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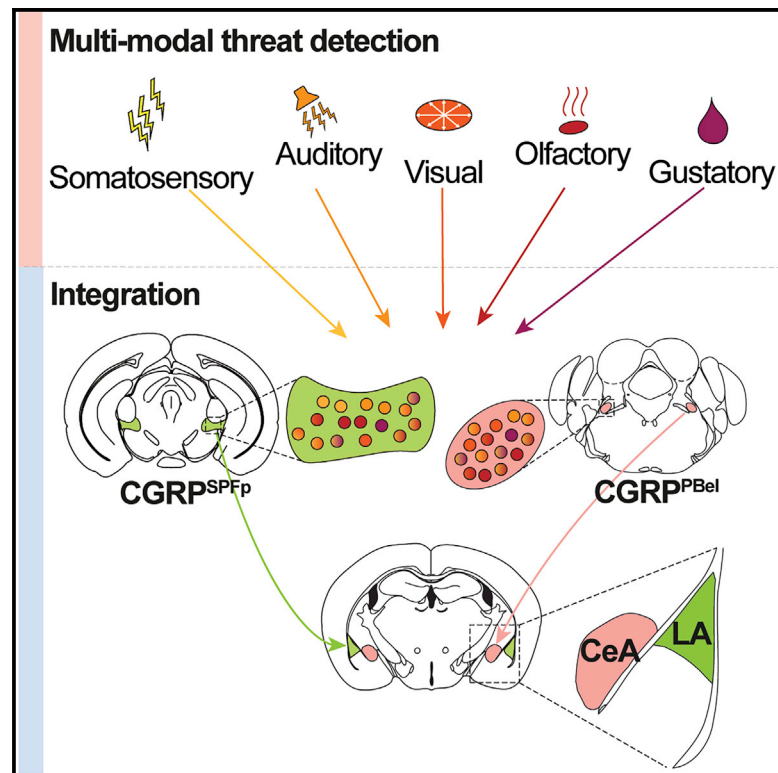
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Authors

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A central alarm system that gates multi-sensory innate threat cues to the amygdala

Graphical abstract



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In brief

Kang et al. report that two CGRP-expressing neuronal populations in the thalamus and brainstem are activated by innate threat cues from various sensory modalities and are required for their perception. These CGRP neurons relay multi-sensory threat cues to the lateral and central amygdala, forming two parallel pathways for aversive memory formation.

Highlights

- $\text{CGRP}^{\text{SPFp}}$ and $\text{CGRP}^{\text{PBel}}$ neurons mediate multi-sensory innate threat perception
- These neurons send aversive information to different subregions of the amygdala
- $\text{CGRP}^{\text{SPFp}} \rightarrow \text{LA}$ and $\text{CGRP}^{\text{PBel}} \rightarrow \text{CeA}$ circuits are crucial for aversive memory formation



Article

A central alarm system that gates multi-sensory innate threat cues to the amygdala

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SUMMARY

Perception of threats is essential for survival. Previous findings suggest that parallel pathways independently relay innate threat signals from different sensory modalities to multiple brain areas, such as the midbrain and hypothalamus, for immediate avoidance. Yet little is known about whether and how multi-sensory innate threat cues are integrated and conveyed from each sensory modality to the amygdala, a critical brain area for threat perception and learning. Here, we report that neurons expressing calcitonin gene-related peptide (CGRP) in the parvocellular subparafascicular nucleus in the thalamus and external lateral parabrachial nucleus in the brainstem respond to multi-sensory threat cues from various sensory modalities and relay negative valence to the lateral and central amygdala, respectively. Both CGRP populations and their amygdala projections are required for multi-sensory threat perception and aversive memory formation. The identification of unified innate threat pathways may provide insights into developing therapeutic candidates for innate fear-related disorders.

INTRODUCTION

To survive in nature, animals must quickly react to danger by detecting aversive information from multiple sensory modalities, such as the shadow of an eagle or the sound of a rattlesnake. Multi-sensory threat stimuli are thought to be detected and processed in parallel by a wide range of relay stations (Silva et al., 2016). To date, separate neural pathways have been reported for processing aversive somatosensory (Barsy et al., 2020; Choi et al., 2020; Han et al., 2015; Sato et al., 2015), visual (Salay et al., 2018; Wei et al., 2015; Zhou et al., 2019), auditory (Barsy et al., 2020; Li et al., 2021; Zhang et al., 2018), gustatory (Carter et al., 2013; Kim et al., 2017; Wang et al., 2018), and olfactory (Rosen et al., 2015; Tong et al., 2020) stimuli and generating immediate defensive behavioral/physiological responses. A common neural substrate that integrates aversive cues from various sensory modalities would be beneficial for survival because essentially all threats in nature exist as multi-modal aversive sensory cues. In addition, it would facilitate efficient information processing and timely initi-

ation of behavioral responses by delivering unified scalable alarm signals to the brain. However, the existence of such pathways has yet to be established.

One candidate brain region for the integration of multi-sensory threat information is the amygdala, a well-known limbic structure crucial for initiating behavioral responses to environmental stimuli and forming associative memories (Janak and Tye, 2015; LeDoux, 2000, 2012). The role of the amygdala in processing aversive information of multiple sensory modalities has been suggested. For example, during canonical Pavlovian threat conditioning, the amygdala is activated by both aversive somatosensory stimuli (e.g., foot shock) and paired auditory cues (Ren and Neugebauer, 2010; Simons et al., 2014; Veinante et al., 2013). Furthermore, lesioning the amygdala attenuates the responsiveness to sensory threats from visual (Bach et al., 2015; Blanchard and Blanchard, 1972; Dal Monte et al., 2015), auditory (Scott et al., 1997), and somatosensory (Blanchard and Blanchard, 1972) modalities, and humans with bilateral damage to this structure exhibit no fear responses to any external threat except CO₂ (Feinstein et al., 2011). Finally, the formation of



aversive memories involving multi-sensory innate threats is amygdala dependent (Blanchard and Blanchard, 1972), supporting the existence of pathways that convey multi-modal aversive sensory cues to the amygdala. However, the neural circuitry that encodes multi-sensory innate aversive cues (other than foot shock) to the amygdala during aversive learning remains uninvestigated.

The amygdala receives dense presynaptic inputs from the posterior thalamus (Craig et al., 1994, 2000; Gauriau and Bernard, 2004; Price, 2002) and the parabrachial nucleus (Usunoff et al., 2006), relay centers that process innate and learned signals of several sensory modalities (e.g., auditory [Yasui et al., 1992], somatosensory [Nakamura and Morrison, 2008], and gustatory [Jarvie et al., 2021]). Subregions of these areas, the parvocellular subparafascicular nucleus (SPFp; also referred to as the posterior thalamus or posterior intralaminar nucleus in the thalamus) (D'Hanis et al., 2007; Doboily et al., 2005; Kruger et al., 1988) and external lateral parabrachial nucleus (PBel), highly express calcitonin gene-related peptide (CGRP), a neuropeptide associated with aversion and nociception (Palmiter, 2018; Russell et al., 2014; Russo, 2015; Shinohara et al., 2017; Yu et al., 2009). In particular, CGRP neurons in the PBel mediate aversive memory formation by transmitting somatosensory and visceral signals to the central amygdala (CeA) (Campos et al., 2018; Chen et al., 2018; Han et al., 2015). CGRP neurons in the medial SPFp are involved in sexual behaviors (Coolen et al., 2003a, 2003b), whereas those in the posterolateral SPFp have been reported to send projections to the lateral amygdala (LA) and striatal amygdala (AStr), implicating that they may be involved in aversive learning (D'Hanis et al., 2007; LeDoux et al., 1985; Yasui et al., 1991). However, the functional role of these neurons in aversive learning has not been investigated.

Given these findings, we hypothesized that CGRP neurons relay multi-sensory innate threat information to the amygdala and that these circuits are involved in generating appropriate behavioral responses and forming aversive memories of innate threat cues.

Here, we report two distinct neural populations in the thalamus and the brainstem, the CGRP^{SPFp} and CGRP^{PBel} neurons, that project to non-overlapping amygdalar subnuclei. These two CGRP populations encode aversive sensory stimuli from various modalities (visual, auditory, somatosensory, gustatory, and olfactory) by recruiting distinct local networks. Silencing CGRP^{SPFp} or CGRP^{PBel} neurons and their terminals in the LA or CeA attenuate the perception of aversive multi-sensory stimuli. Lastly, both the CGRP^{SPFp}→LA and CGRP^{PBel}→CeA circuits are critical for aversive memory formation. The discovery of parallel multi-sensory innate threat pathways supports the critical role of the amygdala in priming behavioral responses to a broad scope of environmental challenges.

