

Hypothalamic representation of the imminence of predator threat detected by the vomeronasal organ in mice

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Abstract

Animals have the innate ability to select optimal defensive behavioral outputs with an appropriate intensity in response to predator threat in specific contexts. Such innate behavioral decisions are thought to be computed in the medial hypothalamic nuclei that contain neural populations directly controlling defensive behavioral outputs. The vomeronasal organ (VNO) is one of the major sensory input channels through which predator cues are detected with ascending inputs to the medial hypothalamic nuclei, especially to the ventromedial hypothalamus (VMH). Here, we show that cat saliva contains predator cues that signal imminence of predator threat and regulate the robustness of freezing behavior through the VNO in mice. Cat saliva activates neurons expressing the V2R-A4 subfamily of sensory receptors, suggesting the existence of specific receptor groups responsible for freezing behavior induced by the predator cues. The number of VNO neurons activated in response to saliva correlates with the freshness of saliva and the intensity of freezing behavior, while the downstream neurons in the accessory olfactory bulb (AOB) and VMH are quantitatively equally activated by fresh and old saliva. Strikingly, however, only the number of VMH neurons activated by fresh saliva positively correlates with intensity of freezing behavior. Detailed analysis of the spatial distribution of fresh and old saliva-responding neurons revealed a neuronal population within the VMH that is more sensitive to fresh saliva than old saliva. Taken together, this study demonstrates that predator cues in cat saliva change over time and differentially activate the sensory-to-hypothalamus pathway. More specifically, the imminent predator signal predominantly activates V2R-A4 receptors, which results in the activation of freezing-correlated neurons in the VMH. In contrast, the less imminent predator signal activates V2R-A4 receptors to a lesser extent, which in turn results in the activation of distinct populations of neurons in the VMH that are not correlated to freezing.

eLife assessment

This **valuable** study shows that, in mice, fresh cat saliva elicits a greater defensive response compared to old cat saliva. Additionally, the authors implicate the vomeronasal organ (VNO) and ventromedial hypothalamus (VMH) as part of a circuit that underlies this process. While the study has potential, the results are somewhat preliminary, and as such the evidence presented is **incomplete**.

Introduction

Prey animals make defensive behavioral decisions through prompt and careful analysis of their environment when encountering predators. Immediacy of the threat, determined by perceived distance from predators, is one of the key factors influencing defensive behavioral decisions, which regulates the intensity of behavioral outputs (**Figure 1** [↗](#)) (Blanchard and Blanchard, 1989 [↗](#)). Predator-derived sensory signals, or predator cues, are detected through multiple sensory systems, including the visual system, auditory system, and olfactory system. These individual sensory cues are sufficient to induce innate defensive behavior such as fear-related freezing and flight, as well as anxiety-related risk assessment behavior. In nocturnal animals including rodents, olfaction is one of the major sensory modalities by which predator-derived chemical cues are detected (Apfelbach et al., 2005 [↗](#)). Rodents respond to a variety of predator-derived chemical cues, such as odors from fox feces, snake skin, cat fur, and cat collars, in both the field and laboratory environments (Apfelbach et al., 2005 [↗](#); Kats and Dill, 1998 [↗](#); Papes et al., 2010 [↗](#); Pérez-Gómez et al., 2015 [↗](#); Takahashi, 2014 [↗](#)). However, how animals use chemosensory cues to make appropriate behavioral decisions is not well understood.

The accessory olfactory system is one of the major sensory input channels through which predator-derived chemical cues are detected. It has been shown that some molecules derived from predator species activate sensory neurons in the VNO and induce defensive behavior (Isogai et al., 2011 [↗](#); Papes et al., 2010 [↗](#); Pérez-Gómez et al., 2015 [↗](#); Samuelsen and Meredith, 2009 [↗](#); Tsunoda et al., 2018 [↗](#)). Moreover, a number of studies have revealed main components of the sensory neural circuits that mediate behavioral responses to chemical predator cues in mice (Blanchard et al., 2003 [↗](#); Canteras, 2002 [↗](#); Choi et al., 2005 [↗](#); Dielenberg and McGregor, 2001 [↗](#); Markham et al., 2004 [↗](#); Petrovich et al., 2001 [↗](#)). Predator signals detected by the VNO are transferred to the accessory olfactory bulb (AOB). The output neurons in the AOB (the mitral cells) then send the signals to the extended amygdala consisting of the bed nucleus of the stria terminalis (BNST) and the posteroventral part of medial amygdaloid nucleus (MeApv), which connects with the dorsomedial part of the ventromedial hypothalamus (VMHdm) that contains neural populations directly controlling defensive behavioral outputs (Gross and Canteras, 2012 [↗](#); LeDoux, 2012 [↗](#); Takahashi, 2014 [↗](#)). It is, therefore, reasonable to hypothesize that the mouse VNO detects predator cues and sends sensory signals to the VMH, triggering appropriate defensive behavioral decisions. However, the direct evidence to support this hypothesis is still missing.

In this study, we investigated sensory-to-hypothalamic neural circuitries that regulate defensive behavior induced by non-volatile chemical cues in cat saliva with different freshness. We examined the role of the VNO in behavioral responses and sought sensory receptor genes for the predator cues. Moreover, by comparing downstream neural populations activated by cat saliva with different freshness, we investigated how the imminence of predator threat is processed in the accessory olfactory circuitry to regulate behavioral outputs.

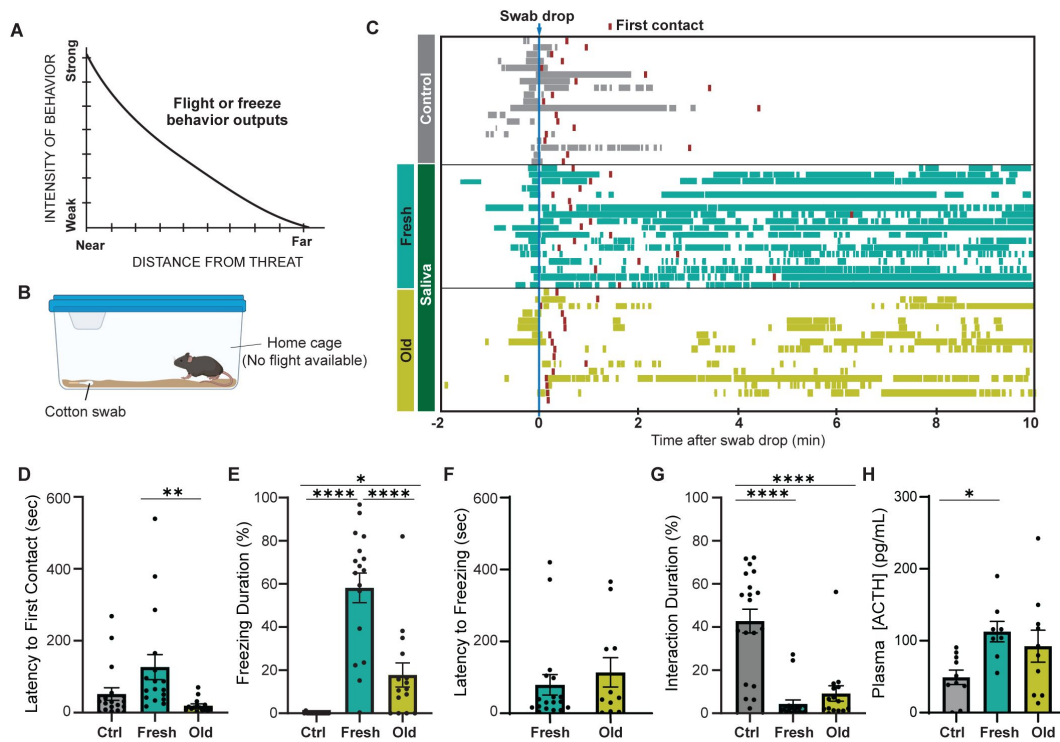


Figure 1

Predator cues in cat saliva induce context-dependent defensive behavior in mice

(A) A schematic diagram illustrating the elicitation of "freeze" or "flight" defensive behaviors in relation to distance. (Adapted with permission from (Blanchard and Blanchard, 1989))

(B) A schematic illustration depicting a test chamber for evaluating defensive behaviors.

(C) Raster plots displaying freezing episodes of individual mice exposed to a clean control swab (gray), fresh saliva (green), or old saliva (yellow). The introduction of a clean, fresh, or old saliva swab into a mouse's cage is denoted as time 0. A red line indicates the first interaction with a swab for each mouse.

(D) Latency to the first episode of interaction with a swab.

(E) The percentage of total freezing episodes after the first contact.

(F) Latency to the first freezing episode in mice exhibited freezing after the first interaction.

(G) The percentage of total interaction episodes.

(H) Plasma ACTH concentrations following exposure to clean, fresh, or old saliva swabs.

In (D) - (H), the values are presented as means $\pm$ S.E.M., with individual dots representing individual mice. Statistical analysis was performed using One-Way ANOVA with Tukey's multiple comparisons. *, **, and **** denote significance levels of $p < 0.05$, 0.01 , and 0.0001 , respectively.

Results

Predator cues in cat saliva induce context-dependent defensive behavior in mice

To explore whether mice exhibit different intensities of defensive behavior in response to chemosensory predator cues with varying threat immediacy, we examined their response to cat saliva with different degrees of freshness, which potentially represents immediacy of the predator threat. Cat saliva has been considered as a source of predator cues found on cat fur and collars (Papes et al., 2010). Fresh saliva is assumed to indicate a more immediate threat to the prey animal, which likely triggers stronger defensive behavior. We exposed mice to a cotton swab containing cat saliva in their home cage where they know a flight route is not available (Figure 1B). We found that mice approached and sniffed the cotton swab, regardless of the presence of saliva, as part of their exploratory behavior towards a novel object in their home cage (Supplemental movies 1-3, Figure 1C; red boxes indicate the first contact). The latency to the first contact was slightly longer for saliva collected immediately before introduction to the mouse cage (fresh saliva) when compared to that for saliva collected a few hours prior to the behavior test (old saliva) (Figure 1D). The difference in latencies was statistically significant, indicating that mice can perceive the qualitative change of cat saliva over time (Figure 1D). Baseline freezing episodes were observed in all groups of mice when the sample was introduced into the cage due to the experimenter's intervention (Figure 1C). However, the groups exposed to cat saliva continued to exhibit freezing behavior throughout the 10-minute observation period, while mice exposed to a clean swab did not exhibit freezing behavior after the first contact (Figure 1C, E). Moreover, we found a correlation between the freshness of cat saliva and duration of freezing behavior in mice. Mice exposed to fresh saliva exhibited considerably longer freezing episodes, accounting for over 50% of the total duration of observation, whereas those exposed to old saliva exhibited freezing episodes for less than 20% of the total duration on average (Figure 1E). We found no significant difference in the latency to the first freezing episode from the first contact with a swab between mice exposed to fresh and old saliva (Figure 1F). Importantly, mice exposed to a swab containing either fresh or old saliva significantly avoided interacting with the swab, while mice exposed to a clean control swab spent a significant amount of time interacting with it, including direct sniffing, playing, and chewing (Figure 1G). Additionally, the stress hormone adrenocorticotrophic hormone (ACTH) in plasma showed a trend towards two-fold upregulation after exposure to both fresh and old saliva (Figure 1H). These results suggest that mice can still recognize predator cues in old saliva, but freezing behavior in response to the cues attenuates quickly over time.

