Imaging in the Whole Mount Olfactory Bulb

To image activity in the olfactory bulb a bolus injection of Fluo-4 AM was performed. For orientation during this procedure we previously electroporated neurons in the MOE and VNO with Alexa-594 dextran (for details see Hassenkläver and Manzini, 2013, 2014). After killing the animals the connective tissue covering the ventral side of the telencephalon was removed using fine scissors and the preparation was mechanically fixed in a recording chamber and covered with bath solution. A solution containing a saturated solution of Fluo-4 AM was filtered, centrifuged, and then the supernatant was used to fill borosilicate pipettes (resistance ~10–15 MΩ). The pipette, controlled via a micromanipulator, was carefully introduced into the olfactory bulb in close proximity of the axon terminals and the Fluo-4 AM solution was ejected by pressure application. After 45 min incubation at room temperature, post-synaptic Ca$^{2+}$-responses were imaged using a multi-photon microscope (Nikon A1R-MP, Nikon). Fluorescence image stacks containing glomeruli and neurons of the main and accessory olfactory bulb were acquired with a fast resonant scanning unit. Image stacks were acquired at 0.7–1 Hz with an interval of 3–5 µm between the acquired planes.

### Analysis of Ca$^{2+}$ Imaging Data

Fluorescence changes for individual regions of interest, i.e., individual neurons or glomeruli, are given as ∆F/F values. The fluorescence changes were calculated as ∆F/F = (F − F₀)/F₀, where F was the fluorescence averaged over the pixels of a sensory neuron, while F₀ was the average fluorescence of that sensory neuron prior to stimulus application. A response was assumed if the following criteria were met: (i) the maximum amplitude of the calcium transient had to be higher than the maximum of the prestimulus intensities; (ii) the onset of the...
response had to be within 10 frames after stimulus application. The analysis of confocal imaging data was carried out as described in earlier work of our lab (Sansone et al., 2014a). For dose-response analysis, responses to each concentration were pooled and fitted using GraphPad Prism (GraphPad Software). Detection threshold for every cell was determined as the lowest concentration at which a response meeting above criteria occurred. The overlaps of reactivity between the single components of sulfated steroid mixtures were visualized as area-weighted Venn diagrams created with Python/matplotlib-venn. To determine reactive regions in the multi-photon imaging data, difference image stacks were calculated by subtracting the mean background activity before stimulus application from the mean peak response. Difference image stacks were converted to binary data by adaptive thresholding. Mean traces of calcium transients were extracted from volumetric regions of interest derived from these binary image stacks. For visualization, the binary data was smoothed and rendered as a 3D volume. All the Ca$^{2+}$ imaging data were analyzed using custom-written programs in MATLAB (Mathworks, Natick, USA).

### Olfactory Stimuli and Solutions

All the stimuli were dissolved in bath solution or DMSO according to the specific solubility and were applied at a final concentration of 100–200 μM (additional dilutions were used in dose-response experiments). The stimuli used were: sulfated steroids (Steraloids, see Supplementary Table 1), non-sulfated steroids (β-estradiol, pregnanolone, allopregnanolone, progesterone; Sigma); a mixture of 19 amino acids (Sigma; Manzini and Schild, 2004), tadpole and adult frog breeding water (diluted 1:1 in bath solution). Sulfated steroids were applied as mixtures in some of the experiments (E mix: E0588, E1050, E2734; P mix: P3817, P8168, P8200). Amino acids are well characterized odorants in *Xenopus laevis* (Manzini and Schild, 2004; Glicen et al., 2013; Sansone et al., 2014a). To collect excretion products, we kept either 80–100 tadpoles or 10–20 post-metamorphic frogs in a beaker filled with 500 ml tap water for 12–24 h. After removing the animals, the water was filtered with a Falcon cell strainer (40 μm nylon mesh; BD Biosciences) to remove particles and stored at +4°C. Bath solution consisted of (in mM): 98 NaCl, 2 KCl, 1 CaCl$_2$, 2 MgCl$_2$, 5 glucose, 5 Na-pyruvate, 10 HEPES. The composition of the Ca$^{2+}$-free bath solution was (in mM): 98 NaCl, 2 KCl, 2 MgCl$_2$, 5 glucose, 5 Na-pyruvate, 10 HEPES, 2 EGTA. High K$^+$ bath solution consisted of (in mM): 17 NaCl, 80 KCl, 1 CaCl$_2$, 2 MgCl$_2$, 5 glucose, 5 Na-pyruvate, 10 HEPES. All bath solutions were adjusted to pH 7.8 and had an osmolarity of 230 mOsmol/l.