RESULTS

Multi-sensory threat stimuli activate CGRP^{SPFp} and CGRP^{PBel} neurons

To investigate how CGRP^{SPFp} and CGRP^{PBel} neurons respond to multi-sensory innate threat stimuli, we recorded the activities of

both populations with miniscope single-cell calcium imaging (Figures 1A and 1B) while presenting multi-sensory innate threat stimuli to freely moving mice (Figure 1C). We expressed jGCaMP8m (Zhang et al., 2020) Cre-dependently in the SPFp or PBel of *Calca*^{Cre} mice (The *Calca* gene encodes CGRP) and implanted a gradient-index (GRIN) lens to facilitate *in vivo* recording of single neuron activities. Mice were presented with aversive threat stimuli of somatosensory (0.2 mA, 2 s foot shock), auditory (85 dB, 2 s sound burst, mimicking thunder blast), visual (2 s expanding looming disk, mimicking the rapid approach of avian predators), olfactory (exposure to cotton tip with 10 μ L of trimethylthiazoline [TMT], a constituent of fox feces), and gustatory modalities (presentation of 10 μ L of 0.5 mM quinine solution to water-restricted mice).

Out of 80 CGRP^{SPFp} (Figure 1D) and 80 CGRP^{PBel} neurons (Figure 1F) recorded, most increased their activity upon presentation of aversive, but not control, stimuli (Figures S2A and S2B), with many neurons responding to more than one sensory modality (Figures 1E and 1G; Tables S1 and S2). Interestingly, we observed different clusters of neurons that preferentially responded to aversive stimuli of each sensory modality. Thus, we sought to further compare the neurons recruited by five modalities using a manifold-based algorithm called convergent cross-mapping (CCM) (Sugihara et al., 2012). Here, CCM is used as a more sophisticated measure than cross-correlation to describe the similarity between the manifolds of the embedded time series of each neuron pair instead of just the cross-correlation of the raw time series, allowing us to compare the neurons recruited across different sensory modalities.

We first calculated the prediction score between each pair of neurons using their peri-stimulus activity traces. A prediction score ρ ($\sum \text{obs}_{(1,2,\dots,n)} / \sum \text{pred}_{(1,2,\dots,n)}$) of 1 indicates that the activity of the input neuron can perfectly predict the future activity of the output neuron. We then performed hierarchical clustering on the prediction score matrix for each of the five stimuli (Figures S2C and S2D, diagonal matrices); clusters represented activated network motifs, since they were composed of neurons that shared the “broadcasters” and “receivers.” However, when we used the same index in the “seed” clustergram to plot the other four heatmaps (Figures S2C and S2D), the clusters were no longer preserved and the heatmaps became dissimilar to each other as quantified by the distance matrix (Figures S2E and S2F), thus indicating that network motifs recruited by each of the five sensory modalities are also distinct. Together, these results suggest that aversive stimuli of all five modalities tested activate CGRP^{SPFp} or CGRP^{PBel} neurons and that each stimulus recruits unique combinations of neurons with distinct patterns of information sharing.

CGRP neurons receive inputs from sensory relay areas

To explore the possible sources of multi-sensory inputs to the CGRP^{SPFp} and the CGRP^{PBel} neurons, we performed the monosynaptic retrograde rabies tracing experiment (Kim et al., 2016) (Figure 2A). Both CGRP^{SPFp} and CGRP^{PBel} neurons received inputs from various sensory relay areas, including the superior colliculus (for visual information), inferior colliculus (for auditory information), vestibular nucleus (for proprioceptive information), spinal dorsal horn (for somatosensory information), trigeminal

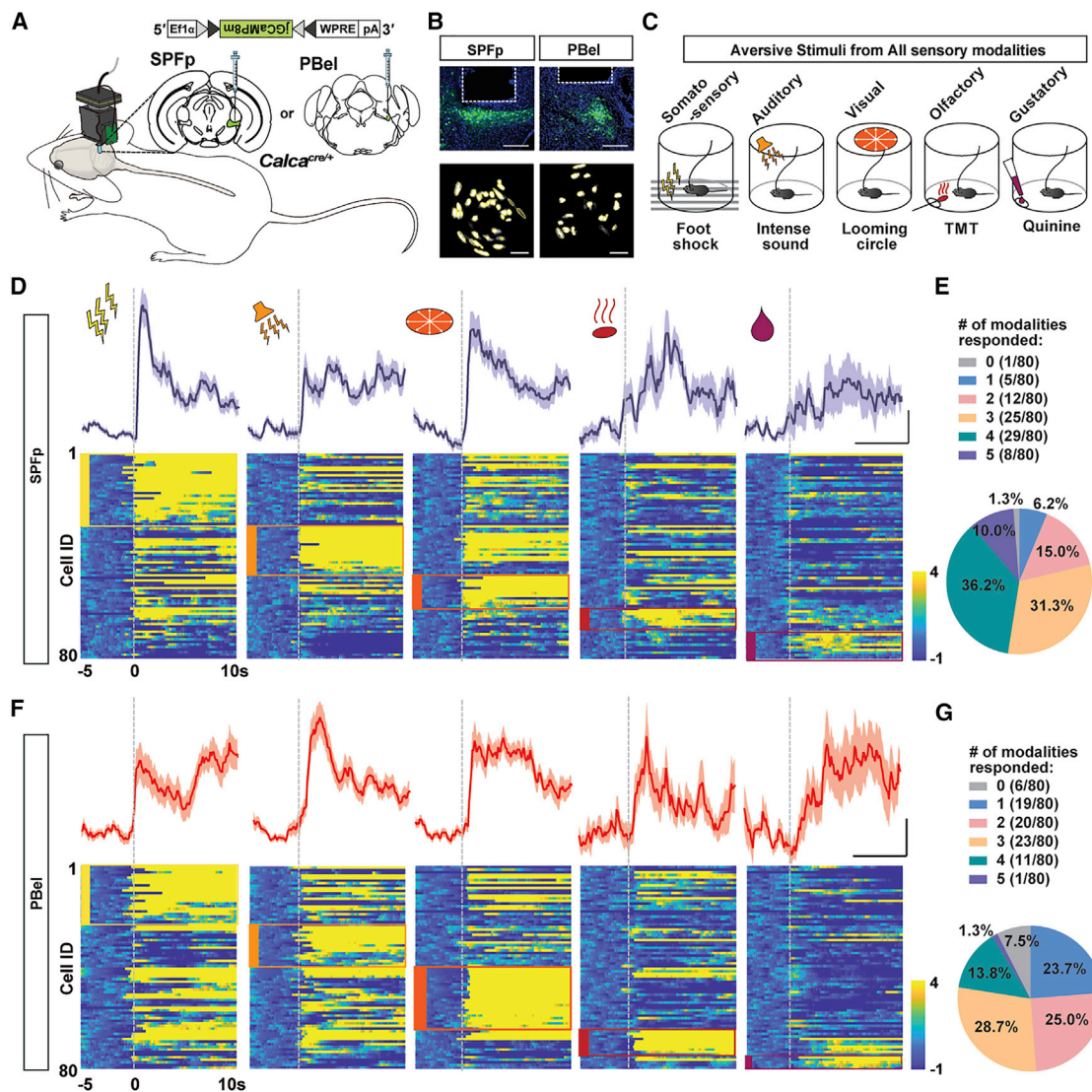


Figure 1. Individual CGRP^{SPFP} and CGRP^{PBel} neurons respond to multi-modal innate threat stimuli

(A–C) Schematic (A), histological verification (B), and behavioral design (C) of single-cell calcium imaging of CGRP^{SPFP} and CGRP^{PBel} neurons with miniature microscope during innate threat stimuli presentation. Scale bars in (B), 200 μ m (top) and 100 μ m (bottom).

(D and F) Bottom: heatmaps of Z-scored CGRP^{SPFP} (D) and CGRP^{PBel} (F) single-cell activity aligned at the onset of stimulus delivery (dotted line) across five different modalities. Clusters of neurons that preferentially respond to certain stimulus types are labeled with boxes of corresponding color. Top: averaged activity from neurons that best respond to each sensory modality. Scale: 5 s/2 Z-scored $\Delta F/F$. n = 80 cells/3 mice in each group.

(E and G) Percentage of CGRP^{SPFP} (E) and CGRP^{PBel} (G) neurons that respond to different numbers of modalities. Data are presented as mean \pm SEM.