Taken together, this behavior study indicates that cat saliva contains predator cues that signal the imminence of predator threat and differentially induce defensive behavioral outputs in mice. More specifically, fresh saliva is recognized as an immediate danger and induces more robust freezing behavior, while old saliva is recognized as a trace of danger and induces less robust freezing behavioral responses.

Cat saliva activates VNO neurons and modulates freezing behavior

As the freezing behavior was induced when mice made direct contact with the saliva-containing swab, we postulated that direct contact with the predator cues in cat saliva is necessary for freezing behavior induction. To examine the necessity of direct contact, we placed a cotton swab with or without cat saliva into a porous container and investigated mouse's behavior toward the container. The pores on the container allow volatile chemicals in cat saliva to diffuse out, while nonvolatile chemicals stay inside of the container with the swab. Interestingly, we found that when we introduced a container containing a swab with fresh saliva into the home cage, the mice no longer exhibited freezing behavior towards the container, but instead interacted with it

similarly to the control swab-containing container (**Figure 2A**, B). This suggests that freezing behavior requires direct contact with the saliva-containing swab and that the freezing-inducing cues in cat saliva detected in this scheme are nonvolatile.

Mice have the ability to detect nonvolatile chemical cues through the VNO (Mohrhardt et al., 2018). Ethologically relevant chemical cues from different animals, such as pheromones and predator cues, are sucked up into the VNO and activate VNO sensory neurons when animals contact the chemical source (Luo et al., 2003). We, therefore, explored the role of the VNO in inducing freezing in response to cat saliva using a VNO-deficient laboratory mouse line that lacks the primary signal transduction ion channel in VNO neurons (transient receptor potential cation channel, subfamily C, member 2-knockout line or Trpc2-KO line) (Leybold et al., 2002). In Trpc2-KO mice, chemosensory signals detected by sensory receptor proteins are expected to be attenuated before being relayed to the brain, due to the lack of the downstream transduction channel in VNO neurons (Leybold et al., 2002; Liman et al., 1999; Stowers et al., 2002). Thus, Trpc2-KO mice have been widely utilized to examine roles of the VNO in various behaviors (Beny-Shefer et al., 2017; Ferrero et al., 2013; Haga et al., 2010; Kimchi et al., 2007; Leybold et al., 2002; Papes et al., 2010; Pérez-Gómez et al., 2015; Stowers et al., 2002). Upon exposure to a fresh saliva-containing swab, Trpc2-KO mice failed to exhibit freezing behavior, in contrast to their wild-type littermates who displayed freezing responses. Additionally, the Trpc2-KO mice interacted with the saliva swab with a similar level of intensity as the control swab, while the wild-type littermates actively avoided it (**Figure 2C**, D). These findings suggest that VNO activation, disrupted in Trpc2-KO mice, is necessary for the freezing behavior observed in response to cat saliva in mice.

To investigate if VNO neurons are indeed activated by cat saliva, we analyzed immunoreactivity (IR) of phosphorylated ribosomal protein S6 subunit, pS6, in the VNO of mice exposed to cat saliva. As the cellular pS6 level is greatly upregulated upon activation of VNO neurons, it has been used as a sensitive marker to examine neural activation in the VNO (Carvalho et al., 2020; Itakura et al., 2022; Osakada et al., 2022; Tsunoda et al., 2018). We observed that exposure to both fresh and old cat saliva induced an increase in pS6-IR in the basal VNO neurons that express Gao protein (**Figure 2E**, G). Conversely, pS6-IR was rarely observed in the apical VNO neurons (**Figure 2F**). Notably, we found that the number of pS6-positive neurons was more than three times higher in mice exposed to fresh saliva compared to that exposed to old saliva (**Figure 2G**). Strikingly, when the duration of freezing was plotted as a function of pS6-IR signals in the VNO of individual animals, we observed a significantly positive correlation ($p = 0.0108$, $R^2: 0.2608$) between the numbers of active sensory neurons and behavioral outputs (**Figure 2H**), suggesting that more intense freezing behavior is induced when more VNO neurons are activated by cat saliva.

Taken together, these results indicate that freezing-inducing predator cues are detected by Gao-positive sensory neurons in the VNO, and these sensory neurons convey the imminence of predator threat to the higher brain center.

Cat saliva activates the V2R-A4 subfamily of the vomeronasal receptors

The significant reduction of the freezing response to aged saliva may reflect attenuation of signal inputs to a specific population of sensory receptors, or alternatively, it may reflect a change of sensory receptor repertoires due to qualitative changes of predator cues over time. To test these possibilities, we examined sensory receptors expressed in the neurons activated by fresh and old saliva. The basal VNO neurons express sensory receptors that belong to vomeronasal receptor type 2 (V2R) family of G protein-coupled receptors in mice (**Figure 3A**) (Mohrhardt et al., 2018). The mouse V2R family consists of over 120 intact genes, which are divided into V2R A1-6, A8-10, and B-D subfamilies based on sequence homology (**Figure 3A**) (Francia et al., 2014);

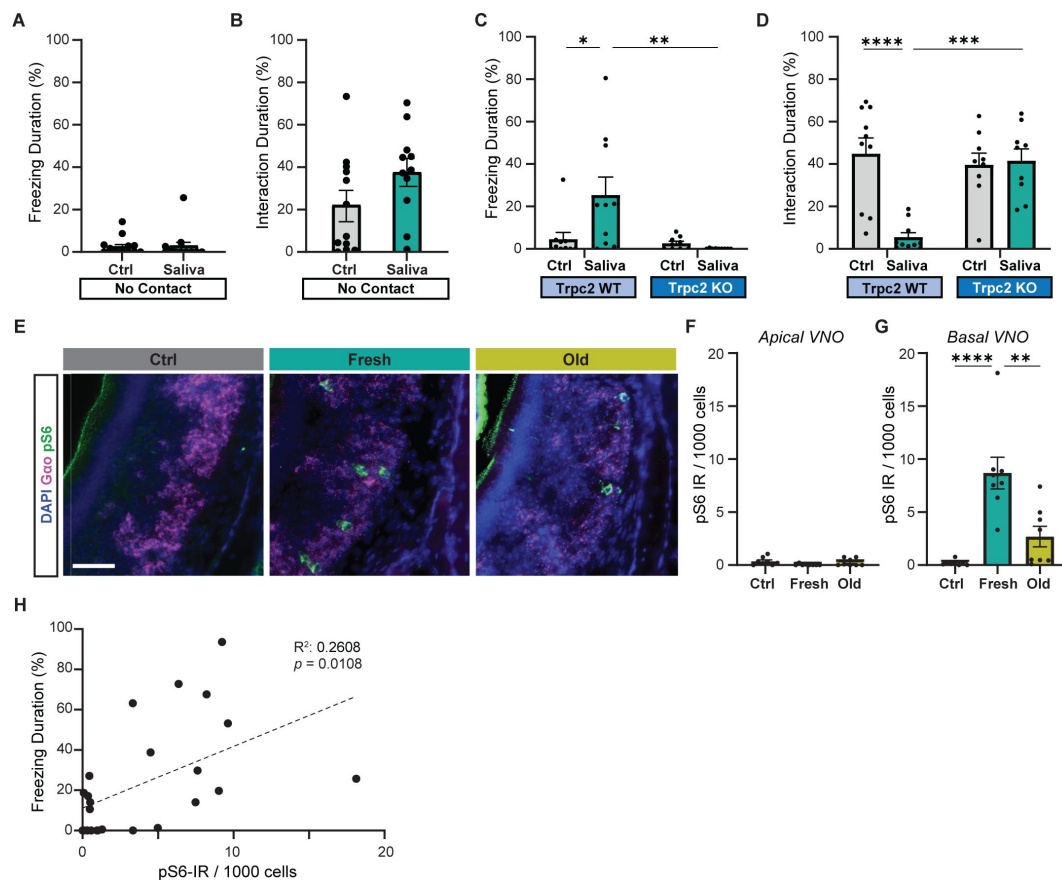


Figure 2

Cat saliva activates VNO neurons and modulates freezing behavior

(A) The percentage of total freezing episodes directed towards control or fresh saliva swabs in cassettes that prevent direct interaction with the swabs.

(B) The percentage of total interaction episodes with cassettes containing control or fresh saliva swabs.

(C) The percentage of total freezing episodes directed toward control or fresh saliva swabs in Trpc2-WT or -KO mice.

(D) The percentage of total interaction episodes with control or fresh saliva swabs in Trpc2-WT or -KO mice.

(E) Representative images illustrating the expression of pS6 (green) and Gao (magenta) in the VNO of mice exposed to control, fresh, or old saliva swabs. DAPI was used as a nuclear counterstain. Scale bar: 50 μ m.

(F, G) The number of pS6-IR positive cells in 1000 VNO cells in the apical (F) and basal (G) regions of the VNO neuroepithelium.

(H) The percentage of total freezing episodes in relation to the number of pS6-positive cells in the VNO of individual mice. Correlation analysis was conducted using Pearson's Correlation.

(A - D, F, G) the values are presented as means $\pm$ S.E.M., with individual dots representing individual mice. Statistical analysis was performed T-tests (A, B), Two-Way ANOVA (C, D), and One-Way ANOVA (F, G) with Tukey's multiple comparisons. *, **, ***, and **** denote significance levels of $p < 0.05$, 0.01 , 0.001 , and 0.0001 , respectively.

Silvotti et al., 2011 [↗](#), 2007 [↗](#)). With the exception of the V2R C group that is co-expressed with other groups of V2Rs in the basal VNO neurons, each sensory neuron expresses only one member of the V2R receptor, allowing individual neurons to respond to specific molecules (Mohrhardt et al., 2018 [↗](#)).

To investigate which subfamilies of V2R are expressed in cat saliva-activated VNO neurons, we designed *in situ* hybridization (ISH) probes that detect individual V2R subfamilies and performed dual staining of V2R ISH with pS6-IR (**Supplemental Figure 1** [↗](#)). We found that the majority of pS6-IR signals induced by cat saliva co-localized with signals of the ISH probe against the V2R-A4 subfamily (**Figure 3B** [↗](#), **Supplemental Figure 1** [↗](#)). Approximately 45% of pS6-IR signals induced by fresh saliva (83 out of 181) co-localized with V2R-A4 probe signals, with less than 1 % of pS6-IR (1 out of 120) co-localized with A8-4 probe signals (**Figure 3C** [↗](#)).

Notably, V2R-A4 probe signals also co-localized with approximately 45% of pS6-IR signals induced by old saliva (37 out of 86). None of the other probe signals co-localized with pS6-IR induced by both fresh and old saliva. These results indicate that members of the V2R-A4 subfamily are potential sensory receptors for the predator cues contained in both fresh and old cat saliva, and varying levels of imminence of predator threat stimulate the same type of sensory receptors in a quantitatively different manner to regulate the freezing behavior.

