The Vomeronasal Organ Mediates Interspecies Defensive Behaviors through Detection of Protein Pheromone Homologs

Fabio Papes,1,2,3 Darren W. Logan,1,3 and Lisa Stowers1,*
1Department of Cell Biology, The Scripps Research Institute, 10550 North Torrey Pines Road, La Jolla, CA 92037, USA
2Department of Genetics and Evolution, State University of Campinas, Campinas SP 13083-970, Brazil
3These authors contributed equally to this work
*Correspondence: stowers@scripps.edu
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SUMMARY

Potential predators emit uncharacterized chemosignals that warn receiving species of danger. Neurons that sense these stimuli remain unknown. Here we show that detection and processing of fear-evoking odors emitted from cat, rat, and snake require the function of sensory neurons in the vomeronasal organ. To investigate the molecular nature of the sensory cues emitted by predators, we isolated the salient ligands from two species using a combination of innate behavioral assays in naive receiving animals, calcium imaging, and c-Fos induction. Surprisingly, the defensive behavior-promoting activity released by other animals is encoded by species-specific ligands belonging to the major urinary protein (Mup) family, homologs of aggression-promoting mouse pheromones. We show that recombinant Mup proteins are sufficient to activate sensory neurons and initiate defensive behavior similarly to native odors. This co-option of existing sensory mechanisms provides a molecular solution to the difficult problem of evolving a variety of species-specific molecular detectors.

INTRODUCTION

The ability of prey to innately recognize the odor of a potential predator provides a strong selective advantage; however, the neural mechanisms that permit chemical eavesdropping on other species, interpret the cues, and initiate a defensive response are unknown. Inbred rodents, which have been isolated in the laboratory from other species for hundreds of generations, are known to respond with a fear-like defensive behavior to cat odors (Dielenberg et al., 2001; Dielenberg and McGregor, 2001; Takahashi et al., 2005; Vyas et al., 2007). This innate response suggests that the neural mechanisms of detection in the receiving animals are genetically determined. Evolving an innate capacity to respond to chemosignals from a variety of species is a mechanistic challenge. To maximize the specificity of the warning, the receiver may sense potential threats by detecting specific ligands from all other animals. Given the sensory circuitry needed to detect and process each cue and the probability that each individual may only encounter a small subset of potential predator odors in its lifetime, this strategy would require a significant genetic investment that may go largely unutilized. An alternate, simpler, mechanism may involve other animals emitting a common odor, perhaps as a consequence of carnivore metabolism, that activates a general predator sensory circuit in the receiving prey (Fendt, 2006). Identifying the signaling ligands from multiple distantly related species is an essential step toward elucidating general mechanisms generating interspecies communication.

Kairomones, such as those that elicit fear behavior, are cues transmitted between species that selectively disadvantage the signaler and advantage the receiver (Wyatt, 2003). Known kairomones have mainly been identified in insect communication, although these models have not provided insight into the organization of the neural response in the receiving animals (Stowe et al., 1995). It is thought that subsets of sensory neurons are genetically determined to mediate innate behavior. These are likely to be distinguished from canonical olfactory neurons by distinctive locations in the nasal cavity, alternate projections to the brain, and expression of atypical molecular features. The vomeronasal organ (VNO), a specialized chemosensory epithelium of terrestrial vertebrates, contains sensory neurons displaying all three of these unique olfactory characteristics and is confirmed to function in the detection of pheromones (Tirindelli et al., 2009). In addition to detecting pheromones, VNO neurons have been shown to respond to regular chemical odorants in vitro, but the biological significance of this activity has not been determined (Sam et al., 2001; Trinh and Storm, 2003). In reptiles, the VNO initiates a defensive response to predators and facilitates the tracking of prey (Halpern and Frumin, 1979; Miller and Gutzke, 1999; Wang et al., 1993). In mammals, the identity of kairomones and detecting sensory neurons remains mostly unknown. An exception to this is trimethylthiazoline (TMT), the prominent pungent compound isolated from fox feces that causes aversion in rodents (Buron et al., 2007). Detection of TMT occurs through unidentified neurons in the main olfactory epithelium (MOE) (Kobayakawa et al., 2007). It is not known whether the MOE neural response is specific to TMT or...
represents a common model for the processing of other innate cues that promote interspecies behavior.

The identities of receptors and circuits that initiate innate behavior in response to olfactory ligands are largely unknown. The isolation of ligands of known function would provide the means to precisely stimulate brain circuits leading to specific behaviors. Recent progress has been made toward identifying the molecular nature of pheromone cues by purifying individual ligands. Small volatile molecules, sulfated steroids, peptides, and small proteins all display hallmarks of mammalian pheromones (Nodari et al., 2008; Tirindelli et al., 2009). Among these ligands, the major urinary proteins (Mups) are abundantly excreted (milligram quantities per milliliter) in mouse urine and are additionally secreted by mammary, salivary, and lachrymal glands (Finlayson et al., 1965; Szoka and Paigen, 1978). Mups emitted by mice have been demonstrated to act as pheromone carrier proteins, environmental pheromone stabilizers, and as genetically encoded pheromones themselves (Chamero et al., 2007; Hurst et al., 1998; Marchlewksa-Koj et al., 2000; Mucignat-Caretta et al., 1995). In several mammals (such as mouse, rat, horse, and lemur) there is evidence for lineage-specific Mup gene expansion consistent with a function in interspecies communication; however, genome analyses have shown that most other mammalian species encode a single ancestral Mup of unknown biological function (Logan et al., 2008; Mudge et al., 2008). Interestingly, these Mup orthologs are primary sources of animal allergens, indicating that they are both highly stable and eminently transmissible between species in the environment (Virtanen et al., 1999).

In order to investigate the neural code that warns of danger, we first devised a robust and quantifiable odor-based behavioral assay and then used a combination of genetic and cellular analyses to identify the responding sensory organ and neural activity. Importantly, we studied mouse odor responses to five different animal species (rat, cat, snake, rabbit, and mouse), which enabled us the comparative means to identify general mechanisms of kairomone information coding. We show that VNO-defective animals, TrpC2−/−, do not sense the olfactory ligands that initiate defensive behavior. We purified and identified the kairomone activities from rat and cat and found that they each encode species-specific Mup homologs. Previously, intraspecies Mups have been shown to function as pheromones (Chamero et al., 2007). Our findings suggest that the stabilization and expansion of Mup chemosensation resulted in the co-option of function to include both inter- and intraspecies communication.

RESULTS

The Vomeronasal Organ Mediates Predator Odor-Elicited Defensive Behavior

Field and laboratory studies have shown that predator odors elicit a range of defensive behaviors from prey species (Apfelbach et al., 2005; Blanchard et al., 2001; Dielenberg and McGregor, 2001). We first utilized a simple and robust odor-driven behavioral assay to quantify the effect of odors from multiple predator species on inbred C57BL/6J mice. All natural odors and control odors were presented in a single trial to naive mice not previously exposed to other species. Mice were analyzed to determine whether exposure to odors elicited the combination of three outputs: avoidance behavior; risk assessment behavior, which is a stereotypical cautious investigative approach characterized by a low-lying extended body posture (see Extended Experimental Procedures and Movie S1 available online for a detailed description of behavior analyses); and the release of the stress response adrenocorticotropic hormone (ACTH). Together, these defensive behaviors and neuroendocrine change are considered to model responses associated with fear in rodents (Apfelbach et al., 2005; Blanchard et al., 2001; Rosen, 2004). The novelty of control odors evoked investigation without significant risk assessment behavior or release of ACTH (Figures 1A and 1C). We next investigated the innate response to native odors obtained from three species that are natural mouse predators: cat (neck swab), snake (shed skin), and rat (urine). In contrast to the controls, we found that wild-type mice displayed significant innate avoidance and risk assessment behaviors as well as an increase in the stress hormone ACTH when exposed to odors from all three species (Figures 1A–1D; Figures S1A and S1B). When similarly assayed, another complex natural odor, rabbit urine, did not induce avoidance or risk assessment behavior (Figure 1E), nor did it increase ACTH levels (Figure S1C), indicating that the defensive responses we observed are not generally directed to all complex novel odor stimuli. Together, our behavioral assays confirm that mice show robust defensive behavior upon first encounter with complex natural odors from three diverse species. This innate response suggests that cognitive neurons that sense odors from multiple potential predators in the receiving animal are genetically hardwired to activate a fixed-action pattern of defensive behavior.