See also Figures S1 and S2.

spinal nucleus (for somatosensory information from the face), and the hypothalamus (e.g., zona incerta [ZI] and lateral hypothalamus [LHA] for multi-sensory information) (Figures 2B–2D, S3A, and S3B; Tables S3 and S4). Furthermore, these two CGRP populations displayed different input profiles. While CGRP^{SPFP} neurons received dense inputs from the midbrain (superior and inferior colliculus), CGRP^{PBel} neurons received abundant projections from the amygdala (in particular the CeA), hypothalamus (ZI, LHA, and paraventricular nucleus), and brainstem structures (nucleus tractus solitarius and reticular formation).

These different input profiles suggest that these two CGRP populations may serve different roles in innate threat perception.

CGRP neurons are required for multi-sensory innate threat perception

Since both CGRP^{SPFP} and CGRP^{PBel} neurons can be activated by multi-sensory innate threat stimuli, we next explored whether they are required for multi-sensory threat perception. To this end, we silenced both populations by expressing tetanus toxin light chain (TetTox) bilaterally in the SPFP or PBel of *Calca*^{Cre}

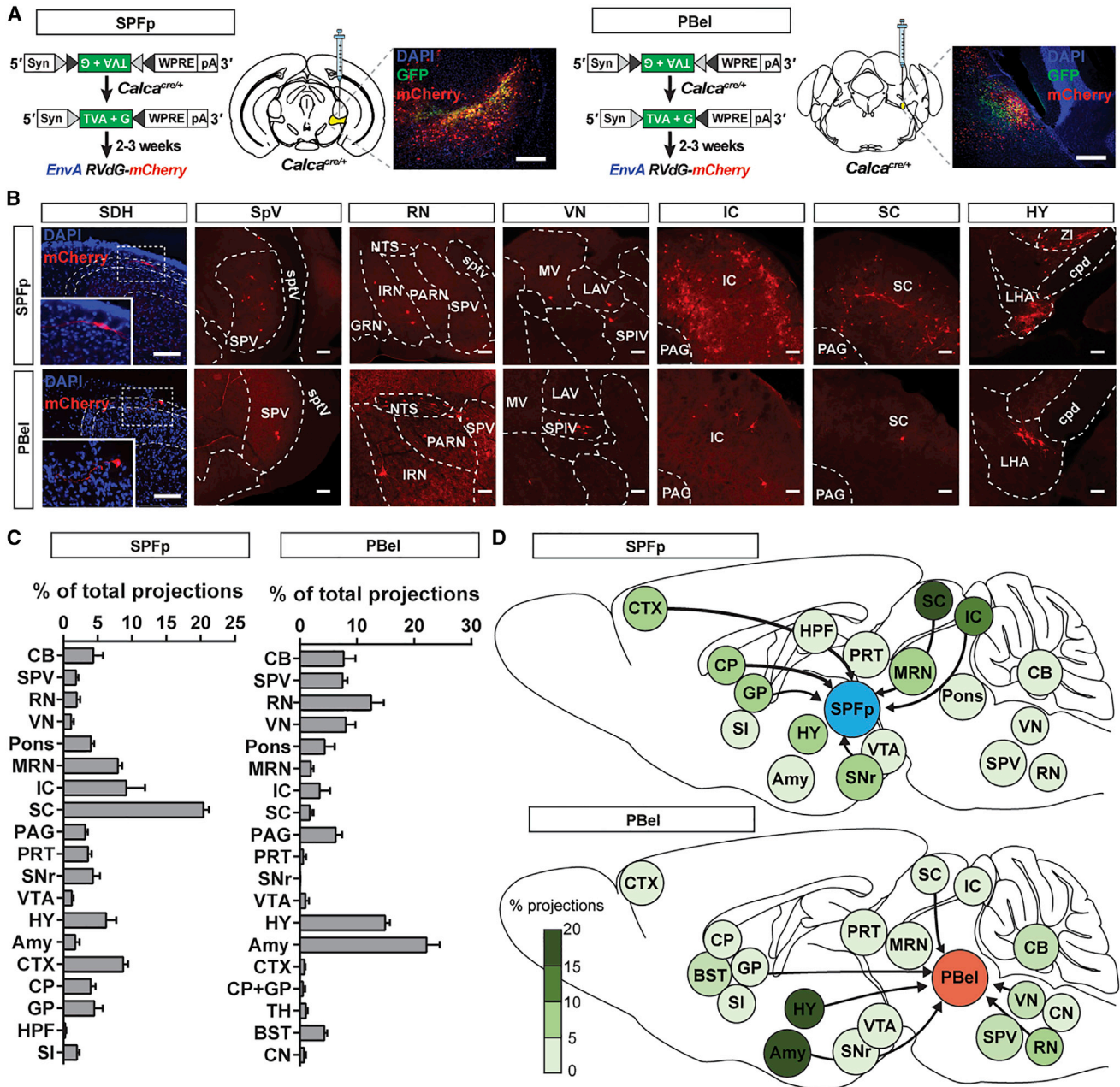


Figure 2. Retrograde tracing from CGRP^{SPFP} and CGRP^{PBel} neurons

(A) Schematic of monosynaptic retrograde rabies tracing experiment and example histology image of the CGRP^{SPFP} (left) and CGRP^{PBel} (right) starter cells. Scale bars, 200 μ m.

(B) Example images of spinal dorsal horn and brain regions that send inputs to CGRP^{SPFP} (top) and CGRP^{PBel} (bottom) neurons. Scale bars, 100 μ m.

(C) Percentage of total projections from each brain region to CGRP^{SPFP} (left) and CGRP^{PBel} (right) neurons. $n = 5$ (CGRP^{SPFP}), $n = 6$ mice (CGRP^{PBel}).

(D) Diagrams of brain-wide inputs to CGRP^{SPFP} (top) and CGRP^{PBel} (bottom) neurons and their relative percentages.

SDH, spinal dorsal horn; CB, cerebellum including interposed nucleus (IP), arbor vitae (arb); SPV, trigeminal spinal nucleus; RN, reticular nucleus including the nucleus tractus solitarius (NTS), intermediate reticular nucleus (IRN), parvocellular reticular nucleus (PARN), gigantocellular reticular nucleus (GRN); VN, vestibular nucleus including medial vestibular nucleus (MV), lateral vestibular nucleus (LAV), spinal vestibular nucleus (SPV); Pons, including the nucleus of the lateral lemniscus, pontine central gray, parabrachial nucleus, pontine reticular nucleus (PRN); MRN, midbrain reticular nucleus; IC, inferior colliculus; SC, superior colliculus; PAG, periaqueductal gray; PRT, pretectal region including anterior pretectal nucleus (APN), medial pretectal area (MPT), nucleus of the posterior commissure (NPC), olivary pretectal nucleus (OP); SNr, substantia nigra; VTA, ventral tegmental area; HY, hypothalamus including lateral hypothalamus (LHA), zona incerta (ZI); Amy, amygdala; CTX, cortex; CP, striatum; GP, globus pallidus; HPF, hippocampus; TH, thalamus; BST, bed nuclei of the stria terminalis; SI, substantia innominata; sptv, spinal tract of the trigeminal nerve; cpd, cerebral peduncle. Data are presented as mean \pm SEM. See also [Figure S3](#); [Tables S2](#) and [S3](#).