### Chromatography and Mass Spectrometry of Animal Breeding Water

Solid-phase chromatography was performed on tadpole and adult frog breeding water (collection as described above), according to a previously employed method for steroids extraction from urine samples (Shackleton and Whitney, 1980; Nodari et al., 2008). Briefly, a Sep-pak C-18 column (Waters Corp) was primed with one volume methanol, followed by one volume 20% methanol/2% acetic acid. After washing with two volumes water, 100 ml of sample (either tadpole or adult frog breeding water) were passed through the column with a syringe, at a flow rate of ∼20 ml/min. Then, one volume 20% methanol/2% acetic acid was passed though the column, followed by 2 ml methanol elution to recover the steroids and steroid conjugates. An API 3000 Triple Quad Mass Spectrometer with an ESI spray ion source in a negative ion mode was then used to analyze the C-18 extracts from adult and tadpole breeding water. The methanol extract was infused at 10 μl/min into the ESI source by syringe pump. The electrospray needle was set at 4.5 kV and the ion source temperature was 180°C. Nitrogen was used as the collision gas. Mass spectrum was collected by precursor ion scan and the collision energy was set at 45 eV for the collision cell (Q2).

### Results

**Sulfated Steroids Activate Sensory Neurons in the Main and the Vomeronasal Epithelium of *Xenopus laevis***

Sulfated steroids from the androgen, estrogen, pregnanolone and glucocorticoid families, potent murine vomeronasal ligands (see Supplementary Table 1 and Nodari et al., 2008; Isogai et al., 2011; Celsi et al., 2012), were tested for their chemosensory activity in larval *Xenopus laevis*. Responses were measured as stimulus-induced calcium increases of sensory neurons in acute slice preparations of the olfactory organ (Figures 1A,B). We found steroid-responsive sensory neurons in the VNO and the MOE (Figure 1C). Neurons responded to pregnanolone-derived (P mix) and estrogen-derived (E mix) steroids (for more information see Materials and Methods). Thereby, in both epithelia P mix responses were more frequent than E mix responses. None of the tested androgen-derived and glucocorticoid-derived steroids were effective (not shown). All cells that responded to sulfated steroids (VNO and MOE) also showed calcium transients upon stimulation with high K$^+$ bath solution, indicating that they were sensory neurons (Dittrich et al., 2014). In both epithelia, sulfated steroid-induced calcium responses were dependent on the presence of extracellular Ca$^{2+}$, indicating that the stimulus-induced intracellular Ca$^{2+}$ increase is mainly due to Ca$^{2+}$ entry through plasma membrane ion channels (Figure 2). Non-sulfated analogs of the employed steroids in most cases (>80%) failed to activate sulfated-steroid-responsive neurons (Figure 3).

### Olfactory Responses to Sulfated Steroids are Conveyed and Processed in the Olfactory Bulb

To gain information about the processing of sulfated steroids in the first relay center of the system, we recorded odorant-induced calcium responses in whole mounts of the olfactory system. Nasal application of P and E mix sulfated steroids elicited transient increases in intracellular Ca$^{2+}$ concentration in the main and accessory olfactory bulb. We detected responses in glomerular tufts and in somata of mitral/tufted cells (Figures 4A–C). Glomerular responses to the E mix were strongly overlapping with the P mix-responsive areas, but...
we also found regions reacting individually to each mixture (Figure 4A). Amino acid-evoked response patterns in the main olfactory bulb were also in part overlapping with the steroid-reactive areas (Figure 4B). Amino acids did not evoke any activity in the accessory olfactory bulb (Figures 4B,C). Signals detected in individual cell somata of the olfactory bulb showed various reactivity patterns to the different odorant stimuli (Figure 4C). Transection of the olfactory nerve abolished all olfactory bulb responses to odorant stimulation (data not shown).