The neurons that eavesdrop on the presence of other species have not been identified, but are expected to be a novel subset of chemosensory neurons in the nasal cavity. Mammalian pheromones that mediate other innate behaviors, such as male-male aggression, are detected by the VNO (Chamero et al., 2007; Leybold et al., 2002; Stowers et al., 2002), and the snake VNO is required to sense and respond to predators (Miller and Gutzke, 1999); therefore, we next investigated the extent to which the mammalian VNO is involved in the innate response toward kairomones. We analyzed the ability of mice lacking TrpC2, the primary signal transduction channel of VNO sensory neurons (VNs), to detect and respond to predator cues. Strikingly, these mutant animals showed no significant defensive behavior responses to any of the three odors from other species (Figures 1B and 1D; Figures S1A and S1B). Instead, TrpC2−/− animals investigated predator odors similarly to control odors, behavior expected if they were unable to detect the ligand(s) that signals caution. To determine the sufficiency of olfactory cues in sensing other animals, we additionally investigated the response of wild-type and TrpC2−/− mutant animals in a more natural environment (Figure S1D). When placed in the presence of a live (anesthetized) rat, wild-type mice spent the majority of the assay retreating in a hiding box and displaying numerous risk assessment episodes. In contrast, the TrpC2−/− animals approached and investigated the rat without significant defensive behaviors (Figure S1D; Movie S2). Remarkably, this suggests that, in the context of our assays, other sensory cues from the
This prompted us to determine whether the VNO neurons in rodents (Kobayakawa et al., 2007). This analysis indicates that TMT does not transmit the same sensory information as cat, rat, and snake odors. Moreover, these additional assays with naphthalene and TMT further demonstrate that TrpC2−/− mutants have intact central circuits that are capable of generating a wild-type display of avoidance behavior. This indicates that the lack of a behavioral response from TrpC2−/− mutants to our kairomones is not due to developmental defects that may affect associated pathways in the brain necessary to execute behavior (Figure 1; Figure S1).

We used two different methods to investigate the extent to which VNs detect odors emitted between species. First, we determined that kairomones directly activate VNs by observing calcium influx in individual dissociated VNs in response to kairomone odors (Figure S1E). This analysis indicates that TMT does not transmit the same sensory information as cat, rat, and snake odors. Moreover, these additional assays with naphthalene and TMT further demonstrate that TrpC2−/− mutants have intact central circuits that are capable of generating a wild-type display of avoidance behavior. This indicates that the lack of a behavioral response from TrpC2−/− mutants to our kairomones is not due to developmental defects that may affect associated pathways in the brain necessary to execute behavior (Figure 1; Figure S1).

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the sensory dendrite, we additionally exposed freely moving behaving mice to predator odor, observed the defensive response, and confirmed corresponding neuronal activation in the VNO epithelium by immunostaining to detect increases in the expression of the immediate early gene c-Fos (Figure 2B) (Morgan and Curran, 1991). The punctate expression of c-Fos throughout the vomeronasal epithelium is consistent with specific activation of a subset of cognate sensory neurons. Together, these results indicate that mammalian VNs directly detect odors from a variety of other species.

In animals exposed to predator odors, we additionally observed a striking amount of c-Fos expression in the accessory olfactory bulb (AOB), to which VNO axons directly project (Figure 2C; Figures S2A–S2H). This activity was entirely absent in the TrpC2−/− mutants. In wild-type animals, immunostaining was concentrated in the granule cell layer, although snake odor reliably induced higher levels of activity and additionally evoked c-Fos expression in the glomerular and mitral cell layers shown to activate sensory neurons that project to either one of the AOB zones (Tirindelli et al., 2009). Interestingly, all three predator odors from our study induced robust activity in both zones, suggesting that they are composed of several ligands capable of activating distinct VNO receptors.

### Purification of a Single Ligand that Evokes Defensive Behavior

Natural animal secretions are typically odorous and expected to be composed of a complex blend of stimuli. We chose an unbiased method to purify and identify the behavior-inducing kairomone ligand(s). We sequentially fractionated the natural odor source and tracked the relevant bioactivity by behavioral analysis to identify kairomone-containing fractions. We first fractionated total rat urine over size-exclusion ultrafiltration columns and tested these fractions for defensive behavior-evoking activity. We found that the low molecular weight fraction (LMW; containing ligands of less than 10 kDa molecular mass)
entirely lacked kairomone activity in our assay, whereas the high molecular weight fraction (HMW; containing ligands of greater than 10 kDa molecular mass) was sufficient to induce prolonged avoidance and repeated risk assessment episodes, similar in quality and quantity to total rat urine (Figure 3A).

When we analyzed the AOB of animals exposed to these two fractions, we found limited c-Fos response in the anterior AOB (aAOB) to the LMW fraction and extensive activation in the granule cell layer of the aAOB to the HMW fraction, which additionally strongly activated the posterior AOB (pAOB) (Figures 3B and 3C). Interestingly, only whole urine and the HMW fraction stimulated detectable c-Fos immunoactivity in the mitral cell layer that contains the output neurons of the pAOB (Figures 3B and 3C), whereas very little c-Fos was detected in this region in response to urine stimuli that did not elicit defensive behavior (Figure S3). VNs that project to the pAOB are known to be activated by peptides and proteins (Munger et al., 2009), consistent with the large molecular mass of our kairomone-containing HMW fraction. Overall, both fractions contain ligands that activate the accessory olfactory system; however, the kairomone activity is found exclusively in the HMW fraction and the functional significance of those in the LMW fraction remains unknown. This finding underscores our strategy of utilizing behavioral analysis to purify ligands of known biological relevance.

To further determine the complexity and identity of potential ligands, we subjected the HMW fraction stimulus to electrospray ionization time-of-flight mass spectrometry (ESI-TOF-MS) and found both the HMW fraction and whole rat urine to contain one prominent protein with a molecular mass of 18,729 kDa as well as many less abundantly expressed proteins (Figure 4A). We used anion-exchange fast protein liquid chromatography (FPLC) to separate the HMW fraction components into 40 fractions over a 0–1 M NaCl gradient (Figure 4B). To streamline the identification of the bioactivity, we first screened the FPLC fractions for the ability to activate VNs as indicated by calcium imaging. Only those fractions containing the prominent protein which eluted between 300 and 450 mM NaCl induced calcium transients in the same neurons as those activated by the HMW fraction (Figures 4D and 4E; Figure S4B). When we analyzed the response at the single-cell level, the FPLC-A fractions were found to induce calcium transients in the same neurons as those activated by the HMW fraction (Figures 4D and 4E; Figure S4B). Importantly, these fractions were fully sufficient to stimulate
robust defensive behaviors when presented to predator odor naive mice and no behavior-promoting activity was present in other tested FPLC fractions (Figure 5A).