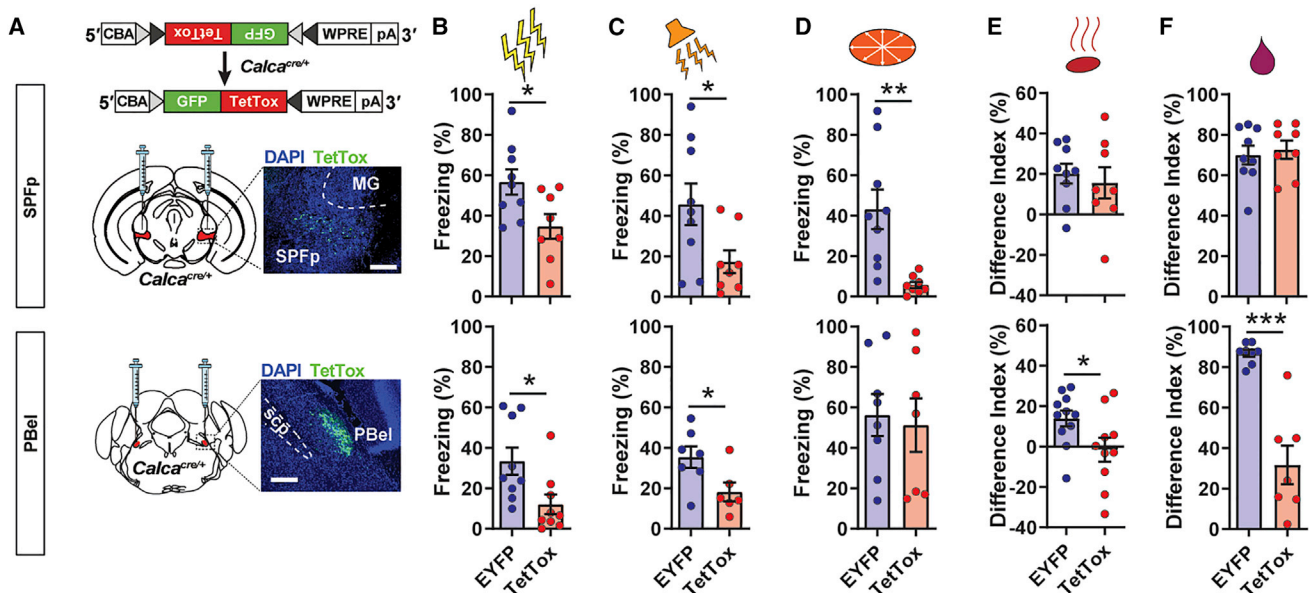


Figure 3. CGRP^{SPFP} or CGRP^{PBel} neurons are required for perception of innate threats of various modalities

(A) Schematic and histological verification of TetTox expression in the CGRP^{SPFP} (top) or CGRP^{PBel} neurons (bottom). Scale bars, 200 μ m.
 (B) Mice expressing TetTox in CGRP^{SPFP} and CGRP^{PBel} neurons displayed less freezing in response to electric foot shock (2 s, 0.6 mA) than EYFP controls. SPFP: n = 9 (EYFP), n = 8 (TetTox) mice; PBel: n = 9 (EYFP), n = 9 (TetTox) mice.
 (C) Mice expressing TetTox in CGRP^{SPFP} and CGRP^{PBel} neurons displayed less freezing in response to intense sound (85 dB) than controls. SPFP: n = 9 (EYFP), n = 8 (TetTox) mice; PBel: n = 7 (EYFP), n = 6 (TetTox) mice.
 (D) Mice expressing TetTox in CGRP^{SPFP} neurons, but not CGRP^{PBel} neurons, displayed less freezing in response to looming stimuli than controls. SPFP: n = 9 (EYFP), n = 8 (TetTox) mice; PBel: n = 8 (EYFP), n = 7 (TetTox) mice.
 (E) The CGRP^{PBel} TetTox group, but not the CGRP^{SPFP} TetTox group, demonstrated reduced avoidance behaviors toward TMT in a two-chamber choice test. SPFP: n = 9 (EYFP), n = 8 (TetTox) mice; PBel: n = 11 (EYFP), n = 10 (TetTox) mice.
 (F) Preference for water over quinine in a two-bottle choice test was intact in the CGRP^{SPFP} TetTox group and greatly reduced in the CGRP^{PBel} TetTox group. SPFP: n = 9 (EYFP), n = 8 (TetTox) mice; PBel: n = 8 (EYFP), n = 7 (TetTox) mice.
 Data are presented as mean \pm SEM. Statistical analyses were performed using two-tailed unpaired t tests (B–F); see also Table S5 for statistical details. *p < 0.05, **p < 0.01, ***p < 0.001.

mice and measured behavioral responses to multi-modal threat stimuli (Figure 3A). Freezing responses to electric foot shock (Figure 3B) and high-intensity sound bursts (Figure 3C) were significantly reduced in both the CGRP^{SPFP} and CGRP^{PBel} TetTox groups compared with the EYFP control groups. While freezing responses to overhead looming stimuli were attenuated in the CGRP^{SPFP} TetTox group compared with controls, the CGRP^{PBel} TetTox group did not differ from controls (Figure 3D).

To assess olfactory threat responses, we next performed a two-chamber preference test in which animals' preference between two chambers containing water and TMT was measured. Silencing CGRP^{SPFP} neurons did not affect animals' avoidance of TMT, whereas this phenotype was abolished in the CGRP^{PBel} TetTox group (Figure 3E). Lastly, gustatory threat response was assessed using a two-bottle choice test in which preference between water and quinine was measured in overnight-water-restricted animals. Quinine aversion was significantly reduced in the CGRP^{PBel} TetTox group but not the CGRP^{SPFP} TetTox group (Figure 3F). Interestingly, although the CGRP^{SPFP} neurons were activated by TMT and quinine (Figure 1D) and CGRP^{PBel} neurons were activated by looming (Figure 1F), inhibition of these populations individually was not

sufficient to attenuate the corresponding threat responses, presumably because of the existence of redundant pathways. These results indicate that CGRP^{SPFP} and CGRP^{PBel} neurons are required for mediating behavioral responses to different sets of multi-sensory threats.

CGRP^{SPFP} and CGRP^{PBel} neurons relay negative valence to the amygdala

The amygdala is implicated in both threat perception and multi-sensory information processing (Bach et al., 2015; Blanchard and Blanchard, 1972; LeDoux, 2000; Ren and Neugebauer, 2010; Scott et al., 1997), so we investigated whether both CGRP populations mediate multi-sensory threat perception via their projections to the amygdala. We first characterized the projections from CGRP^{SPFP} and CGRP^{PBel} neurons to the amygdala by expressing EYFP or mCherry in both populations in the same animals (Figure 4A). Interestingly, EYFP-expressing CGRP^{SPFP} terminals and mCherry-expressing CGRP^{PBel} terminals in the amygdala were mutually exclusive (Figure 4B). While CGRP^{SPFP} synaptic terminals were found in the LA, AStr, and medial amygdala, CGRP^{PBel} terminals were most abundant in the CeA and basomedial amygdala (Figure S3C).