Cat saliva activates neurons in the AOB and VMH

We next aimed to understand how the differential activation of sensory receptor neurons in the VNO by different freshness of cat saliva influences behavioral decisions through downstream neural circuits (**Figure 4A** [↗](#)). We investigated this by examining the induction of cFos, an immediate early gene product, in the projection neurons (mitral cells) and local inhibitory neurons (granule cells) in the AOB of mice exposed to cat saliva. The axons of apical and basal VNO neurons project to the anterior and posterior regions of the AOB (aAOB and pAOB), respectively (Mohrhardt et al., 2018 [↗](#)). We found that the activation of the basal VNO neurons by cat saliva leads to predominant induction of cFos-IR signals in the pAOB, whose glomerulus layer is Gai-IR negative (**Figure 4B-F** [↗](#)). The numbers of cFos-positive mitral and granule cells in the pAOB in saliva-exposed mice are significantly higher than those in mice exposed to the control swab (**Figure 4D** [↗](#), F). However, contrary to our expectations, we did not observe any differences in the numbers of cFos-positive mitral and granule cells in the AOB between fresh and old saliva-exposed mice (**Figure 4D** [↗](#), F). These results suggest that, although the AOB processes saliva signals from the VNO following predator cue detection, the relationship between the degree of AOB activation and freezing behavior intensity is non-linear.

To further investigate how signals of fresh and old saliva are processed in the brain, we examined induction of cFos in the VMH, which receives indirect ascending inputs from the mitral cells in the AOB (Mohrhardt et al., 2018 [↗](#)). Importantly, the dorsomedial and central subdivisions of VMH (VMHdm/c) have been shown to contain subpopulations of neurons that control freezing behavior (Kunwar et al., 2015 [↗](#); Wang et al., 2015 [↗](#)) (**Figure 4A** [↗](#)). Upon exposure to fresh and old saliva, we observed a significant increase in cFos expression in the VMH compared to exposure to control stimulus (**Figure 4G** [↗](#), H). However, there was no significant difference in the number of cFos-positive cells between the two saliva-exposed groups (**Figure 4H** [↗](#)). These results suggest that, although the VMH is activated by cat saliva signals, the relationship between the degree of VMH activation and freezing behavior intensity is non-linear, similar to the situation in the AOB.

Taken together, these results indicate that the relationship between neural activity and behavioral output is more complex in the AOB and VMH than in the VNO. More specifically, the numbers of active neurons and intensity of freezing behavior do not appear to correlate in these regions. This suggests that fresh and old cat saliva may differentially activate functionally distinct neural populations within these regions, which in turn leads to quantitatively different freezing behavioral responses.

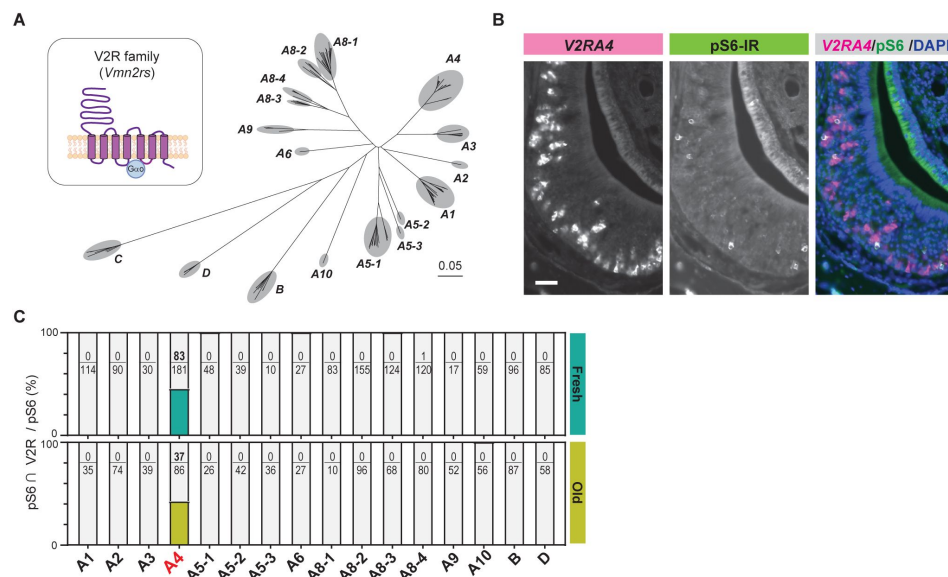


Figure 3

Cat saliva activates the V2R-A4 subfamily of the vomeronasal receptors

(A) Phylogenetic tree of the V2R family and its subfamilies used for V2R cRNA probe design.

(B) Representative images displaying the expression of V2RA4 (magenta) and pS6 (green) in the VNO of a mouse exposed to a fresh saliva swab. DAPI was utilized as a nuclear counterstain. Scale bar: 50 μ m.

(C) The percentage of cells co-labeled with V2R probes and anti-pS6 antibody among the total pS6-positive cells in the VNO of mice exposed to fresh saliva (top) or old saliva (bottom).

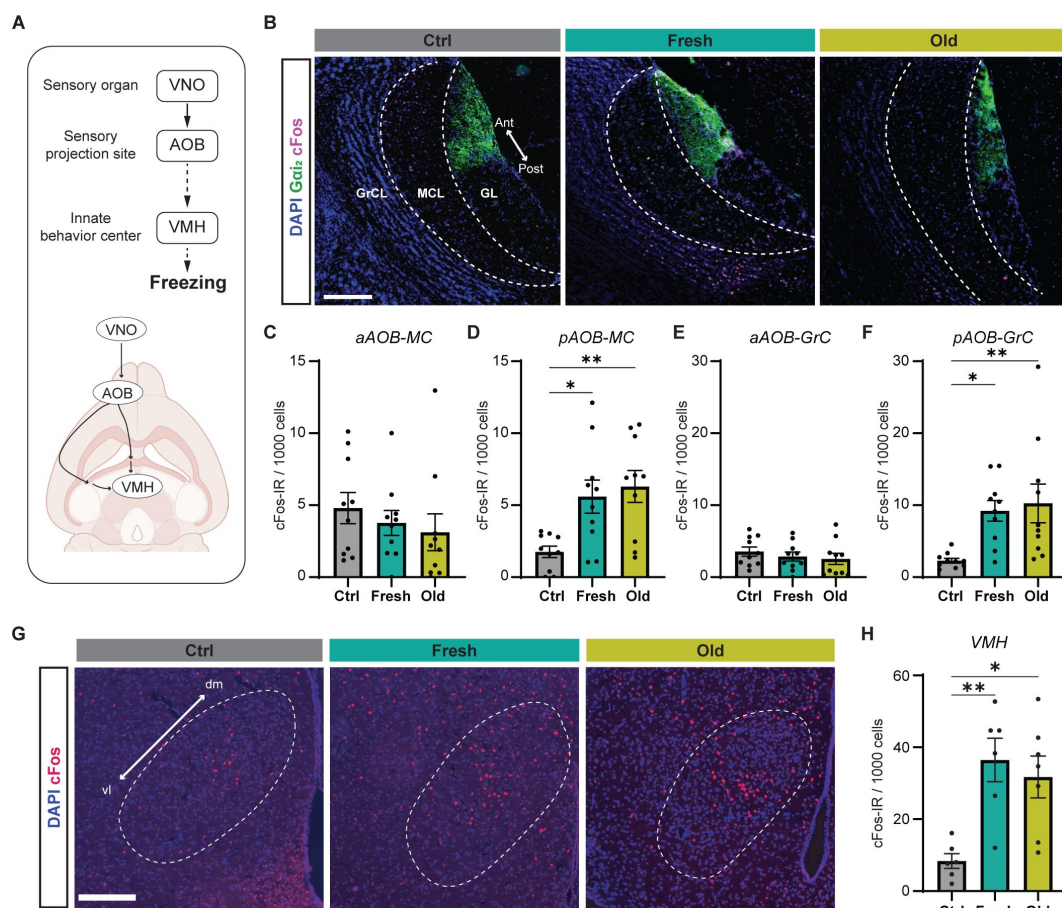


Figure 4

Cat saliva activates neurons in the AOB and VMH

(A) Schematic diagrams illustrating the VNO sensory pathway to the VMH.

(B) Representative images displaying the expression of cFos (magenta) in the mitral cell (MCL) and granule cell (GCL) layers of the AOB of mice exposed to control, fresh, or old saliva swabs. Gai2 (green) visualizes the anterior region of the glomerular layer (GL). DAPI was utilized as a nuclear counterstain. Scale bar: 200 μ m.

(C, D) The number of cFos-IR positive mitral cells per 1000 mitral cells in the anterior **(C)** and posterior **(D)** regions of the AOB.

(E, F) The number of cFos-IR positive granule cells per 1000 granule cells in the anterior **(E)** and posterior **(F)** regions of the AOB.

(G) Representative images displaying the expression of cFos (red) in the VMH of mice exposed to control, fresh, or old saliva swabs. The VMH is delineated by a dashed white line, defined by counterstaining with DAPI. Scale bar: 200 μ m.

(H) The number of cFos-IR positive neurons per 1000 cells in the VMH.

In **(C-F, H)**, the values are presented as means $\pm$ S.E.M., with individual dots representing individual mice. Statistical analysis was performed using One-Way ANOVA with Tukey's multiple comparisons. * and ** denote significance levels of $p < 0.05$ and 0.01, respectively.

Fresh and old cat saliva differentially activate VMH neurons to control defensive behavior

To investigate the potential differences between fresh and old saliva-activated neurons, we analyzed correlations of the duration of freezing behavior and numbers of cFos-IR signals in different cell types in individual animals from different exposure groups. To our surprise, we found that the numbers of cFos-IR signals in AOB mitral and granule cells did not significantly correlate with the duration of freezing behavior in any of the exposure groups (**Figure 5A** [↗](#), B). However, we observed a strong positive correlation between the number of cFos-IR signals and behavior intensity only in the fresh saliva-exposed group in the VMH ($p = 0.0412$, $R^2 = 0.5708$), while no such correlation was observed in the control or old saliva-exposed groups (**Figure 5C** [↗](#)). These findings support our hypothesis that fresh and old saliva may activate distinct populations of neurons in the VMH.

We therefore examined whether fresh and old cat saliva activate neurons in spatially distinct regions within the VMH. We observed the distribution of cFos-IR neurons in five serial brain sections from rostral to caudal VMHdm regions. The VMH region within each section was further divided into 10 bins along the dorsomedial-ventrolateral axis, and cFos-IR signals in each bin were counted (**Figure 6A** [↗](#)). The majority of cFos-IR signals induced by both fresh and old cat saliva were localized in the dorsomedial to central part of the VMH (bins 1 - 5). Strikingly, the numbers of cFos-IR signals induced by fresh and old saliva were significantly different in the section 2 bin 5, as well as the section 3 bin 3 and bin 4, which are adjacent to each other (**Figure 6B** [↗](#)). The cFos-IR signals in the other regions did not show significant difference between fresh and old saliva-exposed mice. These results suggest that there is a distinct population of neurons clustered in a close proximity within the VMH, which is more sensitive to fresh saliva than old saliva. This population of neurons in the VMHdm may play a role in inducing robust freezing behavior to fresh cat saliva. Moreover, these results also suggest that the neurons that induce robust freezing to fresh saliva and less robust freezing to old saliva reside in the same region of the VMH and may use different mechanisms to encode freezing intensity. Overall, our results suggest that the qualitative properties of predator cue inputs, such as their freshness, can induce distinct neural processing and ultimately lead to different defensive behavioral outcomes.