To identify the FPLC-A activity, we used nano-liquid chromatography MS/MS to obtain the sequence of the prominent protein in the behavior-promoting fractions. Interestingly, the resulting peptides identified the protein as an α-2,3-globulin (data not shown). On comparison with the rat genome, we resolved its sequence to the product of a specific gene: Mup13, a homolog of mouse Mup pheromones (Logan et al., 2008). The central hydrophobic binding pocket of all Mups creates a high affinity for small organic ligands, which are themselves known to have chemosignaling functions (Flower, 1996; Leinders-Zufall et al., 2000). To determine whether the kairome activity was produced by the presence of these protein-associated ligands, we incubated the HMW fraction with menadione, which competitively displaces potential rat Mup-bound ligands from the native Mup protein (Chamero et al., 2007). Naive mice responded with defensively displacing potential rat Mup-bound ligands from the native Mup protein (Chamero et al., 2007). Naive mice responded with complete defensive behavior toward menadione-displaced Mups, indicating that native small molecules which may be associated with the active protein fraction do not function as kairomones (Figure S5A). When considered with our fractionation data, it suggests that rat Mup13 protein found in urine is transmitted between species and generates neural activation of the vomeronasal system to signal fear. To validate these findings, we cloned, expressed, and purified a recombinant fusion maltose-binding protein (rMBP) in dissociated VNs isolated from wild-type (black bars) or TrpC2-/- (white bars) male mice and assayed by calcium imaging. The ordinate shows the normalized response compared to the rat HMW activation level.

n = 4–16 experiments. ***p < 0.001. ANOVA followed by Tukey-Kramer HSD post hoc analysis against no stimulus control (0.509% ± 0.177%; 0.256 normalized response).

(D) Representative calcium transients from eight isolated VNs, sequentially stimulated with rMBP, rMup-Rn13, the FPLC-A peak, and the HMW fractions of rat urine. Axis bars: X = 60 s; Y = 3×(F340/380 nm). Images of a representative responding cell are presented below the traces, pseudocolored dark to light to indicate calcium influx.

(E) Comparative percent activity of dissociated VNs stimulated with rMBP, rMup-Rn13, the FPLC-A peak, and the HMW fraction as assayed by calcium imaging. Each bar denotes the percentage of all imaged cells exhibiting a calcium spike in response to the stimuli marked with a plus sign and not exhibiting a response to the stimuli marked with a minus sign. All cells were exposed to all four stimuli, except for control cells, which were exposed to the indicated number of repetitive pulses of rMBP (white bars). Note the percentage of cells activated by all three rat stimuli (first bar), which is significantly above the number of cells responding to three pulses of control rMBP (second bar), and no cells responded to rat rMBP stimuli (third bar).

n = 10–11. ***p < 0.001. ANOVA followed by Tukey-Kramer HSD post hoc analysis against the respective rMBP control. Mean ± SEM. See also Figure S4.

Figure 4. Activation of Subsets of Vomeronasal Sensory Neurons by Purified Putative Kairomones

(A) ESI-MS analysis identifying the major protein constituent in the HMW of rat urine (arrowhead). (B) Further fractionation of the HMW by anion-exchange FPLC. Fractions from the shaded area were combined to form FPLC-A peak and bioactivity was compared to fractions gathered from the nonshaded areas (ctrl fractions).

(C) Quantification of response to fractions of rat HMW, recombinant rat Mup13 (rMup-Rn13), and recombinant maltose-binding protein (rMBP) in dissociated VNs isolated from wild-type (black bars) or TrpC2-/- (white bars) male mice and assayed by calcium imaging. The ordinate shows the normalized response compared to the rat HMW activation level.

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We analyzed the sensitivity of VNs to function as a ligand that directly activates VNO sensory neurons cellular calcium transients (Figure S4C). This concentration is relatively high levels of the ligand to be necessary to initiate intracellular calcium transients or c-Fos induction in the AOB to rMup-Rn13 (Figure 5A) and a significant increase in ACTH (Figure S5B). Together, these analyses demonstrate the sufficiency of rat Mup as a kairomone signal.

**Cat Mup Functions as a Kairomone**

We found it notable that our lab mice innately respond to odors from three different species through sensory neurons of the VNO. Prey species could achieve a similar behavioral response to a variety of potential predators simply by detecting a single ligand common to all carnivorous animals (Fendt, 2006). In order to identify the molecular logic of how prey species respond with defensive behavior to a variety of other species, we aimed to isolate a second predator kairomone. Unfortunately, odor stimuli analyzed in our study are obtained by briefly swabbing medical gauze on a cat’s neck or isolating recently shed snake skin, and we found neither of these preparations to be readily amenable to fractionation. However, we did find cat saliva, a potential source of fur chemosignals, sufficient to induce c-Fos expression in the AOB and initiate defensive behavior (experimental logic in “stimuli” of Extended Experimental Procedures; Figures 6A and 6B). The submandibular salivary gland is known to secrete copious amounts of Feld4, the cat homolog of the rat and mouse Mups (Smith et al., 2004). Feld4 is a prominent cat allergen, indicating that it is stable and transmissible between species (Smith et al., 2004). Therefore, we considered this Mup protein as the potential source of the cat kairomone bioactivity.

Interestingly, a native odor sample that failed to elicit defensive behavior (rabbit urine; Figure 1E) does not appear to contain protein as the potential source of the cat kairomone bioactivity. To directly test whether cat Feld4 is detected as a kairomone, we expressed and purified a recombinant fusion protein between MBP and Feld4 in E. coli (rMup-Feld4). When assayed, rMup-Feld4 predominantly accounted for the native kairomone activity (Figure 6A). We used calcium imaging analysis to identify the responding sensory neurons. rMup-Feld4 failed to produce calcium transients in MOE neurons (Figure S6N); however, the recombinant protein was sufficient to directly initiate calcium transients in a subpopulation of VNs and generate a c-Fos response in the AOB (Figures 6B and 6C; Figures S6A–S6K). This activity is dependent on VN signaling, as TrpC2 \(^{-/-}\) VNs failed to produce calcium transients or c-Fos induction in the AOB to rMup-Feld4 (Figure 6C; Figures S6D–S6M). Importantly, whereas TrpC2 \(^{-/-}\) animals do not display significant defensive responses toward recombinant Feld4 (Figure 6D), this ligand is sufficient to promote significant defensive behaviors and ACTH release in wild-type mice (Figure 6A; Figure S6O). Our analysis indicates that detection of cat Feld4 through the VNO sensory system induces defensive behavior.
Intra- and Interspecies Mups Are Functionally Distinct

Given that mouse Mups have a different biological significance for mice than rat or cat Mups, we analyzed the neuronal and behavioral responses to Mups from different species. We compared the response of dissociated VN neurons to each of these cues by calcium imaging to determine whether they activate identical populations of neurons. We found four independent populations of responding cells, some that detect multiple Mup variants and some that were reproducibly and specifically activated by individual recombinant ligands (Figure 7A; Figure S7H). Among these, there were ensembles of neurons that displayed calcium transients solely to either mouse Mup variants (Figure 7A), seventh bar) or Mups from cat and rat (Figure 7A, fifth bar). These VN responses are likely driven by sensory receptor tuning to sequence variance of the individual Mups (Figures S7A and S7C). We expect these populations to be biologically relevant because of the significant number of activated cells compared to our negative control stimulus, (rMBP, Figure 7A), as well as the reproducibility of the specific responses when a single stimulus was repetitively pulsed (Figure S7B). This analysis reveals that each Mup stimulus has the potential to encode a different quality of information (Figure S7H). However, on its own, VN activation profiles do not reveal the underlying neural code that generates behavior.