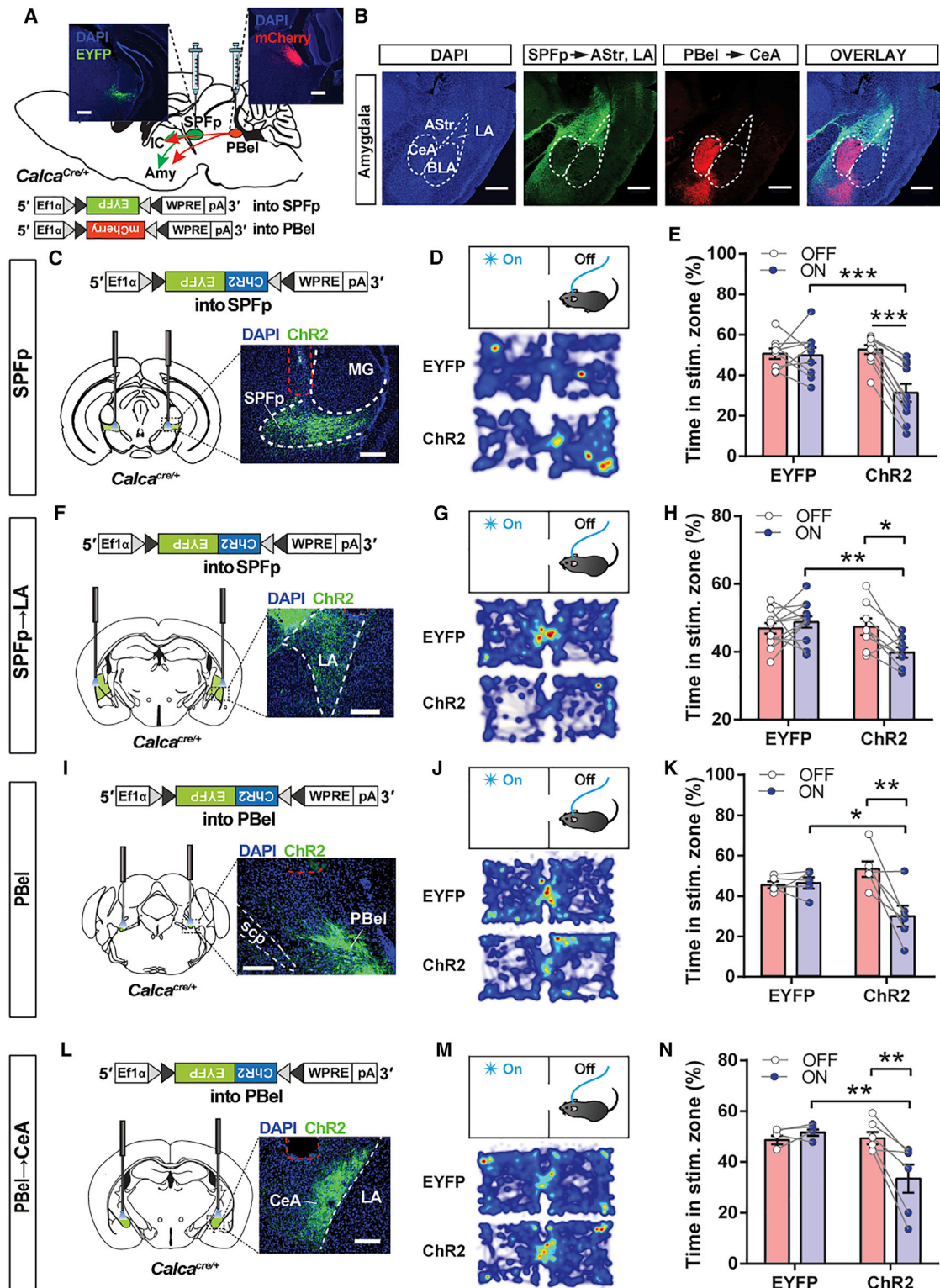


Figure 4. CGRP^{SPFP} and CGRP^{PBel} neurons relay negative valence to the LA and CeA

(A) Schematic and representative images of Cre-dependent expression of EYFP in the SPFP and mCherry in the PBel of a *Calca^{Cre}* mouse. Scale bars, 200 μ m. (B) Axonal projections from the CGRP^{SPFP} and CGRP^{PBel} neurons in the amygdala are mutually exclusive. Scale bars, 500 μ m.

(legend continued on next page)

Non-amygdalar projections from both CGRP populations were also distinct. CGRP^{SPFP} neurons projected to the auditory cortex and dorsal posterior insular cortex (pIC) and did not project to the somatosensory cortex, whereas CGRP^{PBel} neurons projected to the bed nuclei of the stria terminalis, the ventral posteromedial nucleus of the thalamus (parvocellular part), the parasubthalamic nucleus, and the ventral pIC (Figure S3D). Together, these data suggest that the two populations of CGRP neurons are uniquely positioned to anatomically receive aversive sensory stimuli from a wide range of modalities and relay them to the amygdala.

The CGRP^{PBel→CeA} circuit encodes unconditioned noxious stimuli (i.e., foot shock) during aversive learning (Bowen et al., 2020; Han et al., 2015), yet the projections of CGRP^{SPFP} neurons to the amygdala have not been functionally characterized. Since CGRP^{SPFP} neurons project to the LA (Figure 4B), a key area of neural plasticity during threat learning (LeDoux, 2000), we sought to characterize the functional connectivity of this circuit and some other regions (Astr and pIC) via slice electrophysiology and optogenetics. We first expressed channelrhodopsin-2 (ChR2) in the CGRP^{SPFP} neurons and performed electrophysiology recording in the Astr, pIC, and LA (Figures S4A and S4B). We observed optogenetically evoked excitatory (EPSCs) and inhibitory postsynaptic currents (IPSCs) in Astr, pIC, and LA neurons (Figure S4C; pIC trace not shown). Both currents were blocked by the AMPA receptor antagonist CNQX, suggesting the existence of glutamatergic synapses (Figure S4C). Moreover, while most synapses contained both excitatory and inhibitory components (Figure S4D), the onset of IPSCs lagged 4–5 ms relative to EPSCs, indicating the existence of a feedforward inhibitory circuit (Figures S4E–S4G). The amplitudes of optogenetically evoked EPSCs (Figure S4H) or IPSCs (Figure S4I) in the three regions were similar.

Next, we tested whether CGRP^{SPFP} neurons encode negative valence and relay it to the downstream regions, including the LA, using a real-time place aversion (RTPA) test. Mice were placed in a two-chamber arena, one side of which was paired with photostimulation of CGRP^{SPFP} cell bodies (Figure 4C) or CGRP^{SPFP} terminals, including the CGRP^{SPFP→LA} projection (Figures 4F and S4J). In both experiments, the ChR2 group, but not EYFP controls, avoided the photostimulation-paired chamber (Figures 4D, 4E, 4G, 4H, S4J, S5A, and S5B). We used a similar RTPA paradigm to investigate whether CGRP^{PBel} neurons

similarly encode negative valence and transmit this information to the CeA. Photostimulation of CGRP^{PBel} cell bodies (Figures 4I–4K and S5C) and CGRP^{PBel→CeA} terminals (Figures 4L–4N and S5D) both induced aversion to the light-paired chamber. Together, these results suggest that CGRP^{SPFP} and CGRP^{PBel} neurons relay negative valence to non-overlapping amygdala subnuclei.

CGRP^{SPFP→LA} and CGRP^{PBel→CeA} projections are required for multi-sensory threat perception

Given that both CGRP neuronal populations, together with their projections to the amygdala, are required for multi-sensory innate threat responses, we next investigated whether CGRP^{SPFP→LA} and CGRP^{PBel→CeA} circuits are also necessary for generating threat responses to multi-sensory aversive signals. We used, respectively, a photoswitchable Gi-coupled opsin that directly inhibits transmitter release, parainopsin (PPO) (Copits et al., 2021) (Figure 5A), and a light-driven outward proton pump, archaerhodopsin (ArchT) (Figure 5F), to optogenetically inhibit the axon terminals of CGRP^{SPFP→LA} and CGRP^{PBel→CeA} projections while presenting the same threat stimuli used in the cell-body inhibition experiment.

Silencing the CGRP^{SPFP→LA} pathway attenuated freezing responses to auditory (Figure 5B) and visual (Figure 5C) stimuli compared with controls but did not alter responses to olfactory (Figure 5D) or gustatory (Figure 5E) stimuli. Meanwhile, silencing the CGRP^{PBel→CeA} pathway attenuated defensive behavior following the presentation of auditory (Figure 5G), olfactory (Figure 5I), and gustatory (Figure 5J) stimuli, but not visual stimuli (Figure 5H). The somatosensory stimulus (electric foot shock) was excluded here because it will be discussed in the following section. These results are consistent with the cell-body silencing experiment (Figure 3), indicating the critical roles of both CGRP^{PBel→CeA} and CGRP^{SPFP→LA} circuits in generating innate responses to multi-sensory threat cues.

CGRP^{SPFP→LA} and CGRP^{PBel→CeA} projections are required for threat memory formation but not expression

Since both CGRP^{SPFP→LA} and CGRP^{PBel→CeA} circuits are critical for generating behavioral responses to multi-sensory threat signals, we next tested whether they are similarly important for threat memory formation. We used miniscope single-cell

(C and D) Viral expression schematic, histological verification (C), and heatmap (D) of real-time place aversion (RTPA) experiment with CGRP^{SPFP} cell-body photostimulation. Scale bars, 200 μ m.

(E) ChR2-expressing mice, but not EYFP controls, avoided the chamber paired with CGRP^{SPFP} cell-body photostimulation. $n = 9$ (EYFP), $n = 10$ (ChR2) mice. Data indicate the percentage of time spent in the photostimulated zone during the baseline (OFF, 0–10 min) and the last 10 min of photostimulation period (ON, 20–30 min).

(F and G) Viral expression schematic, histological verification (F), and heatmap (G) of CGRP^{SPFP→LA} terminal photostimulation for RTPA experiment. Scale bars, 200 μ m.

(H) ChR2-expressing mice, but not EYFP controls, avoided the chamber paired with CGRP^{SPFP→LA} terminal photostimulation. $n = 12$ (EYFP), $n = 8$ (ChR2) mice.

(I and J) Viral expression schematic, histological verification (I), and heatmap of RTPA experiment (J) with CGRP^{PBel} cell-body photostimulation. Scale bars, 200 μ m.

(K) ChR2-expressing mice, but not EYFP controls, avoided the chamber paired with CGRP^{PBel} cell-body photostimulation. $n = 5$ (EYFP), $n = 6$ (ChR2) mice.

(L and M) Viral expression schematic, histological verification (L), and heatmap of RTPA experiment (M) with CGRP^{PBel→CeA} terminal photostimulation. Scale bars, 200 μ m.