Discussion

This study provides evidence for the role of cat saliva as a source of predator cues that signal imminent predator threat and regulate the intensity of freezing behavior through the accessory olfactory pathway in mice (**Figure 7** [↗](#)). Notably, both fresh and old saliva activate neurons expressing the V2R-A4 subfamily of sensory receptors, suggesting the existence of a specific receptor group responding to the predator cues. Importantly, the number of vomeronasal neurons activated in response to saliva correlates with the freshness of saliva and the intensity of freezing behavior. Moreover, the number of VMH neurons activated by immediate threat (fresh saliva) exhibits a positive correlation with the intensity of behavioral outputs, suggesting their direct involvement in the modulation of freezing behavior. In contrast, the activation of VMH neurons by less immediate threat (old saliva) does not show such a correlation. These contrasting responses to different freshness levels of cat saliva suggest the presence of distinct populations of neurons in the VMH controlling the intensity of freezing behavior. These findings establish a framework for understanding the neural mechanisms underlying innate defensive behavior decisions.

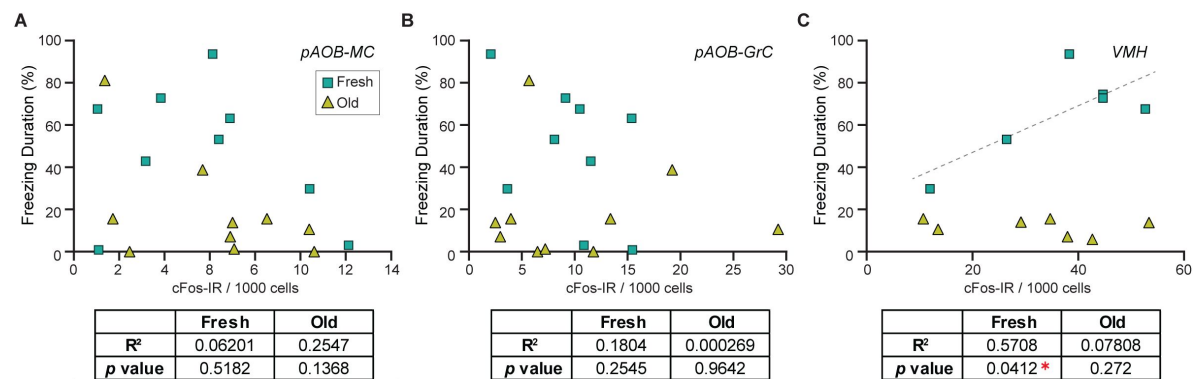


Figure 5

Fresh and old cat saliva differentially activate VMH neurons to control defensive behavior

(A-C) Percentage of the total freezing episodes as a function of the number of cFos-positive posterior AOB mitral cells (A) and granule cells (B), as well as VMH neurons (C). Green, and yellow symbols represent values from mice exposed to control, fresh saliva, and old saliva, respectively. Correlation was analyzed using Pearson's Correlation. * denotes statistical significance with $p < 0.05$.

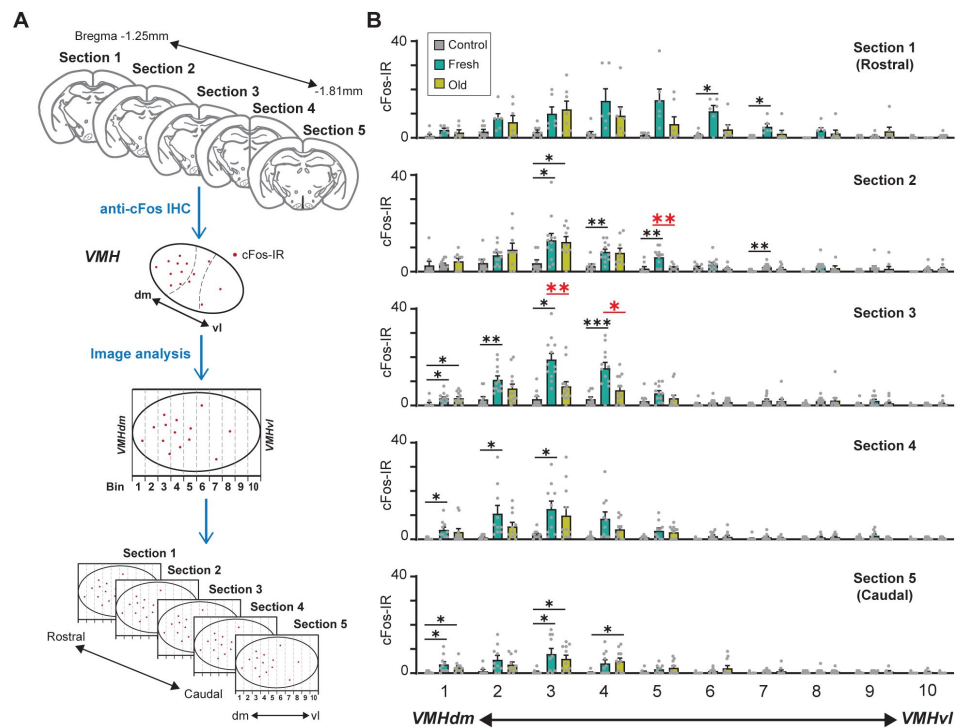


Figure 6

A population of VMH neurons is more sensitive to fresh saliva than old saliva

(A) Schematic diagram illustrating the cFos-IR distribution analysis. Each VMH region in five consecutive sections (140 μ m apart) was prepared along the dorsomedial (dm)-ventrolateral (vl) axis and divided into 10 bins that are vertical to the dm-vl axis. cFos-IR signals in each bin were counted.

(B) The number of cFos-IR-positive cells in each bin in the VMH sections. The values are presented as means + S.E.M., with individual dots representing values from individual VMH. Statistical analysis was performed using Two-Way ANOVA with Tukey's multiple comparisons. Black * and ** denote significance levels of $p < 0.05$ and 0.01 , respectively, between control and fresh or old saliva conditions. Red * and ** denote significance levels of $p < 0.05$ and 0.01 , respectively, between fresh and old saliva conditions.

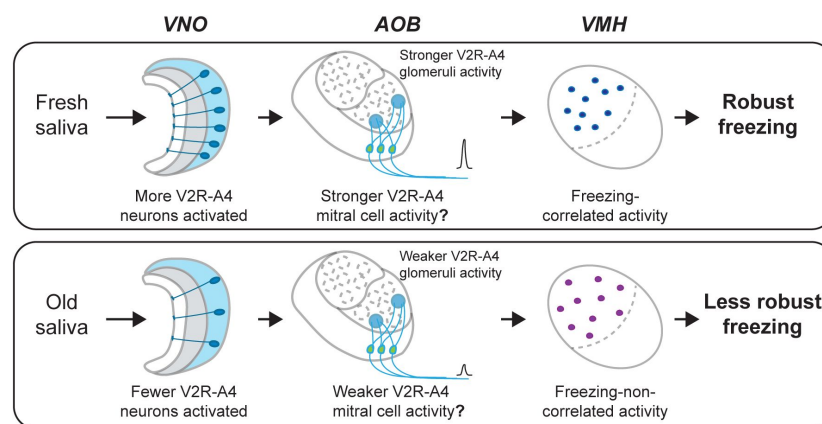


Figure 7

A model of the VMH-to-VMH pathway processing imminence of predator threat

The circuitry is selectively activated based on the varying imminence of predator cues. Fresh saliva activates more V2R-A4-expressing VNO neurons (blue), which results in stronger activation of the mitral cells innervating V2R-A4 glomeruli. This eventually triggers freezing-correlated neural activity in the VMH. In contrast, old saliva activates fewer V2R-A4-expressing VNO neurons (blue), which results in weaker activation of the mitral cells innervating V2R-A4 glomeruli, leading to freezing-non-correlated activity in the VMH.

The VNO as the sensor of predator cues that induce fear-related behavior

This study demonstrates that the VNO is involved in detecting and processing chemosensory predator signals that induce fear-related freezing behavior. Previous studies have shown that the VNO mediates defensive behaviors against predators that are rather related to anxiety state, such as risk assessment, area avoidance, and reduced exploratory and locomotion behaviors (Isogai et al., 2011 [↗](#); Papes et al., 2010 [↗](#); Pérez-Gómez et al., 2015 [↗](#); Tsunoda et al., 2018 [↗](#)). The difference between these studies and our current study is likely due to different sample types and their freshness, as we observed that fear-related behavior is correlated with freshness of predator samples. It is interesting to assume that the biological sources of predator cues, such as saliva, urine, and tears, contain varieties of unidentified cues that convey time-sensitive signals through the mouse accessory olfactory system. This is consistent with the fact that VNO neurons expressing more than half of vomeronasal receptors and a significant fraction of AOB mitral cells have been shown to be activated by predator-derived samples (Ben-Shaul et al., 2010 [↗](#); Isogai et al., 2011 [↗](#)). Therefore, it is important to investigate how the sensory pathway in the accessory olfactory system processes varieties of predator cues, which eventually leads to defensive behavioral decisions in higher brain centers.

V2R-A4 subfamily as the receptor for predator cues in cat saliva

Screening of sensory receptors in the VNO neurons activated by fresh and old saliva demonstrated that members of the V2R-A4 subfamily are the candidate receptors for the predator cues in cat saliva. The V2R-A4 subfamily has been shown to be expressed in the VNO neurons that are activated by biological samples derived from various mammalian predators (Carvalho et al., 2020 [↗](#); Isogai et al., 2011 [↗](#); Tsunoda et al., 2018 [↗](#)). Therefore, it is likely that specific members of the V2R-A4 subfamily are the receptors for the predator cues in cat saliva that induce freezing behavior.

Dual staining of vomeronasal receptors with immediate early gene products has been effectively utilized to identify receptors for several pheromones (Carvalho et al., 2020 [↗](#); Haga et al., 2010 [↗](#); Isogai et al., 2018 [↗](#), 2011 [↗](#); Itakura et al., 2022 [↗](#); Kimoto et al., 2005 [↗](#); Osakada et al., 2022 [↗](#), 2018 [↗](#); Tsunoda et al., 2018 [↗](#)). However, due to very high sequence homologies among members of the V2R-A4 subfamily (Francia et al., 2014 [↗](#)), alternative approaches need to be considered to further isolate the receptors responsible for the detection of predator cues in cat saliva. Potential approaches include receptor mRNA profiling with pS6 immunoprecipitation from the VNO of mice stimulated by cat saliva (Isogai et al., 2018 [↗](#); Jiang et al., 2015 [↗](#)), as well as receptor mRNA profiling from isolated single cells activated by cat saliva in GCaMP imaging using the VNO slices *in vitro* (Haga-Yamanaka et al., 2014 [↗](#); Wong et al., 2020 [↗](#)). These approaches are technically challenging and may provide false positive results. Therefore, identified receptors need to be further confirmed by examining necessity and sufficiency for detecting cat saliva using genetically modified mouse lines (Haga et al., 2010 [↗](#); Haga-Yamanaka et al., 2014 [↗](#); Isogai et al., 2018 [↗](#); Itakura et al., 2022 [↗](#); Osakada et al., 2022 [↗](#), 2018 [↗](#); Tsunoda et al., 2018 [↗](#)).