Next, we took advantage of the fact that the defensive behavior in response to kairomones is context independent: the response occurs when stimuli are solely presented on a cotton gauze. In contrast, aggressive behavior promoted by mouse urine pheromones is context dependent: initiated only when coincidentally detected with another mouse. Behavior in response to mouse Mups out of context has not been evaluated. We analyzed freely moving behaving animals for avoidance time and risk assessment episodes in response to exposure to native mouse Mups presented on cotton gauze. As expected, removed from the context of another mouse, mouse Mup pheromones did not initiate aggressive behavior. Interestingly, they equally showed no signs of initiating defensive behavior (Figure 7B). To control for the possibility of mouse Mup habituation or contextual learning of self-expressed Mups, we additionally assayed the response to Mups from a different mouse strain (heterogenic Swiss), which excretes different Mup variants (Cheetham et al., 2009). Cues from Swiss mice activate subsets of VNs tuned to strain differences (Figure 7C; Figure S7E), and we found this pheromone stimulus to be equally unable to induce defensive behaviors or the release of ACTH (Figure 7B; Figure S7D). We assayed the ability of the pheromone- and kairomone-responding VNs to additionally detect our control complex natural stimulus, the HMW fraction of rabbit urine, and found it not to activate kairomone-responsive VNs (Figures S7F and S7G). This VN response is consistent with the lack of defensive behavior observed in response to rabbit urine (Figure 1E). Taken together,
Figure 7. Kairomones and Pheromones Encode Different Functions
(A) Left: representative calcium transients from isolated VNs, sequentially stimulated with recombinant maltose-binding protein (rMBP), rMup-Feld4, rMup-Rn13, and recombinant mouse Mup pheromones (a pool of mouse rMup3, 8, 17, 24, and 25). Axis bars: X = 60 s; Y = 3 x (F340/380 nm). Boxes indicate application and duration of stimulus. Right: comparative percent activity of dissociated VNs stimulated with recombinant rat and cat kairomones and mouse Mups as assayed by calcium imaging. Each bar denotes the percentage of all imaged cells exhibiting a calcium spike in response to the stimuli marked with a plus sign and not exhibiting a response to the stimuli marked with a minus sign. All cells were exposed to all four stimuli, except for control cells, which were exposed to the indicated number of repetitive pulses of rMBP (white bars). Note the presence of populations of cells responsive to kairomones only (fifth bar) and responsive to all Mups (first bar), which are significantly above controls exposed to pulses of rMBP. n = 10–11 experiments.
(B) Avoidance and risk assessment behaviors are triggered only in the presence of the rat-derived HMW fraction (blue bars, top panels) and cat swab (blue bars, bottom panels) interspecies Mups, but not in the presence of C57BL/6J mouse HMW fraction (white bars, top panels) or Swiss strain urine (white bars, bottom panels), which contain mouse Mups. n = 11–12.
(C) Venn diagram showing populations of cells responsive to kairomones (rMup-Rn13 and rMup-Feld4) and/or Mup-containing HMW fractions from C57BL/6J and Swiss mouse urine, as assayed by calcium imaging (n = 10–11 experiments; see also Figure S7E for complete documentation of % activated VNs). Statistical significance of each population (represented by each intersect), against respective rMBP control pulses, is color coded.
(D) Model for the proposed co-option of semiochemicals. Left: schematic representation of chronograph of Mup ligand evolution. Center: following stabilization of detection of ancestral ligand, genomic duplication and drift enabled Mups to be detected as kairomones (purple) or pheromones (red). Right: Mups have undergone neofunctionalization to instruct odor (ctl in B) is PBS-soaked gauze.
See also Figure S7.

our analyses indicate that there is a functional difference between cat and rat Mups that are detected as kairomones and mouse Mups that are detected as pheromones. This difference is likely initiated by Mup-specific activation of VN ensembles.

DISCUSSION

Accessory Olfaction Function Is Not Limited to Pheromones
Multiple olfactory subsystems are present in the mammalian nasal cavity, including the MOE, VNO, septal organ, and Grueneberg ganglion. The functional significance of this anatomical segregation has not been determined in mammals (Munger et al., 2009). Since its discovery almost two centuries ago, it has been speculated that the VNO serves to detect pheromones (Tirindelli et al., 2009). More recent studies using genetic tools, electrophysiological recordings, and calcium imaging assays have confirmed this function (Chamero et al., 2007; Holy et al., 2000; Leypold et al., 2002; Luo et al., 2003; Stowers et al., 2002). The estimated number of VNO sensory receptors (>250) vastly exceeds both currently identified pheromone ligands and the expected range of social behaviors mediated by pheromones, leaving potential coding space for other types of olfactory cues (Shi and Zhang, 2007; Young et al., 2005; Young and Trask, 2007). We now show that mouse VNs detect nonpheromonal ligands and that the accessory olfactory system is functionally necessary to initiate innate, stereotypic defensive behaviors and endocrine surges in response to odors from other species.
Chemical Detection of Threatening Environments

We were able to purify kairomone ligands from two different species and found them both to be Mup homologs. Mups are endowed with several characteristics that serve as good protein kairomones (Wyatt, 2003). First, the receiving animal must detect a ligand that is fixed in the genome of the signaler. Mup genes have been retained in the genomes of all sequenced placental mammals (except for humans), suggesting that they likely possess an advantageous ancestral function (Logan et al., 2008). The primary function of Mups in the signaling animal is not known; however, recent reports indicate that at least one Mup has beneficial metabolic effects by decreasing hyperglycemia and glucose intolerance in type 1 and type 2 diabetic mice (Zhou et al., 2009). Second, they must be easily detected by the receiver. In all known cases, Mups are secreted or excreted into the environment where they are extremely stable, resistant to degradation, and easily transmissible between individuals. This is demonstrated by the fact that many major respiratory allergens are either Mups or related lipocalins (Virtanen et al., 1999). For example, over 60% of humans who are allergic to cats test positive for Feld4-specific IgE (Smith et al., 2004). Third, genome analysis has shown that the Mup gene complement has undergone multiple species-specific evolutionary expansions followed by selective constraint (Logan et al., 2008; Mudge et al., 2008). The organization of olfactory receptor genes in genomic clusters susceptible to duplication (Lane et al., 2002) and the anatomical organization of the olfactory bulb into modular glomerular units (Mombaerts, 2001) provide a system suitable for expansion of olfactory detectors that can encode a novel function (neofunctionalization).

Co-option of Semiochemicals: One Mechanism, Multiple Functions

How does a detection system that responds defensively to a variety of species upon first exposure evolve? Isolation of Mup homologs of distinct behavioral consequences from two different species provides great insight. We have previously shown that one of the VNO’s functions is the detection of aggression-promoting Mup pheromones (Chamero et al., 2007). We have now determined that Mups also function to communicate between species via the VNO. Although we do not know the ancestral function of Mups, their detection may have become stabilized to sense one’s own production of Mups, to protectively detect other species, or to communicate within a species (Figure 7D, top panel). Once the ancestral ligand/receptor detection pairing was constrained in the genome, duplication followed by neofunctionalization in an evolutionary mouse lineage may have, for example, enabled sensation of additional species emitting Mups of more divergent sequence (Conant and Wolfe, 2008; McLennan, 2008), initiating an interspecies defense system that increased fitness and further stabilized Mup detection (Figure 7D, middle panel). Finally, the Mup gene cluster expanded and cognate olfactory receptors diversified to provide for intraspecies communication (Figure 7D, bottom panel). In this scenario, Mup detection is co-opted from kairomone to pheromone (Figure 7D).