(N) ChR2-expressing mice, but not EYFP controls, avoided the chamber paired with CGRP^{PBel→CeA} terminal photostimulation. $n = 5$ (EYFP), $n = 6$ (ChR2) mice. Data are presented as mean \pm SEM. Statistical analyses were performed using repeated-measures two-way ANOVA with post hoc Sidak's multiple comparison; see also Table S5 for statistical details. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. See also Figures S5 and S6.

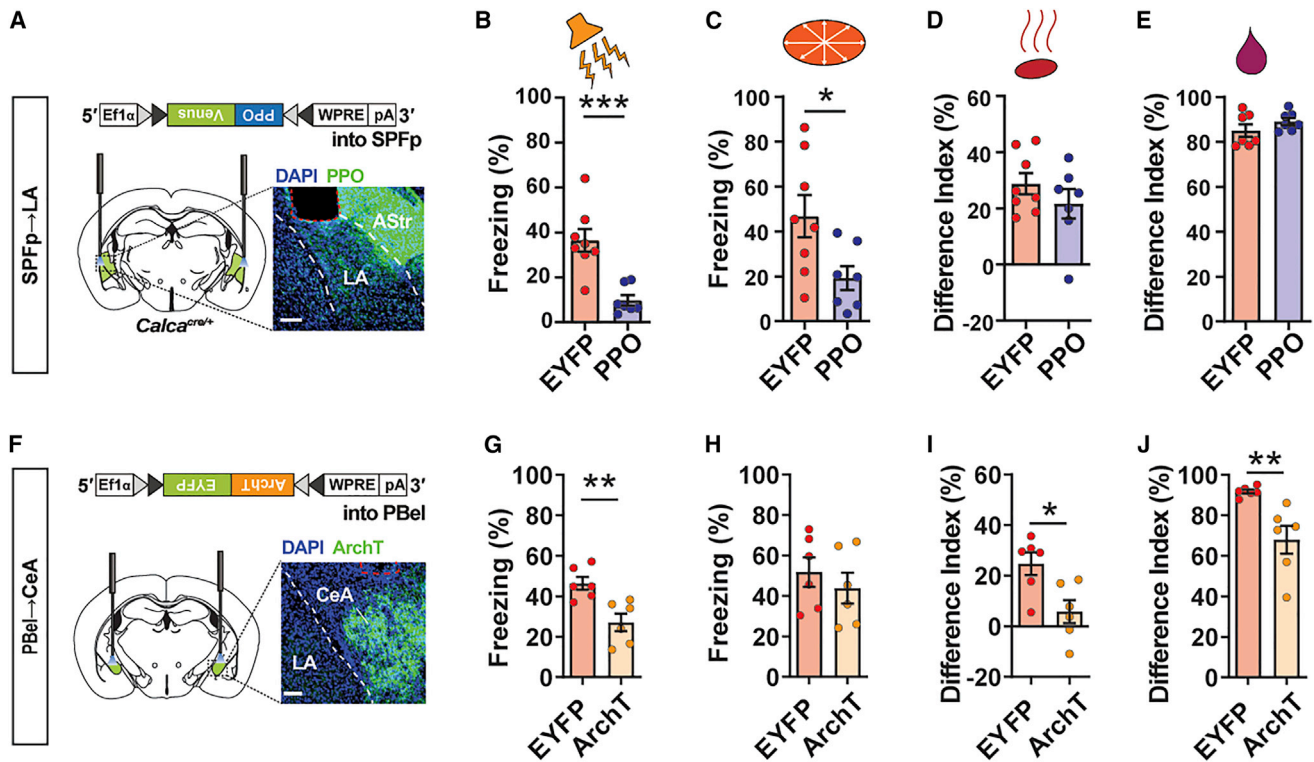


Figure 5. CGRP^{SPFP→LA} and CGRP^{PBel→CeA} projections are required for multi-sensory innate threat perception

(A) Schematic for silencing CGRP^{SPFP→LA} circuit with PPO. Scale bars, 200 μm.
 (B–E) Silencing the CGRP^{SPFP→LA} circuit reduced behavioral responses to aversive auditory (B) and visual (C) stimuli, but not olfactory (D) or gustatory (E) stimuli. Auditory, visual, and olfactory: n = 8 (EYFP), n = 7 (PPO) mice; gustatory: n = 7 (EYFP), n = 7 (PPO) mice.
 (F) Schematic for silencing CGRP^{PBel→CeA} circuit with ArchT. Scale bars, 200 μm.
 (G–J) Silencing the CGRP^{PBel→CeA} circuit reduced behavioral responses to aversive auditory (G), olfactory (I), and gustatory (J) stimuli, but not visual (H) stimuli. n = 6 (EYFP), n = 6 (ArchT) mice.
 Data are presented as mean ± SEM. Statistical analyses were performed using two-tailed unpaired t tests; see also Table S5 for statistical details. *p < 0.05, **p < 0.01, ***p < 0.001.

calcium imaging to quantify the recruitment of CGRP^{SPFP} neurons during a traditional Pavlovian threat-conditioning paradigm that coupled auditory cues with foot shock. Animals displayed minimal freezing during habituation (Figure S6A), but prominent freezing behavior during both the conditioning phase and the cue-dependent retrieval test (Figures S6B and S6C). CGRP^{SPFP} neurons were activated by foot shock (Figure 6A, middle panel) but not by tones during habituation or the cue test (Figures 6A and 6B). These data suggest that CGRP^{SPFP} neurons encode somatosensory threats (unconditioned stimulus [US]) during threat conditioning but do not play a role in their association with conditioned stimuli.

We next performed a projection-specific optogenetic conditioning test wherein CGRP^{SPFP} terminal photostimulation, including the CGRP^{SPFP→LA} projection, was used as the US in lieu of foot shock. In the context-dependent retrieval test, the CGRP^{SPFP→LA} ChR2 group exhibited more freezing than the control while the CGRP^{SPFP→Astr} or CGRP^{SPFP→pIC} ChR2 group did not (Figures 6C, S4K, and S7A). All the projection ChR2 groups formed memory in the cue-dependent test, but only the CGRP^{SPFP→LA} and CGRP^{SPFP→pIC} ChR2 group froze more than the control during the tone exposure. Moreover, the

CGRP^{SPFP→LA} ChR2 group exhibited more freezing than the CGRP^{SPFP→Astr} ChR2 group (Figures 6D and S4L). These results indicate that CGRP^{SPFP→LA} activation was sufficiently aversive to act as the US and induce a fear memory rather than stimulating the other projection regions. Thus, we did not further pursue Astr and pIC in the terminal inhibition experiments. Lastly, we silenced the CGRP^{SPFP→LA} pathway with PPO during the conditioning phase of the traditional threat-conditioning paradigm. Both groups showed similar aversive behavior (jumping and freezing) during the conditioning (data not shown), indicating that inhibition of the amygdalar projection was not able to attenuate the immediate response to the strong somatosensory stimuli. However, compared with controls, PPO-expressing animals displayed reduced freezing behavior in both the context- and cue-dependent fear retrieval tests (Figures 6E, 6F, and S7B). When silencing the same circuit only during the retrieval sessions after threat conditioning, no reductions of freezing were observed (Figures S7E–S7G), indicating that these circuits are necessary for relaying US information but not for memory retrieval.

CGRP^{PBel} neurons were similarly activated by only the US during threat conditioning (Figures 6G, 6H, and S6D–S6F). Some