**Tuning Properties of Sulfated Steroid Sensitive Neurons**

We applied the individual components of the P and E mix (see Materials and Methods and Supplementary Table 1) in addition to the mixtures to obtain response profiles of individual neurons of the VNO and the MOE (Figure 5A). Similar responses for the P mix sulfated steroids were obtained in cells of the MOE and the VNO, i.e., in both epithelia most sensory neurons responded to more than one compound of the mixture. This suggests that sensory neurons in both olfactory subsystems are broadly tuned to P mix sulfated steroids (Figures 5D,E). None of the P mix compounds was detected with much higher frequency than the others (Figure 5G). In contrast, the response profiles for E mix steroids in the MOE showed a different trend. One specific compound, i.e., E0588, activated the large majority of cells. The recorded responses to P8168 were concentration dependent and the dose-response curves were fitted by the Hill equation (Figure 5B). The calculated EC_{50} values and Hill slopes for the two epithelia were slightly different (EC_{50}: MOE, 86 µM, VNO, 49 µM; Hill slope: MOE, 1.10, VNO, 1.87). The detection thresholds, i.e., the sensitivity, for P8168 of individual sensory neurons in the MOE and the VNO were significantly different (Figure 5C; Chi-squared test, **p < 0.01). On average, sensory neurons in the MOE were more sensitive to P8168 than those in the VNO. Specifically, in the MOE the majority of the tested sensory neurons had a threshold in the range of 1–10 µM, whereas the majority of VNO neurons had thresholds in the range of 10–50 µM (for exact numbers see legend of Figure 5).

**Sensory Neurons in the Main and the Vomeronasal Epithelium Respond to the Same Sulfated Steroid with Different Sensitivity**

To gain insight in possible sensitivity differences to sulfated steroids in sensory neurons of the VNO and MOE, we determined dose-response curves of individual cells for P8168, a single steroid of the P mix. The recorded responses to P8168 were concentration dependent and the dose-response curves were fitted by the Hill equation (Figure 5B). The calculated EC_{50} values and Hill slopes for the two epithelia were slightly different (EC_{50}: MOE, 86 µM, VNO, 49 µM; Hill slope: MOE, 1.10, VNO, 1.87). The detection thresholds, i.e., the sensitivity, for P8168 of individual sensory neurons in the MOE and the VNO were significantly different (Figure 5C; Chi-squared test, **p < 0.01). On average, sensory neurons in the MOE were more sensitive to P8168 than those in the VNO. Specifically, in the MOE the majority of the tested sensory neurons had a threshold in the range of 1–10 µM, whereas the majority of VNO neurons had thresholds in the range of 10–50 µM (for exact numbers see legend of Figure 5).

**Chromatography and Mass Spectrometry Reveal the Presence of Sulfated Compounds in Xenopus Breeding Water**

Sulfated steroids were shown to be excreted in mouse urine and to activate receptor neurons of the VNO (Hsu et al., 2008; Nodari et al., 2008). To obtain some insight on whether *Xenopus laevis* also excretes sulfated steroids, we performed steroid extraction from tadpole and adult frog breeding water via solid-phase chromatography. A well-established protocol for steroid extraction from urine samples (Shackleton and Whitney, 1980; Nodari et al., 2008) was employed to recover and concentrate free steroids and steroid conjugates contained in the breeding water. These extracts were then analyzed by precursor ion mass spectrometry (see Materials and Methods) for ions that could dissociate to produce fragments with a mass-to-charge ratio.
This analysis revealed the presence of multiple sulfated compounds (200 µM) are shown. All tested neurons, in both MOE and VNO, showed no response after 3 min incubation in Ca\(^{2+}\)-free bath solution. After 3 min of washout, all responses recovered. Mean calcium responses (±SEM), expressed as percentage of control (MOE/P8168, 9 cells, 2 slices; VNO/P8168, 9 cells, 2 slices; MOE/EO588, 9 cells, 6 slices) are shown in standard (colored columns) and Ca\(^{2+}\)-free bath solution (gray columns; 3 min after application of Ca\(^{2+}\)-free bath solution). All responses were virtually abolished in Ca\(^{2+}\)-free bath solution (*p < 0.01; paired Student’s t-test).