Differential processing of fresh and old saliva signals in the VNO-to-VMH pathway

Our correlation analyses between the number of activated neurons and behavior intensity suggest that sensory signals encoding different imminence of predator threat are differentially processed in the VNO-to-VMH pathway. The differential behavioral responses do not appear to result from global neuromodulations of the neural circuit by stress hormones, as differential endocrinological responses were not observed (**Figure 1H** [↗](#)). This suggests that more specific neural processing happens in the circuitry. In the circuitry downstream of the VNO, sensory signals can converge and diverge from the AOB before reaching the VMH. The axon terminals of hundreds of VNO neurons expressing the same receptor converge into a few specific glomerulus structures

(Belluscio et al., 1999 [↗](#); Del Punta et al., 2002 [↗](#); Rodriguez et al., 1999 [↗](#)), and, in some cases, neurons expressing receptors in the same subfamily send axons into the same glomeruli (Wagner et al., 2006 [↗](#)). Individual AOB mitral cells extend multiple primary dendrites into multiple glomeruli (Imamura et al., 2020 [↗](#); Larriva-Sahd, 2008 [↗](#); Takami and Graziadei, 1991 [↗](#); Wagner et al., 2006 [↗](#); Yonekura and Yokoi, 2008 [↗](#)). In turn, efferent axons of the mitral cells project to multiple brain substrates, namely the BNST, the MEA, and the posteromedial cortical amygdaloid nucleus (PMCo) (Davis et al., 1978 [↗](#); von Campenhausen and Mori, 2000 [↗](#)). These structural features indicate that sensory inputs to the AOB mitral cells are converged, and outputs from the mitral cells may be diverged.

These features of the ascending neural pathway within the AOB may contribute to the sorting of sensory signals derived from fresh and old saliva. Fresh saliva activates more V2R-A4-expressing VNO neurons than old saliva, which may result in stronger activation of V2R-A4 glomeruli in fresh saliva-exposed mice compared to those exposed to old saliva. This differential activation of V2R-A4 glomeruli by fresh and old saliva may lead to activation of the mitral cells innervating the V2R-A4 glomeruli with different excitation levels. We assume that such differential activations of the mitral cells between fresh and old saliva result in the differential activation of VMH neurons (Figure 7 [↗](#)). The methodology that we undertook in this study, cFos-immunohistochemistry, only provides information about neurons responding to cat saliva above a given threshold. It is therefore important to examine how the mitral cells respond to fresh and old saliva using more quantitative techniques, such as *in vivo* electrophysiological recording, in future studies.

Differential activation of VMH neurons potentially underlying distinct intensities of freezing behavior

Our findings indicate that the imminent predator cues present in cat saliva activate a specific subpopulation of VMH neurons, whose overall activity positively correlates with the intensity of freezing behavioral outputs (Figure 5C [↗](#)). In contrast, the less imminent predator cues activate a subpopulation of VMHdm/c neurons, whose overall activity does not correlate with the intensity of freezing behavior (Figure 5C [↗](#)). These findings suggest that induction of robust freezing behavior is determined by a distinct set of VMH neurons, and intensity of freezing behavior is encoded separately, possibly by the intensity of the neuronal firing, which cannot be detected by the cFos-IR used in this study. The VMHdm/c region contains several subpopulations of neurons that coordinate various defensive behavioral responses (Silva et al., 2016 [↗](#)). Previous studies have demonstrated that optogenetic stimulation of VMHdm/c neurons induces freezing/immobility responses, suggesting that these neurons mediate freezing behavior in response to predator cues (Kunwar et al., 2015 [↗](#); Wang et al., 2015 [↗](#)). In our current study, the distribution of cFos-IR in the VMH revealed a population of neurons clustered in close proximity that are more sensitive to fresh saliva than old saliva (Figure 6B [↗](#)). Therefore, we propose that imminent predator cues in cat saliva may activate these specific neurons with certain excitation levels to directly induce robust freezing behavior, while less imminent predator cues in cat saliva may use a different mechanism to regulate freezing behavior.

What distinguishes the subpopulation of VMHdm/c neurons activated by fresh saliva from those activated by old saliva? One possibility is that these neurons stimulate distinct downstream neural circuitries. The VMHdm/c neurons project to two major nuclei, the anterior hypothalamic nucleus (AHN) and the periaqueductal gray (PAG), which directly influence different aspects of defensive behavior (Canteras et al., 1994 [↗](#); Wang et al., 2015 [↗](#)). Specifically, selective activation of the VMHdm/c-to-PAG pathway has been shown to cause freezing/immobility behavior, while activation of the VMHdm/c-to-AHN pathway rather promotes avoidance and induces immobility to a lesser extent (Wang et al., 2015 [↗](#)). It is possible that fresh saliva-activated neurons send inputs to the PAG, whereas old saliva-activated neurons send inputs to the AHN and/or other projection sites. Thus, the number of old saliva-activated cells may correlate with different aspects of defensive responses, rather than freezing behavior.

Given the observation from a previous study that neurons projecting to the PAG also send collateral projections to the AHN, and vice versa, neuronal populations activated by fresh and old saliva may or may not overlap. If the two sets of neurons overlap, it is possible that the strength of presynaptic inputs to the VMHdm/c differs between fresh and old saliva stimulation, which leads to differential activation of downstream pathways. Previous studies have shown that optogenetic stimulation of the VMHdm/c with different stimulus frequencies can induce different types of defensive behaviors (Wang et al., 2015 [DOI](#)), suggesting that distinct activity patterns of fresh and old saliva-responsive neurons in the VMHdm/c may contribute to differences in defensive responses. Investigating these activity patterns would be an intriguing direction for future research.

In summary, our study proposes a predator defensive circuitry that involves the accessory olfactory system and the VMHdm/c. Understanding the precise neural mechanisms of how different degrees of imminence of predator threat are processed in this circuit will provide valuable insights into the decision-making processes underlying defensive behaviors in animals.

Materials and methods

Animals

The experimental procedures were approved by the Institutional Animal Care and Use Committee at the University of California, Riverside (UCR) and were conducted in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals. C57BL/6N mice originally obtained from Charles River (Catalog# 027) were bred at the UCR vivarium. The *Trpc2* knock-out mouse line was generously provided by Dr. Richard Axel Lab at Columbia University. All mice were housed in the vivarium under a reversed light cycle (lights on at 21:00; lights off at 9:00) for a minimum of 2 weeks prior to the behavior and histology experiments. Cats used in this study were obtained from the Nutrition and Pet Care Center at the University of California, Davis (a class A dealer) and were also housed in the UCR vivarium.

Collection of saliva from cats

A person familiar with the cat gently held it for all procedures. A sterile cotton swab was inserted into the cat's mouth where the cat chewed and licked on the swab for at least 20 seconds or until the swab is saturated with saliva. The swabs containing saliva were then placed in microcentrifuge tubes and stored until used. Fresh saliva was collected within 15 minutes prior to its use in the mouse behavioral study, while old saliva was collected 4 hours prior to the study. The old saliva was kept at room temperature until it was used.

Behavioral assay

All behavioral experiments were conducted during the animals' dark cycle. Mice were placed in their experimental room and allowed approximately 4 hours to acclimate to the ambient conditions. A cat saliva-containing swab was manually introduced into the animal's home cage to assess the defensive behavioral responses of resident mice. Recording of mouse behaviors commenced 2 minutes prior to introducing the sample and concluded 10 minutes thereafter. If tissues were required for histology, the animals were euthanized after 60 minutes, while for plasma collection, euthanasia took place after 15 minutes. Following the introduction of the sample, the behavioral room remained vacant of any individuals until the animal was euthanized. Mouse behavior was captured by night vision cameras equipped with infrared lights.

Behavioral analysis

Recordings of each behavioral session were saved and provided in a blinded manner to a researcher for analysis. The analysis was performed using a behavioral analysis program called Behavioral Observation Research Interactive Software (BORIS) (Friard and Gamba, 2016). Freezing behavior was defined as immobility lasting more than 2 seconds and was scored accordingly. Interaction was scored when there was physical contact with the swab. Animals that exhibited inactivity in their home cage for more than 50% of the pre-drop analysis duration in each experiment were excluded from the data set.

Histology and Immunohistochemistry (IHC)

After being exposed to the cat specimen or control swab for 60 minutes, the animals were euthanized with an overdose of sodium pentobarbital. Intracardial perfusions were performed using 4% paraformaldehyde (PFA) in PBS. The brains were extracted and post-fixed in 4% PFA overnight at 4°C. Subsequently, the brains were cryoprotected by sequentially immersing them in 15% and 30% sucrose solutions in PBS. The VNOs were also extracted and underwent the same post-fixation procedure, followed by decalcification with EDTA pH 8.0. Cryoprotection of the VNOs was achieved using a 30% sucrose solution in PBS. The cryoprotected tissues were then embedded in OCT (Optimal Cutting Temperature) media and frozen over liquid nitrogen. Tissues embedded in OCT were sectioned at 14 µm (VNO) and 20 µm (brain) thicknesses using a cryostat set at -25°C. The sections were placed onto positively charged microscope slides. The slides were stored at -80°C until stained. A standard frozen IHC protocol was employed. For VNO sections, staining was performed using a rabbit anti-pS6 antibody (Cell Signaling Technology, #4858) at a dilution of 1:200, incubated overnight at 4°C, followed by a secondary antibody, Goat anti-rabbit IgG Alexa Fluor 647 (Life Technologies #a21245), for 60 minutes at room temperature. For AOB sections, staining was carried out using a mouse anti-Gai2 antibody (Millipore #MAB3077MI) and a guinea pig anti-cFos antibody (Synaptic Systems #226 308) at dilutions of 1:500 and 1:1000, respectively. This was followed by incubation with a goat anti-mouse IgG2b Alexa Fluor 546 (Life Technologies #a21143) and a Goat anti-guinea pig IgG Alexa Fluor 647. For brain sections, staining was performed using a guinea pig anti-cFos antibody at dilutions of 1:1000 incubated overnight at 4°C, followed by Goat anti-guinea pig IgG Alexa Fluor 647. Finally, the slides were mounted using MS Shield Mounting Medium With 4,6-diamidino-2-phenylindole (DAPI) and DABCO™ (EMS #17989-20).

Enzyme-linked Immunosorbent Assay (ELISA)

Plasma collected from mice exposed to cat saliva or control swabs was frozen at -80°C before initiating the ELISA protocol. The ACTH ELISA kit (MD Bioproducts) was utilized, following the instructions provided with the kit.