Interestingly, defensive behavior can entail freezing, fighting, or fleeing, depending on the context of the stimuli (Eilam, 2005). Although pheromone-mediated aggression (fighting) and kairomone-promoted defensive fleeing are mutually exclusive behaviors, the controlling neural circuits may share common mechanisms. Our current findings initiate several fundamental questions. How does the receiving animal differentiate those Mups emitted from a conspecific, which do not elicit defensive behavior, from those from species that do? Is the activity from all kairomones integrated into a common neural circuit that serves as a master control of defensive behavior? The purification and identification of salient ligands with intrinsic activity now provide the molecular tools to detect and manipulate the precise neural code that governs behavior.

EXPERIMENTAL PROCEDURES

Mice

Wild-type animals were 8-week-old male C57BL/6J mice, unless otherwise noted. Female mice showed identical responses as analyzed by c-Fos expression and behavior (data not shown). TrpC2−/− and TrpC2+/− littermates were obtained from heterozygous mating couples, which were produced by backcrossing the TrpC2−/− knockout line (Stowers et al., 2002) into the C57BL/6J background for at least four generations. To ensure the identification of innate behavior, animals had no previous exposure to odors from other animal species. All procedures were approved by the Institutional Animal Care and Use Committee.

Behavioral Assays

Individually caged mice were habituated for 2 hr in the dark over 2 consecutive days and assayed on day 3. See Extended Experimental Procedures for collection and preparation of predator odor and control stimuli. Mice were assayed and filmed for 1 hr in the dark. Movies were scored blindly for approach and avoidance times during the first 30 min of exposure; risk assessment episodes were quantified for the first 15 min of assay (see details in Extended Experimental Procedures and Movie S1). Either unpaired t tests or one-way ANOVA were applied. The number of risk assessment episodes was additionally scored for ten subsequent 3 min sessions and statistically compared to controls by one-way ANOVA. Error bars indicate SEM.

Calcium Imaging

Transient increases in free Ca2+ concentration in dissociated VNO neurons were determined by ratiometric Fura-2 fluorescence as described (Chamero et al., 2007). The HMV and FPLC-A fractions of rat urine and recombinant rat and cat Mups were imaged at 3.33 µg/ml (Figure 4C) or 10 µg/ml (elsewhere) in imaging buffer unless otherwise specified (see Extended Experimental Procedures for details on rat urine fractionation and production of recombinant Mups). Control fractions were diluted to the same extent as the FPLC-A fraction, irrespective of actual protein concentration in the fraction. Pooled mouse Mups were imaged at a total of 27.7 µg/ml as described (Chamero et al., 2007). Protein concentrations were calculated by Bradford assay and adjusted for recombinant maltose-binding protein (rMBP) content. The rMBP control was imaged at 6.66 µg/ml (Figure 4C) or 20 µg/ml (elsewhere) in imaging buffer. Statistical significance was tested using one-way ANOVA followed by the Tukey-Kramer HSD post hoc analysis. Error bars indicate mean ± SEM. Further details on stimuli used, number of experiments, imaged cells per experiment, and percentages of activated cells are given in Extended Experimental Procedures and Table S1.

Additional methods can be found in the Extended Experimental Procedures.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Extended Experimental Procedures, seven figures, one table, and two movies and can be found with this article online at doi:10.1016/j.cell.2010.03.037.
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REFERENCES


Supplemental Information

EXTENDED EXPERIMENTAL PROCEDURES

Stimuli
Cat stimulus was obtained by rubbing medical gauze against the fur of a cat. One milliliter of cat saliva was used as cat stimulus; four 5 × 5 cm pieces of skin shed by a pet snake were used as the snake stimulus; rat stimulus was obtained by placing 1000 μl of rat urine on medical gauze; Swiss and C57BL/6J mouse stimuli were 200 μl urine, which had an equivalent amount of Mup-sized proteins, as judged by Coomassie staining of PAGE gels and Bradford analysis (not shown); rabbit stimulus was 1000 μl urine on medical gauze. HMW and LMW fractions as well as the FPLC-A peak, obtained during fractionation of rat urine (see below), were used at equivalent concentrations of rat urine protein used in the same exposure experiment. Recombinant proteins were used at equivalent concentrations to HMW fraction protein concentration. Control mice were exposed to clean medical gauze attached to the binder clip. Where recombinant proteins (see below) or other liquid preparations were used as stimuli, the control gauzes were wetted with the equivalent volume of 1× PBS. It is known that the kairomones found in cat fur have a stronger effect on rodents when collected from the cat’s neck region, and it has been suggested that this region is constantly licked by the animal, perhaps transmitting kairomones from the saliva to the fur (Anderson and Baer, 1981). Fortuitously, one of our more playful volunteer cats chewed the gauze as we swabbed her fur and we found it to elicit more substantial c-Fos expression throughout the accessory olfactory system. This led us to directly test cat saliva for kairomone activity. Responses to 5 μl, 50 μl, or 80 μl of 15.7 M trimethylthiazoline (TMT; as published in Kobayakawa et al., 2007) were similar; 5 μl were used as stimulus in Figure S1. We chose naphthalene (scent related to the smell of fire, naturally released from burning wood) as a generally aversive odorant because it is relatively stable and nonreactive and therefore additional types of sensory information (pain, tingling, or nociceptive thermosensation) are unlikely to be conveyed upon acute exposure, thus providing a good olfactory-only control. For Figure 1F, naphthalene (0.5 g pellet) was presented completely enclosed in a gauze mesh to prevent oral contact.

Fractionation of Rat Urine, Small-Molecule Displacement, and MS Analyses
Five hundred microliters of freshly collected rat urine were placed on a 10 kDa cutoff YM10 ultrafiltration microcolumn (Millipore) and spun at 10000 g for 35 min at 4°C. The filtrate was saved and the retentate was washed four times with the same volume of ice-cold PBS in the same YM10 column and then resuspended in 500 μl of PBS to produce the HMW fraction. The filtrate saved in the first centrifugation step was applied over a new YM10 filtration device and the flowthrough was named the LMW fraction. The HMW fraction was desalted and applied to a 5 ml Hi-Trap Q High Performance anion-exchange FPLC column (GE Life Sciences). Fractions were eluted over a 0–500 mM NaCl gradient and then desalted prior to animal presentation. For the displacement of small organic molecules bound to macromolecular components in the HMW fraction, 500 μl of this fraction were incubated in an uncovered Eppendorf tube with 8 μl of a 4 mg/ml solution of menadione in ethanol for 30 min at room temperature with gentle shaking. The solution was then spun over a YM10 column and the retentate containing the high molecular weight components devoid of their small organic molecules was washed 3 times with PBS containing 0.4 mg/ml of menadione, twice with PBS and resuspended in PBS to produce the HMW fraction treated with menadione (Figure SSA). HMW, LMW and the menadione-treated HMW fraction were extracted with an equal volume of dichloromethane and the lower phase was subjected to GC-MS analysis, as published previously. Additionally, the HMW fraction was subjected to ESI-MS analysis and MS proteomics analysis to determine the masses and peptidic composition of the sample, as described before (Chamero et al., 2007). The same ESI-MS conditions were also used to analyze rabbit urine.