Sulfated steroids are urinary excretion products known to act as natural vomeronasal ligands in mice (Nodari et al., 2008; Isogai et al., 2011; Celsi et al., 2012). They account for up to 80% of the VNO neuronal responses by female urine and are thought to signal information about the sex and physiological status (Nodari et al., 2008). The present study now demonstrates that sulfated steroids are also detected by the olfactory system of *Xenopus laevis*. Interestingly, sulfated steroid are sensed and processed by both the main and the vomeronasal system in *Xenopus*, providing an unprecedented evidence of parallel processing of the same olfactory cues in two amphibian olfactory subsystems. Moreover, we found that *Xenopus* excretions contain sulfated compounds that activate a large subset of sulfated steroid-sensitive sensory neurons. This finding supports the idea that sulfated steroids could mediate intraspecific olfactory communication in amphibians.

It has long been assumed that the main and the vomeronasal systems are sensitive to different sets of chemosensory signals. While the main system was thought to solely detect common odors, the vomeronasal system was believed to specifically detect pheromones (Liberles, 2014; but see Hudson and Distel, 1986). Studies in rodents have reported that both systems respond to overlapping sets of stimuli (Sam et al., 2001; Spehr et al., 2006b; Zufall and Leinders-Zufall, 2007). Common odors as well as pheromones have been reported to activate both systems (Leinders-Zufall et al., 2004; Xu et al., 2005; Spehr et al., 2006a). The present study shows that the same set of molecular cues, i.e., sulfated steroids, are detected by two olfactory subsystems in an amphibian species. The frequency of P and E mix-sensitive cells in the MOE and the VNO of *Xenopus* varies considerably. In both epithelia P mix-sensitive cells were more frequent than E mix-sensitive cells. Especially in the VNO the E mix-sensitive cells are a rare subpopulation. These differences are reflected also.

**Discussion**

Sulfated steroids are urinary excretion products known to act as natural vomeronasal ligands in mice (Nodari et al., 2008; Isogai et al., 2011; Celsi et al., 2012). They account for up to 80% of the VNO neuronal responses by female urine and are thought to signal information about the sex and physiological status (Nodari et al., 2008). The present study now demonstrates that sulfated steroids are also detected by the olfactory system of *Xenopus laevis*. Interestingly, sulfated steroids are sensed and processed by both the main and the vomeronasal system in *Xenopus*, providing an unprecedented evidence of parallel processing of the same olfactory cues in two amphibian olfactory subsystems. Moreover, we found that *Xenopus* excretions contain sulfated compounds that activate a large subset of sulfated steroid-sensitive sensory neurons. This finding supports the idea that sulfated steroids could mediate intraspecific olfactory communication in amphibians.
in the olfactory bulb responses. Mucosal application of P mix activated larger bulb areas than the E mix in both the main and the accessory olfactory bulb (Figure 4A). Furthermore, response profile analysis for the single components of the mixtures highlighted differences in tuning properties of individual neurons (see Figures 5D–J). Sensory neurons in both epithelia were broadly tuned to all single components of the P mix. E mix-sensitive cells in the MOE were rather narrowly tuned to one component of the mixture (E0588). The low frequency of E mix-sensitive neurons in the VNO prevented the recording of response profiles to steroids of this group. The low selectivity of the P mix neurons might be due to high structural similarity among the mixture components. A similar broad tuning of individual P steroid-sensitive VNO neurons has been reported in mouse (Nodari et al., 2008; Meeks et al., 2010; Turaga and Holy, 2012). Interestingly, these findings are in contrast with results of another study with isolated mice VNO neurons, where all P sulfated steroid-sensitive vomeronasal neurons were narrowly tuned to one steroid (Celsi et al., 2012). E mix-responsive VNO neurons are rather narrowly tuned also in mice (Nodari et al., 2008).