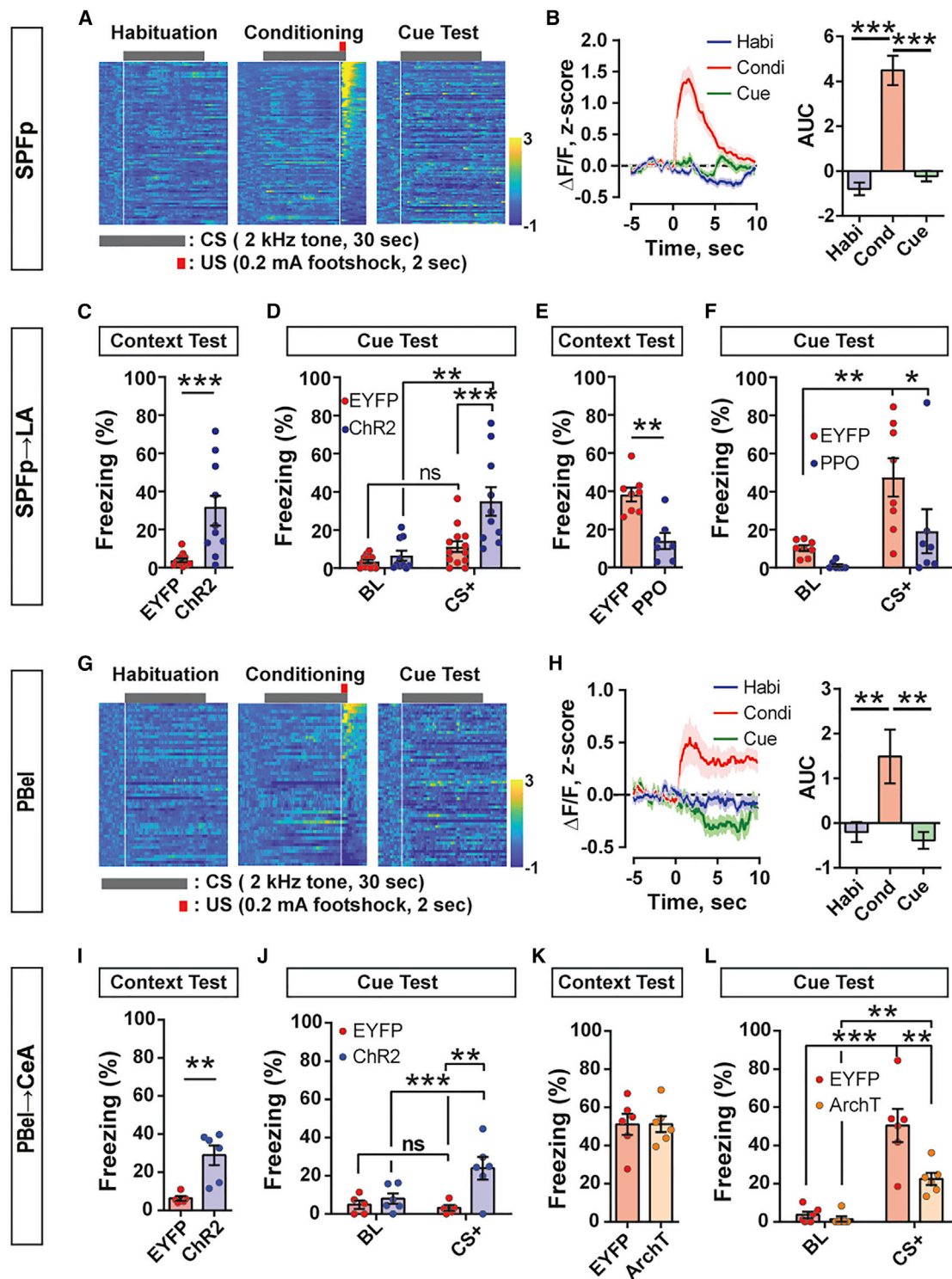


Figure 6. $\text{CGRP}^{\text{SPFP} \rightarrow \text{LA}}$ and $\text{CGRP}^{\text{PBel} \rightarrow \text{CeA}}$ circuits relay unconditioned stimulus information and are crucial for aversive memory formation during Pavlovian threat conditioning

(A) Heatmap of Z-scored $\text{CGRP}^{\text{SPFP}}$ single-cell calcium activity during threat conditioning. $n = 81$ (habituation), $n = 90$ (conditioning), and $n = 93$ cells (cue test) from three mice. Each row represents the averaged value during three to six stimulus presentations in a single cell. White vertical lines indicate the onsets of tone or foot shock used for (B).

(legend continued on next page)

neurons from the same field of view displayed similar activity patterns but were confirmed to be spatially segregated (Figures S6G–S6J). CGRP^{PBeI→CeA} photostimulation was likewise sufficient to generate an aversive memory during optogenetic conditioning, increasing freezing behavior in both the context (Figures 6I and S7C) and cue retrieval tests (Figure 6J). Inhibiting the CGRP^{PBeI→CeA} projection during the conditioning phase of the traditional threat-conditioning paradigm with ArchT reduced freezing behavior during the cue test (Figure 6L), but not the context test (Figures 6K and S7D). Moreover, inhibiting this circuit only during the retrieval tests did not alter freezing responses (Figures S7H–S7J), thus indicating that it is not required for threat memory expression.

Collectively, these data suggest that both CGRP^{SPFP→LA} and CGRP^{PBeI→CeA} circuits are crucial for forming aversive memories by relaying US information.

DISCUSSION

Regardless of sensory modality, it is critical that threat stimuli be processed promptly and reliably to elicit appropriate behavioral responses. The present study characterized two parallel circuits that encode multi-sensory innate threat signals and are required for innate defensive behaviors. These two circuits comprise thalamic (SPFP) and brainstem (PBeI) CGRP neurons that project to complementary subdivisions of the amygdala, a well-known center for generating threat memories and diverse defensive responses. CGRP^{SPFP} neurons send massive projections to AStr and LA, making it difficult to manipulate just one region. However, we have performed experiments with different optic fiber positions (Figures S4M and S4N), allowing us to confidently claim the role of the CGRP^{SPFP→LA} circuit in our study.

Both CGRP^{SPFP} and CGRP^{PBeI} neurons are activated by threat signals of all five modalities tested (Figure 1). Interestingly, different subpopulations of neurons are preferentially recruited by each modality, possibly encoding stimulus identity (Figures 1D, 1F, and S2C–S2F). Although both CGRP^{SPFP→LA} and CGRP^{PBeI→CeA} circuits are necessary for processing somatosensory and auditory threat cues, aversive visual stimuli

require only the former pathway for processing while olfactory and gustatory stimuli require only the latter. This specificity may originate from distinct input patterns (Figure 2). CGRP^{SPFP} neurons receive densest inputs from midbrain visual and auditory centers (i.e., the superior and inferior colliculi [Silva et al., 2016; Zhou et al., 2019]), while CGRP^{PBeI} neurons receive inputs from the nucleus tractus solitarius, a key player in gustatory and olfactory processing (Garcia-Diaz et al., 1988). Additionally, terminal optogenetic inhibition did not influence the immediate defensive responses by foot shock. Based on these distinctions, we speculate that the CGRP^{SPFP→LA} circuit is more specialized for processing exteroceptive signals while the CGRP^{PBeI→CeA} circuit is more specialized for processing interoceptive stimuli. Moreover, we contemplate that both circuits are necessary for aversive memory formation, but they have limited effects in immediate escape responses associated with bursts of locomotor activity by intense stimuli such as jumping and running. Other parallel circuits may be mediating those immediate responses by severe stimuli (Chiang et al., 2020; Silva et al., 2016). Another interesting possibility we have yet to rule out is that both circuits could relay appetitive information to the amygdala in addition to aversive information.

A shared neural substrate for encoding multi-modal aversive sensory signals would confer many advantages for survival. First, given that predators often generate stimuli of multiple sensory modalities simultaneously, the convergence of these threat signals onto a single brain area could facilitate decision-making and the initiation of appropriate responses. CGRP neurons from the SPFP and PBeI may serve as parallel conduits of sensory information from various upstream regions to the amygdala, which can then mediate the behavioral and physiological components of the fear responses via downstream targets (e.g., periaqueductal gray [Rizvi et al., 1991] or hypothalamus [Viviani et al., 2011]). Second, a shared circuit for detecting multi-sensory innate threats could be integrated into the network for aversive memory formation. Here, we demonstrated that: (1) the CGRP^{SPFP→LA} and CGRP^{PBeI→CeA} circuits are necessary for forming threat memories in response to foot shock during Pavlovian threat learning; (2) the same pathways that process

(B) Left: averaged peri-stimulus activity of all recorded CGRP^{SPFP} neurons. Stimulus onset is indicated by white vertical lines in (A). Right: area under the curve (AUC) of all recorded CGRP^{SPFP} neurons. AUCs were calculated from 0 to 5 s after stimulus onset. CGRP^{SPFP} neurons were activated by the unconditioned stimulus (foot shock) and not by the conditioned stimulus (non-aversive tone) during habituation or cue tests. $n = 81$ (habituation), $n = 90$ (conditioning), and $n = 93$ cells (cue test) from three mice.

(C and D) During the optogenetic threat-conditioning test wherein CGRP^{SPFP→LA} circuit photostimulation was used as an unconditioned stimulus (US), the ChR2 group displayed higher freezing levels during both the context (C) and cue (D) retrieval tests than EYFP controls. $n = 13$ (EYFP), $n = 10$ (ChR2) mice. “BL” in the cue test indicates freezing during baseline before the first tone was delivered, and “CS+” is the average freezing during the three tones.