Calcium Imaging
For Figure 2A, approximately 1.5 g snake skin was ground in liquid nitrogen to a fine powder, suspended in 20 ml calcium imaging buffer, incubated with rocking at 4°C for 2 hr, and centrifuged twice to remove debris. Cat odorized gauze was similarly incubated in calcium imaging buffer. The HMW and FPLC-A fractions of rat urine and recombinant rat and cat Mups were imaged at 1:300 dilution, 178 nM rMup-Rn13, 171 nM rMup-Feld4 (Figure 4C) or 1:100 dilution, 534 nM rMup-Rn13, 513 nM rMup-Feld4 (elsewhere) unless otherwise specified. C57BL/6J and Swiss urine HMW fractions were diluted 1:300, as described (Chamero et al., 2007). Rabbit urine HMW fractions were diluted 1:100, irrespective of actual protein concentration in the fraction. The rMBP control was imaged at 159 nM (Figure 4C) or 476 nM (elsewhere). Responses to 100 mM KCl and 10 mM vanillin (in MOE sensory neurons) were imaged as positive controls. To determine whether the number of neurons showing transient increases in response to specific stimuli were significantly different from the number showing spontaneous increases, we measured the spontaneous calcium influx in 4315 neurons adjusted for the length of the stimulus window. As rMBP showed no significant VN activation (see Figure 4C), in all subsequent experiments we used sequential rMBP stimuli as a negative control.

Behavioral Assays
All subjects were sequentially exposed to odor, monitored for behavior, and subsequently processed for immunostaining, ensuring that the cellular response and behavior were analyzed from the same individuals and no animals were reused for behavioral analysis. All behavior was filmed in the dark with infrared video for subsequent analysis.

A number of behavioral and endocrine responses have been observed in rodents toward predators and predator odors (Blanchard et al., 2001; Dielenberg and McGregor, 2001) including increased physical avoidance, vocalization and freezing duration, decreased grooming, increased number of defensive burying and risk assessment episodes (Hebb et al., 2004), as well as elevated plasma
ACTH and corticosterone levels. In laboratory mice, vocalizations are not observed toward predator odors (Blanchard et al., 2001), while freezing is seen just toward repugnant predator odors, such as TMT, and not to other predator odors (Blanchard et al., 2001; Hebb et al., 2004); moreover, increased defensive burying and decreased grooming are also observed toward generally aversive odorants, such as butyric acid (Hebb et al., 2004). In contrast, the combination of prolonged avoidance, risk assessment behaviors and ACTH release is specifically displayed toward many different predators and has been used as a set of outputs to model innate fear responses in rodents (Blanchard et al., 2001; Kobayakawa et al., 2007). These were therefore chosen as the three behavioral/endorcine readouts in our assays. On the test day, mice were exposed for 1 h to the stimulus, which was positioned in one side of the cage attached to a binder clip, to visually confirm its position. Movies were later scored blindly for approach and avoidance times and the number of risk assessment episodes directed toward the stimuli. Approach behavior was defined as the amount of time animals spent within a 15x5 cm rectangular area surrounding the stimulus (area 1 in Figure 1); avoidance behavior is the amount of time the animals spent inside the far 15 x 5 cm region (area 2). Risk assessment behaviors have been consistently used as indicators of defensive behavior in many animals (Blanchard et al., 2001); the kind of risk assessment behavior used in the present study is the flat-back/stretch-attend response typically exhibited by mice and rats exposed to predators (Kemble and Bolwahnn, 1997). Initially, the subject slowly approaches the threat source, then lowers its head, flattens its back while nearing the stimulus, resulting in an extended posture with a raised tail and thursted head (described in detail in Blanchard et al., 1990; see also Movie S1).

For exposure to the live, intact rat (Figure S1), the arena was a square 70 x 70 cm open-field box. One side of the arena contained a covered black hiding box (transparent to infrared wavelengths enabling filming of mouse activity) with a 6 x 6 cm arch to allow entrance. Mice were habituated to the arena as described. On day 3, a rat was anesthetized with an IP injection of a mixture of 7.5 mg/kg xylazine and 7.5 mg/kg ketamine, and placed in the arena opposite the hiding box, with its paws and snout facing the wall (see drawing in Figure S1D); approach behavior was defined as the amount of time animals spent within a 20 x 70 cm rectangular area surrounding the rat, while avoidance behavior is the amount of time the animals spent inside the 20 x 70 cm hiding box. Assay duration: 28 min.

When tested in a 50 x 50 cm open-field apparatus (Med Associates), wild-type and TrpC2−/− mice exhibited statistically equivalent locomotor activity (2240.2 ± 305.8 cm ambulatory distance traveled by wild-type subjects versus 2257.1 ± 330.5 cm by TrpC2−/− mice in a 10 min session; n = 8; mean ± SEM; p = 0.49, one-tailed t test). Moreover, in elevated plus maze tests (Harvard Apparatus; Walf and Frye, 2007), wild-type and TrpC2−/− mice exhibited equivalent levels of general anxiety (36.8 ± 8.3 s spent in the open arms by wild-type subjects compared to 51.96 ± 6.1 s spent by TrpC2−/− mice in a 5 min session; statistically equivalent number of head-out episodes; n = 8; mean ± SEM; p = 0.08, one-tailed t test).

**Fear Conditioning**

Male mice were habituated for 360 s on day 1 to the footshock conditioning apparatus (Harvard Apparatus) in the presence of 20 μl of a 100 mM eugenol solution (control odor) without footshock (FS), in physical context 1. On day 2, mice were footshocked (0.5 mA at 120 s, 180 s, 240 s and 300 s; session ended at 360 s) in the presence of 20 μl of a 100 mM pentyl acetate solution (FS-paired odor), in the same apparatus but with physical context 2. On the test day, freezing behavior was measured during the first 3 min after introduction into the apparatus prior to odor exposure (“pre-odor” session in Figure S2I). During the following 3 min, control or FS-paired odor was presented in physical context 3 and freezing behavior was measured (“odor” session in Figure S2I). Similar results were obtained with pentyl acetate as the control odor and eugenol as the FS-paired odor.

**ACTH Assay**

Measurements of ACTH levels in blood of mice exposed to odors were performed as published (Kobayakawa et al., 2007). Mice were individually housed on day 1, habituated to the assay room on days 3, 4, and 5, and exposed for 10 min to odor stimuli or 30 min restraint on day 6 (within the 10 am to 11 am period). Five microliters of a 100 mM eugenol solution was used as an irrelevant odor and trunk blood was collected. Plasma ACTH concentrations were measured using the ACTH ELISA kit (MD Biosciences). n = 5-6.