The dose-response curves and threshold values for P8168 (VNO and MOE) obtained in Xenopus were in the same range as those obtained in VNO neurons of mice (Nodari et al., 2008; Celsi et al., 2012). Rather unexpectedly the thresholds in the MOE of Xenopus were significantly lower than in the VNO. These differences in sensitivity could indicate that the activation of both systems is not just redundant, but rather signal different information, possibly triggering different behavioral responses. A similar situation is present in the mouse olfactory system. Major histocompatibility complex peptides, chemosensory signals that carry information about individuality (Leinders-Zufall et al., 2004; Spehr et al., 2006a), mediate different behavioral responses via the main and accessory system (Kelliher, 2007).

In Xenopus laevis, strikingly different than in later diverging vertebrates such as rodents, vomeronasal type-1 and more “ancestral” vomeronasal type-2 receptor genes are expressed in the MOE (Date-Ito et al., 2008; Gliem et al., 2013; Syed et al., 2013), together with OR-type and trace amine associated receptor genes (Mezler et al., 1999; Gliem et al., 2013). Sensory neurons of the Xenopus VNO exclusively express later diverging vomeronasal type-2 receptor genes (Hagino-Yamagishi et al., 2004; Syed et al., 2013). This allows one to speculate about the receptor gene families involved in the detection of sulfated steroids in the two olfactory subsystems of Xenopus. It appears very likely that the steroid responses in the VNO are mediated by vomeronasal type-2 receptors. This receptor family has been reported to be involved in the detection of P steroid responses in the rodent accessory olfactory system (Hammen et al., 2014). However, the MOE responses could be generated by any of the receptor families expressed in this epithelium. In the mouse VNO, E steroids have been shown to activate
FIGURE 4 | Sulfated steroids are processed in the olfactory bulb. (A) 3D rendering of reactive volumes in the MOB (left hand side) and AOB (right hand side) upon mucosal application of E and P mix (E mix-responsive areas, blue; P mix-responsive areas, red). Active regions always contained both cell somata and glomerular structures. Schematics of the olfactory bulb in the upper part show the approximate location of the responsive regions. (B) Calcium responses of the glomerular regions indicated by an arrow and green color in A (MOB, green traces; AOB, magenta traces). Amino acid mixture (AA, 100 µM) was applied as a positive control for activation of the MOB. (C) Representative calcium responses of mitral/tufted cells in the MOB (green) and AOB (magenta) upon mucosal application of sulfated steroids (E mix and P mix, 200 µM) and amino acids (AA, 100 µM). In both main and accessory olfactory bulb, mitral/tufted cells were activated by either E or P steroids, or both stimuli. Amino acids only elicited calcium responses in mitral/tufted cells of the MOB. Similar results were obtained in 4 MOBs and 3 AOBs.

vomeronasal type-1 receptors (Isogai et al., 2011), therefore it appears plausible that at least some responses in the Xenopus MOE are mediated by this receptor family. Together, the available data about receptor expression in the two epithelia of Xenopus suggests that different sets of receptors are involved in the steroid detection in the MOE and the VNO. This is supported also by the differences in the dose-response curves and the different thresholds for P8168 in the two epithelia reported in the present study. Our results show that sulfated steroids appear to be a class of conserved chemosensory stimuli in the tetrapod lineage. The evolutionary relevance of these stimuli is further supported by the fact that sulfated steroids (though different than those used in the present study) serve as migratory pheromones in lamprey, a primitive jawless fish (Sorensen et al., 2005). The information about intraspecific chemical communication in adult anurans is very limited, and virtually nothing is known for larval anurans (Belanger and Corkum, 2009; Houck, 2009; Poth et al., 2012; Woodley, 2014). The presence of sulfated compounds, possibly sulfated steroids, in excretion products of larval and adult Xenopus indicates that they...
could serve as intraspecific communication cues. The fact that the majority of neurons sensitive to synthetic sulfated steroids also respond to breeding water is certainly in line with this speculation.