In situ hybridization (ISH) and dual staining

Detection of *GaO* RNA in fixed frozen VNO sections was conducted using the ACD RNAscope® probe targeting GNAO1 (*GaO*) (ACD #444991) and the RNAscope® Multiplex Fluorescent Reagent Kit v2 (ACD #323100). Probe binding was visualized using Akoya Biosciences' Opal 570 (FP1488001KT) Dye at a dilution of 1:750 in RNAscope TSA Buffer. Following RNAscope staining, the slides were incubated in anti-pS6 antibody at 1:200 dilution overnight at 4°C, then detected by goat anti-rabbit IgG AlexaFluor 647 for one hour at room temperature.

Vmn2r (V2R) probes for the following genes were used: *Vmn2r13* for subfamily A1, *Vmn2r118* for A2, *Vmn2r117* for A3, *Vmn2r42* for A4, *Vmn2r69* for A5-1, *Vmn2r65* for A5-2, *Vmn2r76* for A5-3, *Vmn2r120* for A6, *Vmn2r98* for A8-1, *Vmn2r105* for A8-2, *Vmn2r58* for A8-3, *Vmn2r62* for A8-4, *Vmn2r81* for A9, *Vmn2r18* for A10, *Vmn2r24* for B, and *Vmn2r53* for D. Each cRNA probe was synthesized using the PCR-based method (Hua et al., 2018).

For detection of V2R RNA in ISH, VNO tissues were sectioned and placed onto slides up to one week before ISH and stored at -80 °C. Fluorescent ISH was performed following published protocols (Ishii et al., 2004). Briefly, the VNO slides were baked at 60 °C for 5 minutes and fixed with 4% PFA. The slides were then treated with 10 µg/mL Proteinase K (Sigma #3115828001), followed by being acetylated in 0.1 M Triethanolamine (Sigma #90279)-HCl with acetic anhydride (Sigma #320102). V2R cRNA probes were hybridized with tissue sections overnight at 65 °C in a hybridization solution consisting of 50 % formamide, 10 mM Tris-HCl pH 8.0, 200 µg/mL yeast tRNA, 10 % dextran sulfate, 1x Denhardt's, 600 mM NaCl, 0.25 % SDS, 1 mM EDTA pH 8.0. After hybridization, the slides were washed with 2x SSC in 50 % formamide for 30 minutes, then 2x SSC, 0.2x SSC, and 0.1x SSC in H₂O for 20 minutes each at 65°C. The slides were then blocked with blocking reagent (Perkin Elmer) and 0.5 % peroxidase block sequentially. Anti-DIG-POD (Millipore Sigma #11207733910) was used to detect DIG-labeled probes. TSA-Cy3 Plus kit (Akoya #NEL744001KT) was used to detect anti-DIG antibody at 1:200. The slides were incubated with anti-pS6 antibody at 1:200 dilution overnight at 4°C, then detected by goat anti-rabbit IgG AlexaFluor 647 for one hour at room temperature.

Following staining, the slides were mounted with MS Shield Mounting Medium with DAPI and DABCO™.

Image Quantification

Fluorescent images were captured at either 10X or 20X magnification using the Zeiss Axio Imager.M2 microscope. Whole slide images of IHC-stained brains were scanned through fluorescent imaging using Panoramic SCAN (3D Histech) at Reveal Biosciences (San Diego, CA). Automated counting of fluorescent positive cells was performed using a proprietary code developed in our laboratory within the QuPath software.

VMH cFos IR Quantification and Distribution Analysis

All VMH regions were identified by clusters of nuclei visualized through DAPI staining and manually annotated in QuPath using standard coronal mouse brain atlas as reference. Annotations were processed with a proprietary script for immunoreactivity quantification. Distribution analyses were performed in QuPath using a rectangular annotation set parallel to the dorsomedial-ventrolateral axis of each VMH region. This rectangular annotation was then split into 10 equal bins, and immunoreactivity quantification in each bin was automated in QuPath, where the detections were finally manually recorded.

Statistical Analysis

All statistical analyses were performed using Prism software (GraphPad). Behavioral analysis results and cFos-IR distribution in the VMH, involving two variables, were compared using a traditional Two-Way ANOVA with Tukey's multiple comparisons. Behavioral and immunohistological results were statistically compared using One-Way ANOVA with Tukey's multiple comparisons, assuming a parametric normal distribution. The correlation between immunoreactivity and behavior was analyzed using Pearson's Correlation.

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Author contributions

Conceptualization, S.H.-Y.; Methodology, Q.A.N., C.P., and S.H.-Y.; Investigation, Q.A.N., A.R., Y.Y., R.C., and C.S., and S.H.-Y.; Writing – Original Draft, Q.A.N and S.H.-Y.; Writing – Review & Editing, Q.A.N and S.H.-Y.; Supervision, S.H.-Y.; Funding Acquisition,

Q.A.N and S.H.-Y.

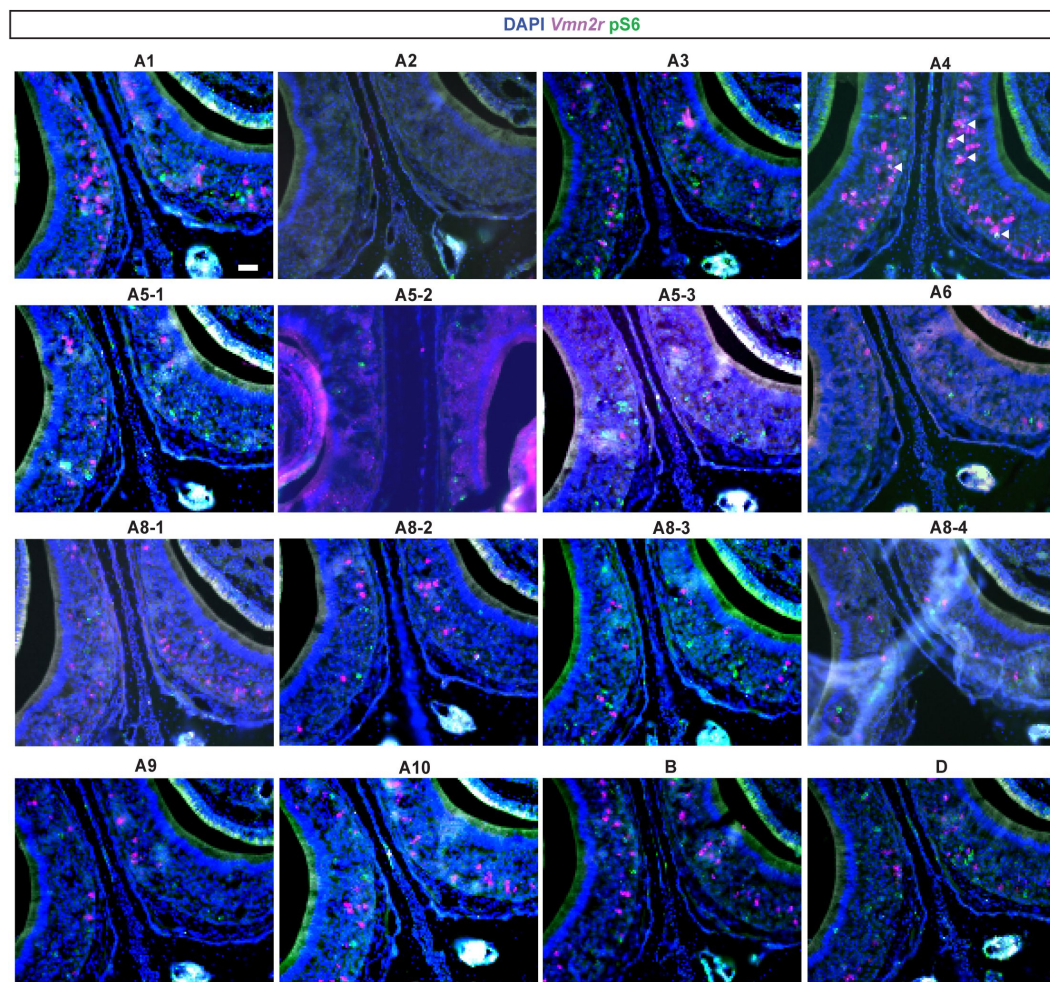
Declaration of interests

The authors declare no competing interests.

Supplemental movie 1 Mouse behavior toward a control swab.

Supplemental movie 2 Mouse behavior toward a swab containing fresh saliva.

Supplemental movie 3 Mouse behavior toward a swab containing old saliva.



Supplemental figure 1

Representative images of *V2R* in situ hybridization and pS6 immunohistochemistry for each *V2R* subfamily.

References

- Apfelbach R, Blanchard CD, Blanchard RJ, Hayes RA, McGregor IS (2005) **The effects of predator odors in mammalian prey species: a review of field and laboratory studies** *Neurosci Biobehav Rev* **29**:1123–1144 <https://doi.org/10.1016/j.neubiorev.2005.05.005>
- Belluscio L, Koentges G, Axel R, Dulac C (1999) **A map of pheromone receptor activation in the mammalian brain** *Cell* **97**:209–220 [https://doi.org/10.1016/s0092-8674\(00\)80731-x](https://doi.org/10.1016/s0092-8674(00)80731-x)
- Ben-Shaul Y, Katz LC, Mooney R, Dulac C (2010) **In vivo vomeronasal stimulation reveals sensory encoding of conspecific and allospecific cues by the mouse accessory olfactory bulb** *Proc Natl Acad Sci U S A* **107**:5172–5177 <https://doi.org/10.1073/pnas.0915147107>
- Beny-Shefer Y, Zilkha N, Lavi-Avnon Y, Bezalel N, Rogachev I, Brandis A, Dayan M, Kimchi T (2017) **Nucleus Accumbens Dopamine Signaling Regulates Sexual Preference for Females in Male Mice** *Cell Reports* <https://doi.org/10.1016/j.celrep.2017.11.062>
- Blanchard DC, Li CI, Hubbard D, Markham CM, Yang M, Takahashi LK, Blanchard RJ (2003) **Dorsal preammillary nucleus differentially modulates defensive behaviors induced by different threat stimuli in rats** *Neurosci Lett* **345**:145–148 [https://doi.org/10.1016/s0304-3940\(03\)00415-4](https://doi.org/10.1016/s0304-3940(03)00415-4)
- Blanchard RJ, Blanchard DC (1989) **Attack and defense in rodents as ethoexperimental models for the study of emotion** *Prog Neuropsychopharmacol Biol Psychiatry* **13** [https://doi.org/10.1016/0278-5846\(89\)90105-x](https://doi.org/10.1016/0278-5846(89)90105-x)
- Canteras NS (2002) **The medial hypothalamic defensive system: hodological organization and functional implications** *Pharmacol Biochem Behav* **71**:481–491 [https://doi.org/10.1016/s0091-3057\(01\)00685-2](https://doi.org/10.1016/s0091-3057(01)00685-2)
- Canteras NS, Simerly RB, Swanson LW (1994) **Organization of projections from the ventromedial nucleus of the hypothalamus: a Phaseolus vulgaris-leucoagglutinin study in the rat** *J Comp Neurol* **348**:41–79 <https://doi.org/10.1002/cne.903480103>
- Carvalho VM de A, Nakahara TS, Souza MA de A, Cardozo LM, Trintinalia GZ, Pissinato LG, Venancio JO, Stowers L, Papes F. (2020) **Representation of Olfactory Information in Organized Active Neural Ensembles in the Hypothalamus** *Cell Rep* **32** <https://doi.org/10.1016/j.celrep.2020.108061>
- Choi GB, Dong H-W, Murphy AJ, Valenzuela DM, Yancopoulos GD, Swanson LW, Anderson DJ (2005) **Lhx6 delineates a pathway mediating innate reproductive behaviors from the amygdala to the hypothalamus** *Neuron* **46**:647–660 <https://doi.org/10.1016/j.neuron.2005.04.011>
- Davis BJ, Macrides F, Youngs WM, Schneider SP, Rosene DL (1978) **Efferents and centrifugal afferents of the main and accessory olfactory bulbs in the hamster** *Brain Res Bull* **3**:59–72 [https://doi.org/10.1016/0361-9230\(78\)90062-x](https://doi.org/10.1016/0361-9230(78)90062-x)