**c-Fos Immunostaining**

Following odor exposure, brains were fixed overnight in 4% paraformaldehyde, equilibrated in 30% sucrose/1× PBS and sectioned on a Leica 1400S vibrating-blade microtome. Forty μm coronal sections were collected for the entire brain and for the VNO. Suitable sections were chosen for subsequent c-Fos immunostaining based on comparisons to a reference brain atlas (Paxinos and Franklin, 2003). Sections were blocked as free-floating sections for 1 hr with 1% blocking reagent (Molecular Probes), followed by incubation with the anti-c-Fos primary antibody (rabbit polyclonal; Ab2 or Ab5; Abcam) diluted 1:5000 in 1% BSA/1× PBS/0.3% Triton X-100 for 36 hr at 4° C under gentle agitation. Sections were washed twice in 1× PBS/0.1% Triton X-100, 15 min each, and incubated for 3 hr at room temperature with Alexa 488-conjugated goat anti-rabbit secondary antibody (Molecular Probes) diluted 1:500 in 1% BSA/1× PBS/0.3% Triton X-100. After two washes in 1× PBS/0.1% Triton X-100, 15 min each, sections were counterstained with To-Pro-3 nuclear stain (Molecular Probes) diluted 1:1000 in 1× PBS, washed twice in 1× PBS, 15 min each, and mounted onto glass microscope slides with Vectashield HardSet (Vector Labs). Dry mounted sections were imaged on an Olympus Fluoview 500 confocal fluorescence microscope. The number of c-Fos-positive nuclei was counted blindly for each individual. For aAOB sections, the number of c-Fos-positive nuclei was counted in two consecutive sections for the aAOB (approximate bregma 3.50 mm) and two consecutive
sections for the pAOB (approximate bregma 3.20 mm) for equivalent areas (approximately 19,000 μm²) in the granule and mitral cell layers. Numbers of c-Fos nuclei were usually highly concordant between the two sections analyzed, and data from only one section from each animal was computed to calculate means and errors. In the rare instances when they were discordant, the cell count from a third consecutive imaged section was used to calculate an average number of c-Fos nuclei for that animal. The differences in mean numbers of c-Fos-positive cells were tested by one-way ANOVA followed by Tukey’s HSD post hoc analysis against the respective controls.

Recombinant Protein Expression

The cDNA for the rat major urinary Mup13 protein (GenBank accession number P02761) was amplified by PCR from male Sprague-Dawley rat liver cDNA using oligonucleotides 5’-ATCGGATCCCATGCAGAAGAAGCTAGTTCCACAAGAG-3’ and 5’-ATCAAGCTTTGTCCTGGCCTGGAGACAG-3’. This amplicon was cloned into pMAL-c2x bacterial expression vector (New England Biolabs) into BamHI and EcoRI restriction sites. Mup13 was expressed as a fusion protein with maltose-binding protein (MBP), following the manufacturer’s recommendations. Protein was eluted from an amylose affinity resin using maltose and then exchanged into 1× PBS using a YM10 column prior to calcium imaging and behavioral assays. Recombinant MBP (rMBP) was used as a control. The same procedure was applied for production of the recombinant cat Mup, except that the corresponding cDNA was synthesized in vitro based on the published sequence of Feld4 (cat Mup; GenBank accession number NM_001009233; Smith et al., 2004). We amplified a second Feld4 allele from cat genomic DNA, that results in an R/K change in the polypeptide at position 43 (numbering in Figure S7A). All studies presented here were conducted with the cat sequence shown in Figure S7, but similar results were obtained with the R43K variant (not shown).

SUPPLEMENTAL REFERENCES


Figure S1. TrpC2 Function Is Necessary for Defensive Behavior in Response to Snake and Rat Odors, Related to Figure 1

(A and B) The left graphs reveal TrpC2 involvement in the display of innate risk assessment behavior in response to rat urine odor (A) and snake shed skin (B). Right panels: histograms of avoidance behavior. TrpC2<sup>-/-</sup> mice exhibit impaired risk assessment behavior, but normal avoidance, toward snake odor. This overlying behavior likely occurs through MOE detection, since snake skin is characteristically pungent as detected by humans.

(C) Plasma ACTH concentration in animals exposed to rabbit urine (rab) is not significantly different from levels in subjects exposed to control odor (ctrl). Physical restraint (restr) is used as a positive control.

(D) See also Movie S2. Wild-type, but not mutant TrpC2<sup>-/-</sup>, mice display defensive behaviors to live rats. Freely moving behaving mice were quantified for approach, avoidance and risk assessment behavior in the presence of an anesthetized rat. Both WT and TrpC2<sup>-/-</sup> animals displayed similar responses to a novel furry control object in the arena (dummy rat).

(E) Avoidance and risk assessment behaviors in animals exposed to control odor and TMT. Blue bars indicate behavior exhibited by another group of animals exposed to cat odor, as a reference. Mice show avoidance but do not perform risk assessment behavior in response to TMT. TrpC2<sup>-/-</sup> animals continue to detect and avoid TMT. Further, TMT does not induce c-Fos expression in the AOB (FP, data not shown). n = 4–8; *p < 0.05; **p < 0.01; ***p < 0.001; one-tailed unpaired t test (avoidance and risk assessment bar graphs and experiment in C) and ANOVA (risk assessment line graphs). Mean ± SEM. Control odor (ctrl) was PBS-soaked gauze (A, C, and E), or clean gauze (B).
TrpC2 +/+ (aAOB) TrpC2 −/− (aAOB)

control odor

cat odor

rat odor

snake odor

TrpC2 +/+ (aAOB) TrpC2 −/− (aAOB)

control odor

cat odor

rat odor

snake odor

Figure S2. AOB Activation in Mice Exposed to Kairomone and Control Odors, Related to Figure 2

(A–H) Low magnification confocal images of coronal anterior AOB, aAOB, sections immunostained for c-Fos in TrpC2+/+ (A, C, E, and G) and TrpC2−/− (B, D, F, and H) mice exposed to control (A and B), cat (C and D), rat (E and F), or snake (G and H) odors. Control odor is clean gauze. c-Fos-positive cells are concentrated in the granule cell layer (gr) and occasionally in the mitral cell layer (mcl) in (C), (E), and (G).

(I–N) Fear conditioning by pairing a regular odorant (pentyl acetate) with footshocks (I; see Extended Experimental Procedures for details) does not lead to c-Fos induction in the AOB (J–M; quantification in N). Control odor is eugenol, which was not paired with footshocks. Preodor (white) and odor (black) bars indicate amount of freezing before and after the onset of odor presentation, respectively, d, dorsal; m, medial. The scale bar represents 100 μm. **p < 0.01; ***p < 0.001; one-tailed unpaired t test (I) or ANOVA followed by Tukey-Kramer HSD post hoc analysis (N). Mean ± SEM.
Figure S3. AOB Activation in Mice Exposed to Complex Natural Odors, Related to Figure 3
(A–F) c-Fos immunostaining images of coronal sections through the anterior (aAOB) and posterior (pAOB) accessory olfactory bulbs, showing activity induced by control complex stimuli (G and E).

(G) Quantification of c-Fos-positive nuclei in the granule and mitral cell layers of the AOB in animals exposed to control native odors. White bars, aAOB; black bars, pAOB. gl, glomerular layer; mcl, mitral cell layer; gr, granule cell layer; d, dorsal; m, medial. The scale bar represents 100 μm. *p < 0.05; **p < 0.01; ***p < 0.001; ANOVA followed by Tukey-Kramer HSD post hoc analysis. Mean ± SEM. Control odor was PBS-soaked gauze.
Figure S4. Calcium Imaging Controls to Stimulation with Recombinant Mup Proteins, Related to Figure 4

(A) Response to KCl is indistinguishable in dissociated VNs from wild-type (WT) and TrpC2-/- animals. n = 6–11 experiments.

(B) Controls to accompany Figure 4E. Comparative percent activity of dissociated VNs stimulated with rMBP, rMup-Rn13, the FPLC-A peak, and the HMW fraction as assayed by calcium imaging. The few activated cells in each population (black bars) are not significantly different from the number of cells responding to pulses of rMBP (white bars). All cells analyzed were sequentially exposed to all four stimuli shown below the graph, except for the control rMBP pulses. A plus sign indicates that the corresponding stimulus induces calcium transients in the population of neurons represented by the bar above; a minus sign indicates that the population is not responsive to the corresponding stimulus. Each stimulus was pulsed for 60 s, with 120 s recovery, for a total duration of 13 min. n = 10-11 experiments.