**Conclusions**

The available functional data on the amphibian accessory olfactory system is very scarce if compared to mammals. In particular, a physiological characterization of vomeronasal responses in anuran amphibians was never performed, due to the lack of known stimuli. In this study, we identified sulfated steroids as a novel class of amphibian olfactory stimuli, which also activate the vomeronasal organ. Sulfated steroids are known to function as migratory pheromones in lamprey and have recently been identified as the predominant vomeronasal ligands in female mouse urine. In mice they are thought to transmit information about the animal’s physiological status, although a clear behavioral output has not been identified yet. Our data show that sulfated steroids are potent stimuli of both the main and the vomeronasal system of the amphibian *Xenopus laevis*. Moreover, breeding water containing sulfated compounds, with mass-to-charge ratios consistent with sulfated steroids, activated a large subset of steroid-sensitive sensory neurons, suggesting their involvement in amphibian intraspecific communication. Taken together, our study shows that sulfated steroids are conserved vomeronasal stimuli functioning in phylogenetically distant classes of tetrapods living in aquatic and terrestrial environments.
FIGURE 6 | Tadpole and frog breeding water contains sulfated compounds and activates neurons sensitive to synthetic steroids. (A) Following chromatographic extraction of steroids from breeding water, precursor ion scan mass spectrometry was used to analyze the abundance of peaks that fragment to produce negative ions with m/z of 80 (SO\(^-\), left column) or 97 (HSO\(^-\), right column). (B) Representative calcium responses of five MOE neurons. The shown cells include the main subgroups of responding cells, i.e., cells responding to both synthetic steroids (200 µM) and breeding water (1:1 dilution), cells responding to amino acids (100 µM) and breeding water, cells responding to breeding water only, and cells responding to synthetic steroids, breeding water and amino acids. (C) Response matrix of neurons in the MOE responding to breeding water (1:1 dilution) and synthetic sulfated steroids (200 µM; 85 cells, 5 slices). The majority of the neurons responding to synthetic steroids also responded upon application of breeding water (16 out of 27 cells). Response intensity is coded by a color gradient. A mixture of amino acids (AA, 100 µM) was applied as a control for slice viability. TW, tadpole breeding water; FW, frog breeding water; TapW, tap water control; ctrl, bath solution control.

habitats. Future studies will be necessary to elucidate whether sulfated steroid responses in fact trigger behavioral responses in larval (and adult) *Xenopus*, and if the different information channels (main and accessory olfactory system) elicit diverse behaviors.

**Author Contributions**

The experiments were conceived and designed by AS, TH, and IM. The experiments were performed by AS, TO and XF. AS, TH, TO, XF, and TEH analyzed the data. AS, TH, TEH and IM wrote the paper. All authors participated in the discussion of the data and in production of the final version of the manuscript.

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**Supplementary Material**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fncel.2015.00373

**References**

Bazáes, A., Olivares, J., and Schmachtenberg, O. (2013). Properties, projections, and tuning of teleost olfactory receptor neurons. *J. Chem. Ecol.* 39, 451–464. doi: 10.1007/s10886-013-0268-1

Belanger, R. M., and Corkum, L. D. (2009). Review of aquatic sex pheromones and chemical communication in anurans. *J. Herpetol.* 43, 184–191. doi: 10.1670/08-054R1.1

Celsi, F., D’Errico, A., and Menini, A. (2012). Responses to sulfated steroids of female mouse vomeronasal sensory neurons. *Chem. Senses* 37, 849–858. doi: 10.1093/chemse/bjs068
Mezler, M., Konzelmann, S., Freitag, J., Rössler, P., and Breer, H. (1999). Expression

Manzini, I., and Schild, D. (2004). Classes and narrowing selectivity of olfactory

Meeks, J. P., Arnson, H. A., and Holy, T. E. (2010). Representation and

Liberles, S. D. (2014). Mammalian pheromones.

Date-Ito, A., Ohara, H., Ichikawa, M., Mori, Y., and Hagino-Yamagishi, K. (2008).