- Del Punta K, Puche A, Adams NC, Rodriguez I, Mombaerts P (2002) **A divergent pattern of sensory axonal projections is rendered convergent by second-order neurons in the accessory olfactory bulb** *Neuron* **35**:1057–1066 [https://doi.org/10.1016/s0896-6273\(02\)00904-2](https://doi.org/10.1016/s0896-6273(02)00904-2)
- Dielenberg RA, McGregor IS (2001) **Defensive behavior in rats towards predatory odors: a review** *Neurosci Biobehav Rev* **25**:597–609 [https://doi.org/10.1016/s0149-7634\(01\)00044-6](https://doi.org/10.1016/s0149-7634(01)00044-6)
- Ferrero DM, Moeller LM, Osakada T, Horio N, Li Q, Roy DS, Cichy A, Spehr M, Touhara K, Liberles SD (2013) **A juvenile mouse pheromone inhibits sexual behaviour through the vomeronasal system** *Nature* **502**:368–371 <https://doi.org/10.1038/nature12579>
- Francia S, Silvotti L, Ghirardi F, Catzeflis F, Percudani R, Tirindelli R (2014) **Evolution of spatially coexpressed families of type-2 vomeronasal receptors in rodents** *Genome Biol Evol* **7**:272–285 <https://doi.org/10.1093/gbe/evu283>
- Friard O, Gamba M (2016) **BORIS : a free, versatile open-source event-logging software for video/audio coding and live observations** *Methods Ecol Evol* **7**:1325–1330 <https://doi.org/10.1111/2041-210x.12584>
- Gross CT, Canteras NS (2012) **The many paths to fear** *Nat Rev Neurosci* **13**:651–658 <https://doi.org/10.1038/nrn3301>
- Haga S, Hattori T, Sato T, Sato K, Matsuda S, Kobayakawa R, Sakano H, Yoshihara Y, Kikusui T, Touhara K (2010) **The male mouse pheromone ESP1 enhances female sexual receptive behaviour through a specific vomeronasal receptor** *Nature* **466**:118–122 <https://doi.org/10.1038/nature09142>
- Haga-Yamanaka S, Ma L, He J, Qiu Q, Lavis LD, Looger LL, Yu CR (2014) **Integrated action of pheromone signals in promoting courtship behavior in male mice** *Elife* **3** <https://doi.org/10.7554/eLife.03025>
- Hua R, Yu S, Liu M, Li H (2018) **A PCR-Based Method for RNA Probes and Applications in Neuroscience** *Front Neurosci* **12** <https://doi.org/10.3389/fnins.2018.00266>
- Imamura F, Ito A, LaFever BJ (2020) **Subpopulations of Projection Neurons in the Olfactory Bulb** *Front Neural Circuits* **14** <https://doi.org/10.3389/fncir.2020.561822>
- Ishii T, Omura M, Mombaerts P (2004) **Protocols for two-and three-color fluorescent RNA in situ hybridization of the main and accessory olfactory epithelia in mouse** *J Neurocytol* **33**:657–669 <https://doi.org/10.1007/s11068-005-3334-y>
- Isogai Y, Si S, Pont-Lezica L, Tan T, Kapoor V, Murthy VN, Dulac C (2011) **Molecular organization of vomeronasal chemoreception** *Nature* **478**:241–245 <https://doi.org/10.1038/nature10437>
- Isogai Y, Wu Z, Love MI, Ahn MH-Y, Bambah-Mukku D, Hua V, Farrell K, Dulac C (2018) **Multisensory Logic of Infant-Directed Aggression by Males** *Cell* **175**:1827–1841 <https://doi.org/10.1016/j.cell.2018.11.032>
- Itakura T, Murata K, Miyamichi K, Ishii KK, Yoshihara Y, Touhara K (2022) **A single vomeronasal receptor promotes intermale aggression through dedicated hypothalamic neurons** *Neuron* **110**:2455–2469 <https://doi.org/10.1016/j.neuron.2022.05.002>

- Jiang Y, Gong NN, Hu XS, Ni MJ, Pasi R, Matsunami H (2015) **Molecular profiling of activated olfactory neurons identifies odorant receptors for odors in vivo** *Nat Neurosci* **18**:1446–1454 <https://doi.org/10.1038/nn.4104>
- Kats LB, Dill LM (1998) **The scent of death: Chemosensory assessment of predation risk by prey animals** *Écoscience* **5**:361–394
- Kimchi T, Xu J, Dulac C (2007) **A functional circuit underlying male sexual behaviour in the female mouse brain** *Nature* **448**:1009–1014 <https://doi.org/10.1038/nature06089>
- Kimoto H, Haga S, Sato K, Touhara K (2005) **Sex-specific peptides from exocrine glands stimulate mouse vomeronasal sensory neurons** *Nature* **437**:898–901 <https://doi.org/10.1038/nature04033>
- Kunwar PS, Zelikowsky M, Remedios R, Cai H, Yilmaz M, Meister M, Anderson DJ (2015) **Ventromedial hypothalamic neurons control a defensive emotion state** *Elife* **4** <https://doi.org/10.7554/eLife.06633>
- Larriva-Sahd J (2008) **The accessory olfactory bulb in the adult rat: a cytological study of its cell types, neuropil, neuronal modules, and interactions with the main olfactory system** *J Comp Neurol* **510**:309–350 <https://doi.org/10.1002/cne.21790>
- LeDoux J (2012) **Rethinking the emotional brain** *Neuron* **73**:653–676 <https://doi.org/10.1016/j.neuron.2012.02.004>
- Leypold BG, Yu CR, Leinders-Zufall T, Kim MM, Zufall F, Axel R (2002) **Altered sexual and social behaviors in trp2 mutant mice** *Proc Natl Acad Sci U S A* **99**:6376–6381 <https://doi.org/10.1073/pnas.082127599>
- Liman ER, Corey DP, Dulac C (1999) **TRP2: a candidate transduction channel for mammalian pheromone sensory signaling** *Proc Natl Acad Sci U S A* **96**:5791–5796 <https://doi.org/10.1073/pnas.96.10.5791>
- Luo M, Fee MS, Katz LC (2003) **Encoding pheromonal signals in the accessory olfactory bulb of behaving mice** *Science* **299**:1196–1201 <https://doi.org/10.1126/science.1082133>
- Markham CM, Blanchard DC, Canteras NS, Cuyno CD, Blanchard RJ (2004) **Modulation of predatory odor processing following lesions to the dorsal premammillary nucleus** *Neurosci Lett* **372**:22–26 <https://doi.org/10.1016/j.neulet.2004.09.006>
- Mohrhardt J, Nagel M, Fleck D, Ben-Shaul Y, Spehr M (2018) **Signal Detection and Coding in the Accessory Olfactory System** *Chem Senses* **43**:667–695 <https://doi.org/10.1093/chemse/bjy061>
- Osakada T *et al.* (2022) **Hemoglobin in the blood acts as a chemosensory signal via the mouse vomeronasal system** *Nat Commun* **13** <https://doi.org/10.1038/s41467-022-28118-w>
- Osakada T, Ishii KK, Mori H, Eguchi R, Ferrero DM, Yoshihara Y, Liberles SD, Miyamichi K, Touhara K (2018) **Sexual rejection via a vomeronasal receptor-triggered limbic circuit** *Nat Commun* **9** <https://doi.org/10.1038/s41467-018-07003-5>
- Papes F, Logan DW, Stowers L (2010) **The vomeronasal organ mediates interspecies defensive behaviors through detection of protein pheromone homologs** *Cell* **141**:692–703 <https://doi.org/10.1016/j.cell.2010.03.037>

- Pérez-Gómez A, Bleymehl K, Stein B, Pyrski M, Birnbaumer L, Munger SD, Leinders-Zufall T, Zufall F, Chamero P (2015) **Innate Predator Odor Aversion Driven by Parallel Olfactory Subsystems that Converge in the Ventromedial Hypothalamus** *Curr Biol* **25**:1340–1346 <https://doi.org/10.1016/j.cub.2015.03.026>
- Petrovich GD, Canteras NS, Swanson LW (2001) **Combinatorial amygdalar inputs to hippocampal domains and hypothalamic behavior systems** *Brain Res Brain Res Rev* **38**:247–289 [https://doi.org/10.1016/s0165-0173\(01\)00080-7](https://doi.org/10.1016/s0165-0173(01)00080-7)
- Rodriguez I, Feinstein P, Mombaerts P (1999) **Variable patterns of axonal projections of sensory neurons in the mouse vomeronasal system** *Cell* **97**:199–208 [https://doi.org/10.1016/s0092-8674\(00\)80730-8](https://doi.org/10.1016/s0092-8674(00)80730-8)
- Samuelsen CL, Meredith M (2009) **The vomeronasal organ is required for the male mouse medial amygdala response to chemical-communication signals, as assessed by immediate early gene expression** *Neuroscience* **164**:1468–1476 <https://doi.org/10.1016/j.neuroscience.2009.09.030>
- Silva BA, Mattucci C, Krzywkowski P, Cuozzo R, Carbonari L, Gross CT (2016) **The ventromedial hypothalamus mediates predator fear memory** *Eur J Neurosci* **43**:1431–1439 <https://doi.org/10.1111/ejn.13239>
- Silvotti L, Cavalca E, Gatti R, Percudani R, Tirindelli R (2011) **A recent class of chemosensory neurons developed in mouse and rat** *PLoS One* **6** <https://doi.org/10.1371/journal.pone.0024462>
- Silvotti L, Moiani A, Gatti R, Tirindelli R (2007) **Combinatorial co-expression of pheromone receptors, V2Rs** *J Neurochem* **103**:1753–1763 <https://doi.org/10.1111/j.1471-4159.2007.04877.x>
- Stowers L, Holy TE, Meister M, Dulac C, Koentges G (2002) **Loss of sex discrimination and male-male aggression in mice deficient for TRP2** *Science* **295**:1493–1500 <https://doi.org/10.1126/science.1069259>
- Takahashi LK (2014) **Olfactory systems and neural circuits that modulate predator odor fear** *Front Behav Neurosci* **8** <https://doi.org/10.3389/fnbeh.2014.00072>
- Takami S, Graziadei PP (1991) **Light microscopic Golgi study of mitral/tufted cells in the accessory olfactory bulb of the adult rat** *J Comp Neurol* **311**:65–83 <https://doi.org/10.1002/cne.903110106>
- Tsunoda M, Miyamichi K, Eguchi R, Sakuma Y, Yoshihara Y, Kikusui T, Kuwahara M, Touhara K (2018) **Identification of an Intra- and Inter-specific Tear Protein Signal in Rodents** *Curr Biol* **28**:1213–1223 <https://doi.org/10.1016/j.cub.2018.02.060>
- von Campenhausen H, Mori K. (2000) **Convergence of segregated pheromonal pathways from the accessory olfactory bulb to the cortex in the mouse** *Eur J Neurosci* **12**:33–46 <https://doi.org/10.1046/j.1460-9568.2000.00879.x>
- Wagner S, Gresser AL, Torello AT, Dulac C (2006) **A multireceptor genetic approach uncovers an ordered integration of VNO sensory inputs in the accessory olfactory bulb** *Neuron* **50**:697–709 <https://doi.org/10.1016/j.neuron.2006.04.033>