(C) Dose-response curve of VNs stimulated with rMup-Rn13 (green circles) and a cat Mup ortholog (rMup-Feld4, red triangles), at concentrations between 0.33 mg/ml and 0.03 mg/ml n = 4–6 experiments/point. Curves fitted to Boltzmann equation: y = A2 + (A1 - A2)/(1 + exp((x - x0)/dx)). n.s., nonsignificant. ANOVA followed by Tukey-Kramer HSD post hoc analysis. Mean ± SEM.
Figure S5. ACTH and c-Fos Induction in the Presence of Recombinant Rat Mup, Related to Figure 5

(A) Menadione displacement of potential small molecules bound to protein component of rat urine HMW fraction does not alter the behavior-inducing activity of this fraction. n = 8–12. *p < 0.05; **p < 0.01; ANOVA followed by Tukey-Kramer HSD post hoc analysis. Mean ± SEM.

(B) Exposure to recombinant Mup13 (rMup-Rn13), but not to recombinant maltose-binding protein (rMBP) alone, leads to ACTH release. Restrained animals (restr) are used as positive control. **p < 0.01; ***p < 0.001; ANOVA followed by Tukey-Kramer HSD post hoc analysis. n.s., nonsignificant. Mean ± SEM.

(C) Quantification of c-Fos-positive nuclei in the granule and mitral cell layers of the AOB in animals exposed to recombinant rMup-Rn13 protein (see corresponding images in Figure 5C). White bars, aAOB; black bars, pAOB. *p < 0.05; **p < 0.01; ***p < 0.001; ANOVA followed by Tukey-Kramer HSD post hoc analysis. Mean ± SEM c-Fos expression is statistically indistinguishable in animals exposed to rat urine and rMup-Rn13 (HSD contrasts between the rat urine and rMup-Rn13 groups yielded p = 0.266 for granule cell layer in aAOB, p = 0.397 for granule cell layer in pAOB, p = 0.286 for mitral cell layer in aAOB and p = 0.103 for mitral cell layer in pAOB). Control odor was PBS-soaked gauze (A and C).
Figure S6. Recombinant Cat Mup Displays Kairomone Activity, Related to Figure 6

(A) Quantification of c-Fos-positive nuclei in the granule and mitral cell layers of the AOB in animals exposed to recombinant rMup-Feld4 protein (see corresponding images in Figure 6B). White bars, aAOB; black bars, pAOB. *p < 0.05; **p < 0.01; ANOVA followed by Tukey-Kramer HSD post hoc analysis. Mean ± SEM c-Fos expression is statistically indistinguishable in animals exposed to cat saliva and rMup-Feld4 (HSD contrasts between the saliva and rMup-Feld4 groups yielded p = 0.522 for granule cell layer in aAOB, p = 0.224 for granule cell layer in pAOB, p = 0.075 for mitral cell layer in aAOB and p = 0.072 for mitral cell layer in pAOB).

(B–M) TrpC2 function is necessary for c-Fos induction in the AOB following exposure to rat (F and G) and cat (J and K) recombinant Mup proteins. TrpC2−/− animals did not show c-Fos staining above control levels (D, E, H, I, L, and M).

(N) Quantification of response to rat urine, rat HMW fraction, recombinant rat Mup13 and cat Feld4 (rMup-Rn13 and rMup-Feld4) and 10 mM vanillin in dissociated sensory neurons isolated from the main olfactory epithelium of wild-type male mice and assayed by calcium imaging. The ordinate shows the percentage of activated cells. n = 7–20 experiments; ***p < 0.001; ANOVA followed by Tukey-Kramer HSD post hoc analysis against no stimulus control (white bar).

(O) Exposure to recombinant Feld4 (rMup-Feld4), but not to recombinant maltose-binding protein (rMBP) alone, leads to ACTH release. Restrained animals (restr) are used as positive control. *p < 0.05; **p < 0.001; ANOVA followed by Tukey-Kramer HSD post hoc analysis. n.s., nonsignificant. Mean ± SEM.

(P) ESI-MS analysis showing the absence of major proteins in rabbit urine.

(Q) Silver-stained gel of equivalent amounts of rabbit, C57BL/6J and Swiss mouse urine samples. C57BL/6J is additionally shown at a 2× total concentration to visualize less concentrated high molecular weight components. Control odor was PBS-soaked gauze (A–M).
Figure S7. Kairomones and Pheromones Activate Both Overlapping and Exclusive Subsets of VNs, Related to Figure 7

(A) Alignment of amino acid sequences of mouse Mups 3, 8, 17, 24, and 25, found in C57BL/6J urine, rat Mup13 and cat Mup (Feld4). Identical and similar residues are highlighted in black and gray, respectively.

(B) Calcium transients from isolated VNs, stimulated twice with either recombinant rat Mup13 (rat), a pool of five recombinant mouse Mups (mouse) or recombinant cat Mup Feld4 (cat). The cat and rat Mups were applied in separate experiments to show the fidelity of reproducibility for each stimulus and confirm that they each activate a subset of VNs that is not also activated by mouse Mups. When they are perfused in the same experiment the majority of VNs that detect rat Mup also detect cat Mup (Figure 7A) (H). Axis bars: X = 60 s; Y = 33 (F340/380 nm). Boxes indicate application and duration of stimulus; above, representative calcium influx from responsive neuron; below, images of corresponding VNs (arrows) during each stimulus application window. Cells are pseudocolored dark-to-light to indicate calcium influx.

(C) Rooted phylogenetic tree of major urinary proteins at the amino acid sequence level. Numbers above each node indicate bootstrap values.

(D) Plasma ACTH concentration in animals exposed to Swiss mouse urine (Swiss) is not significantly different from levels in PBS-exposed subjects (ctrl). Physical restraint (restr) is used as a positive control. ***p < 0.001; n.s, nonsignificant; ANOVA followed by Tukey-Kramer post hoc analysis.

(E) Comparative percent activity of dissociated VNs stimulated with recombinant kairomones, rMup-Rn13 and rMup-Feld4, and urine HMW samples from Swiss or C57BL/6J mouse strains, as assayed by calcium imaging (see corresponding Venn diagram in Figure 7C). All cells analyzed were sequentially exposed to all four stimuli shown below the graph, except for the control rMBP pulses (white bars). Each bar denotes a population of cells responsive to the stimuli indicated below the graph. A plus sign indicates that the corresponding stimulus induces calcium transients in the population of neurons represented by the bar above; a minus sign indicates that the population is not responsive to the corresponding stimulus. Each stimulus was pulsed for 60 s, with 120 s recovery, for a total duration of 13 min. n = 10-11 experiments.

(F) Comparative percent activity of dissociated VNs sequentially stimulated with recombinant kairomones, rMup-Rn13 and rMup-Feld4, C57BL/6J mouse and rabbit HMW samples, as assayed by calcium imaging.

(G) Corresponding Venn diagram, with the number of cells in each population indicated. Each stimulus was pulsed for 60 s, with 120 s recovery, for a total duration of 13 min. n = 10–12 experiments.

(H) Venn diagram showing populations of cells responsive to kairomones (rMup-Rn13 and rMup-Feld4) and recombinant mouse Mup pheromones (a pool of mouse rMup3, 8, 17, 24, and 25), as assayed by calcium imaging (n = 10-11 experiments; see also Figure 7A for documentation of % activated VN cells). Statistical significance of each population in the Venn diagrams (represented by each intersect), against respective rMBP control pulses, is color coded. *p < 0.05; **p < 0.01; ***p < 0.001; ANOVA followed by Tukey-Kramer post hoc analysis. n.s, nonsignificant. Mean ± SEM.