Hamdani, E. H., and Døving, K. B. (2007). The functional organization

Hamnani, A., and Doving, K. B. (2007). The functional organization

Hansen, A., Reiss, J. O., Gentry, C. L., and Burd, G. D. (1998). Ultrastructure of the

Hsu, F. F., Nodari, F., Kao, L. F., Fu, X., Holekamp, T. F., Kao, L. F., Turk, J., et al. (2008). Sulfated steroids as natural ligands of mouse pheromone-sensing neurons. J. Neurosci. 28, 6407–6418. doi: 10.1523/JNEUROSCI.1425-08.2008

Poth, D., Wollenberg, K. C., Vences, M., and Schulz, S. (2012). Volatile amphibian pheromones: macrolides from mantellid frogs from Madagascar. Angew. Chem. Int. Ed. Engl. 51, 2187–2190. doi: 10.1002/anie.201106592

Sam, M., Vora, S., Malnic, B., Ma, W., Novotny, M. V., and Buck, L. B. (2001). Neuropharmacology. Odorants may arouse instinctive behaviours. Nature 412, 142. doi: 10.1038/35084137

Sansone, A., Hassenklöver, T., Syed, A. S., Korschig, S. I., and Manzini, I. (2014a). Phospholipase C and diacylglycerol mediate olfactory responses to amino acids in the main olfactory epithelium of an amphibian. PLoS ONE 9:e87721. doi: 10.1371/journal.pone.0087721

Sansone, A., Syed, A. S., Talantali, E., Korschig, S. I., and Manzini, I. (2014b). TrpC2 is expressed in two olfactory subsystems, the main and the vomeronasal system of larval Xenopus laevis. J. Exp. Biol. 217, 2235–2238. doi: 10.1242/jeb.103465

Shackleton, C. H., and Whitney, J. O. (1980). Use of Sep-pak cartridges for urinary steroid extraction: evaluation of the method for use prior to gas chromatographic analysis. Clin. Chim. Acta 107, 231–243. doi: 10.1016/0009-8981(80)90451-9

Sorensen, P. W., Fine, J. M., Dvornikovs, V., Jeffrey, C. S., Shao, F., Wang, J., et al. (2005). Mixture of new sulfated steroids functions as a migratory pheromone in the sea lamprey. Nat. Chem. Biol. 1, 324–328. doi: 10.1038/nchembio739

Spehr, M., Kellhner, K. R., Li, X. H., Boehm, T., Leinders-Zufall, T., and Zufall, F. (2006a). Essential role of the main olfactory system in social recognition of major histocompatibility complex peptide ligands. J. Neurosci. 26, 1961–1970. doi: 10.1523/JNEUROSCI.4939-05.2006

Spehr, M., Spehr, J., Ukhanov, K., Kellhner, K. R., Leinders-Zufall, T., and Zufall, F. (2006b). Parallel processing of social signals by the mammalian main and accessory olfactory systems. Cell. Mol. Life Sci. 63, 1476–1484. doi: 10.1007/s00018-006-6109-4

Syed, A. S., Sansone, A., Nadler, W., Manzini, L., and Korschig, S. I. (2013). Ancestral amphibian v2rs are expressed in the main olfactory epithelium. Proc. Natl. Acad. Sci. U.S.A. 110, 7714–7719. doi: 10.1073/pnas.1302088110

Tamura, D., and Holy, T. E. (2012). Organization of vomeronasal sensory coding revealed by fast volumetric calcium imaging. J. Neurosci. 32, 1612–1621. doi: 10.1523/JNEUROSCI.5339-11.2012

Weidolf, L. O., Lee, D. E., and Henion, J. D. (1988). Determination of boldenone sulfonconjugate and related steroid sulfates in equine urine by high-performance liquid chromatography/tandem mass spectrometry. Biomed. Environ. Mass Spectrom. 15, 283–289. doi: 10.1002/bms.1200150508

Woodley, S. K. (2014). “Chemical signaling in amphibians,” in Neurobiology of Chemical Communication, ed C. Muscignat-Caretta (Boca Raton, FL: CRC Press), 255–284.

Xu, F., Schaefer, M., Kida, I., Schaefer, J., Liu, N., Rothman, D. L., et al. (2005). Simultaneous activation of mouse main and accessory olfactory bulb by odors or pheromones. J. Comp. Neurol. 489, 491–500. doi: 10.1002/cne.20652

Yan, Y., Rempel, D. L., Holy, T. E., and Gross, M. L. (2014). Mass spectrometry of these terms.

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