Wang L, Chen IZ, Lin D (2015) **Collateral pathways from the ventromedial hypothalamus mediate defensive behaviors** *Neuron* **85**:1344–1358 <https://doi.org/10.1016/j.neuron.2014.12.025>

Wong WM, Cao J, Zhang X, Doyle WI, Mercado LL, Gautron L, Meeks JP (2020) **Physiology-forward identification of bile acid-sensitive vomeronasal receptors** *Sci Adv* **6** <https://doi.org/10.1126/sciadv.aaz6868>

Yonekura J, Yokoi M (2008) **Conditional genetic labeling of mitral cells of the mouse accessory olfactory bulb to visualize the organization of their apical dendritic tufts** *Mol Cell Neurosci* **37**:708–718 <https://doi.org/10.1016/j.mcn.2007.12.016>

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Reviewer #1 (Public Review):

Summary:

Animals in natural environments need to identify predator-associated cues and respond with the appropriate behavioral response to survive. In rodents, some chemical cues produced by predators (e.g., cat saliva) are detected by chemosensory neurons in the vomeronasal organ (VNO). The VNO transmits predator-associated information to the accessory olfactory bulb, which in turn projects to the medial amygdala and the bed nucleus of the stria terminalis, two regions implicated in the initiation of antipredator defensive behaviors. A downstream area to these two regions is the ventromedial hypothalamus (VMH), which has been shown to control both active (i.e., flight) and passive (i.e., freezing) antipredator defensive responses via distinct efferent projections to the anterior hypothalamic nucleus or the periaqueductal gray, respectively. However, whether differences in predator-associated sensory information initially processed in the VNO and further conveyed to the VMH can trigger different types of behavioral responses remained unexplored. To address this question, here the authors investigated the behavioral responses of mice exposed to either fresh or old cat saliva, and further compared the underlying neural circuits that are activated by cat saliva with different freshness.

The scientific question of the study is valid, the experiments were well-performed, and the statistical analyses are appropriate. However, there are some concerns that may directly affect the main interpretation of the results.

Major Concerns:

1. An important point that the authors should clarify in this study is whether mice are detecting qualitative or quantitative differences between fresh and old cat saliva. Do the environmental conditions in which the old saliva was maintained cause degradation of Fel d 4, the main protein known for inducing a defensive response in rodents? (see Papes et al, 2010 again). If that is the case, one would expect that a lower concentration of Fel d 4 in the old saliva after protein degradation would result in reduced antipredator responses. Alternatively, if the authors believe that different proteins that are absent in the old saliva are contributing to the increased defensive responses observed with the fresh saliva, further protein quantification experiments should be performed. An important experiment to differentiate qualitative versus quantitative differences between the two types of saliva would be diluting the fresh saliva to verify if the amount of protein, rather than the type of protein, is the main factor regulating the behavioral differences.
2. The authors claim that fresh saliva is recognized as an immediate danger by rodents, whereas old saliva is recognized as a trace of danger. However, the study lacks empirical tests to support this interpretation. With the current experimental tests, the behavioral differences between animals exposed to fresh vs. old saliva could be uniquely due to the reduced amount of the exact same protein (e.g., Fel d 4) in the two samples of saliva.
3. In Figure 4H, the authors state that there were no significant differences in the number of cFos-positive cells between the two saliva-exposed groups. However, this result disagrees with the next result section showing that fresh and old saliva differentially activate the VMH.

It is unclear why cFos quantification and behavioral correlations were not performed in other upstream areas that connect the VNO to the VMH (e.g., BNST, MeA, and PMCo). That would provide a better understanding of how brain activity correlates with the different types of behaviors reported with the fresh vs. old saliva.

4. The interpretation that fresh and old saliva activates different subpopulations of neurons in the VMH based on the observation that cFos positively correlates with freezing responses only with the fresh saliva lacks empirical evidence. To address this question, the authors should use two neuronal activity markers to track the response of the same population of VMH cells within the same animals during exposure to fresh vs. old saliva. Alternatively, they could use single-cell electrophysiology or imaging tools to demonstrate that cat saliva of distinct freshness activates different subpopulations of cells in the VMH. Any interpretation without a direct within-subject comparison or the use of cell-type markers would become merely speculative. Furthermore, the authors assume that differential activations of mitral cells between fresh and old saliva result in the differential activation of VMH subpopulations (page 13, line 3). However, there are intermediate structures between the mitral cells and the VMH, which are completely ignored in this study (e.g., BNST, medial amygdala).

5. The authors incorrectly cited the Papes et al., 2010 article on several occasions across the manuscript. In the introduction, the authors cited the Papes et al 2010 study to make reference to the response of rodents to chemical cues, but the Papes et al. study did not use any of the chemical cues listed by the authors (e.g., fox feces, snake skin, cat fur, and cat collars). Instead, the Papes et al. 2010 article used the same chemical cue as the present study: cat saliva. The Papes et al. 2010 article was miscited again in the results section where the authors cited the study to make reference to other sources of cat odor that differ from the cat saliva such as cat fur and cat collars. Because the Papes et al. 2010 article has previously shown the involvement of Trpc2 receptors in the VNO for the detection of cat saliva and the subsequent expression of defensive behaviors by using Trpc2-KO mice, the authors should properly cite this study in the introduction and across the manuscript when making reference to their findings.

6. In the introduction, the authors hypothesized that the VNO detects predator cues and sends sensory signals to the VMH to trigger defensive behavioral decisions and stated that direct evidence to support this hypothesis is still missing. However, the evidence that cat saliva activates the VMH and that activity in the VMH is necessary for the expression of antipredator defensive response in rodents has been previously demonstrated in a study by Engelke et al., 2021 (PMID: 33947849), which was entirely omitted by the authors.

7. In the discussion, the authors stated that their findings suggest that the induction of robust freezing behavior is mediated by a distinct subpopulation of VMH neurons. The authors should cite the study by Kennedy et al., 2020 (PMID: 32939094) that shows the involvement of VMH in the regulation of persistent internal states of fear, which may provide an alternative explanation for why distinct concentrations of saliva could result in different behavioral outcomes.

8. The anatomical connectivity between the olfactory system and the ventromedial hypothalamus (VMH) in the abstract is unclear. The authors should clarify that the VMH does not receive direct inputs from the vomeronasal organ (VNO) nor the accessory olfactory bulb (AOB) as it seems in the current text.

Reviewer #2 (Public Review):

In this study, Nguyen et al. showed that cat saliva can robustly induce freezing behavior in mice. This effect is mediated through the accessory olfactory system as it requires physical contact and is abolished in Trp2 KO mice. The authors further showed that V2R-A4 cluster is

responsive to cat saliva. Lastly, they demonstrated c-Fos induction in AOB and VMHdm/c by the cat saliva. The c-Fos level in the VMHdm/c is correlated with the freezing response.

Strength:

The study opens an interesting direction. It reveals the potential neural circuit for detecting cat saliva and driving defense behavior in mice. The behavior results and the critical role of the accessory olfactory system in detecting cat saliva are clear and convincing.

Weakness:

The findings are relatively preliminary. The identities of the receptor and the ligand in the cat saliva that induces the behavior remain unclear. The identity of VMH cells that are activated by the cat saliva remains unclear. There is a lack of targeted functional manipulation to demonstrate the role of V2R-A4 or VMH cells in the behavioral response to cat saliva.

Reviewer #3 (Public Review):

Summary:

Nguyen et al show data indicating that the vomeronasal organ (VNO) and ventromedial hypothalamus (VMH) are part of a circuit that elicits defensive responses induced by predator odors. They also show that using fresh or old predator saliva may be a method to change the perceived imminence of predation. The authors also identify a family of VNO receptors that are activated by cat saliva. Next, the authors show how different components of this defensive circuit are activated by saliva, as measured by fos expression. Though interesting, the findings are not all integrated into a single narrative, and some of the results are only replications of earlier findings using modern methods. Overall, these findings provide incremental advance.

Strengths:

- 1 Predator saliva is a stimulus of high ethological relevance
- 2 The authors performed a careful quantification of fos induction across the anterior-posterior axis in Figure 6.

Weaknesses:

- 1 It is unclear if fresh and old saliva indeed alter the perceived imminence predation, as claimed by the authors. Prior work indicates that lower imminence induces anxiety-related actions, such as re-organization of meal patterns and avoidance of open spaces, while slightly higher imminence produces freezing. Here, the authors show that fresh and old predator saliva only provoke different amounts of freezing, rather than changing the topography of defensive behaviors, as explained above. Another prediction of predatory imminence theory would be that lower imminence induced by old saliva should produce stronger cortical activation, while fresh saliva would activate the amygdala, if these stimuli indeed correspond to significantly different levels of predation imminence.
- 2 It is known that predator odors activate and require AOB, VNO, and VMH, thus replications of these findings are not novel, decreasing the impact of this work.
- 3 There is a lack of standard circuit dissection methods, such as characterizing the behavioral effects of increasing and decreasing the neural activity of relevant cell bodies and axonal projections, significantly decreasing the mechanistic insights generated by this work.
- 4 The correlation shown in Figure 5c may be spurious. It appears that the correlation is primarily driven by a single point (the green square point near the bottom left corner). All correlations should be calculated using Spearman correlation, which is non-parametric and less likely to show a large correlation due to a small number of outliers. Regardless of the correlation method used, there are too few points in Figure 5c to establish a reliable correlation. Please add more points to 5c.

5 Some of the findings are disconnected from the story. For example, the authors show that V2R-A4-expressing cells are activated by predator odors. Are these cells more likely to be connected to the rest of the predatory defense circuit than other VNO cells?

6 Were there other behavioral differences induced by fresh compared to old saliva? Do they provoke differences in stretch-attend risk evaluation postures, number of approaches, the average distance to odor stimulus, the velocity of movements towards and away from the odor stimulus, etc?