(E and F) Inhibiting the CGRP^{SPFP→LA} circuit with PPO during threat conditioning reduced freezing level in the context (E) and cue retrieval tests (F). $n = 8$ (EYFP), $n = 7$ (PPO) mice.

(G) Heatmap of Z-scored CGRP^{PBeI} single-cell calcium activity during threat conditioning. $n = 56$ (habituation), $n = 50$ (conditioning), and $n = 59$ cells (cue test) from three mice. White vertical lines indicate the onsets of tone or foot shock used for (H).

(H) Same design as in (B). CGRP^{PBeI} neurons were activated by the unconditioned stimulus (foot shock), but not by the cues during either habituation or cue tests. $n = 56$ (habituation), $n = 50$ (conditioning), and $n = 59$ cells (cue test) from three mice.

(I and J) During the optogenetic threat-conditioning test wherein CGRP^{PBeI→CeA} circuit photostimulation was used as a US, the ChR2 group displayed higher freezing levels during both the context (I) and cue (J) retrieval tests than EYFP controls. $n = 5$ (EYFP), $n = 6$ (ChR2) mice.

(K and L) Inhibiting the CGRP^{PBeI→CeA} circuit with ArchT during threat conditioning reduced freezing behavior in the cue retrieval test (L) but not the context test (K). $n = 6$ (EYFP), $n = 6$ (ArchT) mice.

Data are presented as mean \pm SEM. Statistical analyses were performed using Kruskal-Wallis one-way ANOVA on ranks with Dunn’s multiple comparisons (B and H), two-tailed unpaired t tests (C, E, I, and K), or repeated-measures two-way ANOVA with post hoc Sidak’s multiple comparison (D, F, J, and L); see also Table S5 for statistical details. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. See also Figures S7–S9.

foot-shock cues relay innate threat cues from other sensory modalities; (3) stimulation of either of these pathways is sufficient to act as a US. These same pathways are therefore likely recruited when forming memories of innate sensory threats. Our findings complement the current understanding of innate threat perception circuitry whereby threat cues from each sensory modality are processed in parallel by different brain areas (Canteras, 2002; Gross and Canteras, 2012; Kunwar et al., 2015; Silva et al., 2013) independently from the Pavlovian threat learning circuit (Silva et al., 2016). Finally, a common neural substrate for multiple threat signals could be an entry point for coordinating threat responses with other behaviors. For example, the amygdala could be modulated by various inputs (Fu et al., 2020) and allow threat signals to be collectively regulated with other behavior demands, e.g., foraging and parental drives (Comeras et al., 2019; Ferreira et al., 2002). However, our terminal inhibition of the CGRP^{PBel→CeA} circuit was not able to attenuate the freezing behavior during the context test of the threat-conditioning experiment. This phenomenon could be due to the intricacies of the contextual and cued threat-conditioning paradigms. Previous studies claim that the contextual fear conditioning requires both hippocampus and amygdala, while cued threat conditioning is amygdala dependent but hippocampus independent (Kim and Fanselow, 1992; Nagy et al., 2006). Therefore, terminal inhibition of the CGRP^{PBel→CeA} circuit may not affect contextual freezing.

Our study suggests that the neuropeptide CGRP may be a marker for circuits that process multi-sensory innate threat stimuli. Importantly, it has also been shown that CGRP exerts functional roles in the amygdala. For example, infusing CGRP receptor antagonist into the CeA blocks Pavlovian auditory fear conditioning (Kocorowski and Helmstetter, 2001). Future developments of *in vivo* CGRP release monitoring approaches are needed to unveil the contributions of CGRP in the circuits reported here. In humans, the PBel and SPFP also abundantly express CGRP (Lacalle and Saper, 2000), suggesting that the dual circuits we reported in mice may also be involved in threat perception-related psychiatric disorders. For example, the CGRP signaling pathway is a proven therapeutic target for migraines (Ashina, 2020), which involve hypersensitivity toward neutral stimuli (Demarquay and Mauguière, 2016; Harriott and Schwedt, 2014). It is, therefore, likely that the CGRP network, including the two populations reported in this study, are also hyperactive during migraine-triggered sensory hypersensitivity episodes. CGRP receptor antagonists or neutralizing monoclonal antibodies currently used for migraines may also be useful for treating other disorders involving multi-sensory stimuli processing abnormalities, including autism (Nelson et al., 2001), post-traumatic stress disorder (Ashina et al., 2020), and chronic pain disorders such as fibromyalgia (Harte et al., 2016; López-Solà et al., 2017). Examining the roles of CGRP signaling in the CGRP^{SPFP→LA} and CGRP^{PBel→CeA} circuits in mediating migraine-related symptoms would be an exciting direction for further research.

Limitations of the study

We would like to point out several aspects of our current study that are worth further investigation. First, results from our study are at discrepancy with a previous study regarding the CGRP^{PBel} neurons' role in the cue retrieval test (Campos et al.,

2018). While our results show that no net calcium activity changes in CGRP^{PBel} neurons were observed during cue retrieval, Campos et al. showed increased CGRP^{PBel} activity in response to the conditioned stimulus during the memory retrieval test, which were decreased as the animal was repeatedly exposed to the tone. We believe that this discrepancy could be due to the intensity of the tone, since our study used a low-intensity conditioned stimulus (73 dB tone) that does not activate CGRP^{PBel} neurons during conditioning. However, we would need to perform a fear conditioning experiment with a higher-intensity auditory conditioned stimulus cue to draw this conclusion. Although we showed that the CGRP^{PBel} neuronal activation is not necessary for memory retrieval (Figures S7I and S7J), future studies will be needed to evaluate whether different conditioned stimulus intensities contribute differentially to the role of CGRP^{PBel} neurons in associative threat learning. Second, we did not identify specific cell types of the LA or CeA neurons that receive inputs from the SPFP or PBel in these circuits. The neurons responsible for these functions in the CeA express CGRP receptor (Han et al., 2015), but their counterparts in the LA have not been characterized, which remains an open question for follow-up studies. Finally, our study shows that the CGRP^{SPFP→LA} and CGRP^{PBel→CeA} circuits serve the parallel US pathways in foot-shock-induced classical threat learning. Considering that these parallel circuits relay multi-sensory threat signals to the amygdala, it is plausible that they play similar roles in forming aversive memories associated with other sensory threat cues. However, this idea was not tested in the current study and requires further investigation.

STAR★METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.celrep.2022.111222>.

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AUTHOR CONTRIBUTIONS

S.H. conceived of the idea and secured funding. S.H., S.J.K., S.L., and M.Y. designed the experiments. S.L., S.H., and S.J.K. wrote the manuscript. S.J.K. performed most CGRP^{SPFP} experiments. S.L. and S.J.K. performed most CGRP^{PBeI} experiments. S.L. set up the miniscope experiment and analyzed the data. M.Y. performed electrophysiology and CGRP^{PBeI} RTPA. D.-I.K. provided viruses. G.M.P. performed the CCM analyses. K.-F.L. provided resources for microscopy. M.R.B. and B.A.C. provided the PPO virus. B.Z.R. edited the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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REFERENCES

Ashina, M. (2020). Migraine. *N. Engl. J. Med.* *383*, 1866–1876.

Ashina, H., Iljazi, A., Al-Khazali, H.M., Eigenbrodt, A.K., Larsen, E.L., Andersen, A.M., Hansen, K.J., Bräuner, K.B., Mørch-Jessen, T., Chaudhry, B., et al. (2020). Efficacy, tolerability, and safety of erenumab for the preventive treatment of persistent post-traumatic headache attributed to mild traumatic brain injury: an open-label study. *J. Headache Pain* *21*, 62.

Bach, D.R., Hurlmann, R., and Dolan, R.J. (2015). Impaired threat prioritisation after selective bilateral amygdala lesions. *Cortex* *63*, 206–213.

Barsy, B., Kocsis, K., Magyar, A., Babiczky, Á., Szabó, M., Veres, J.M., Hillier, D., Ulbert, I., Yizhar, O., and Máttyás, F. (2020). Associative and plastic thalamic signaling to the lateral amygdala controls fear behavior. *Nat. Neurosci.* *23*, 625–637.

Blanchard, D.C., and Blanchard, R.J. (1972). Innate and conditioned reactions to threat in rats with amygdaloid lesions. *J. Comp. Physiol. Psychol.* *81*, 281–290.

Bowen, A.J., Chen, J.Y., Huang, Y.W., Baertsch, N.A., Park, S., and Palmiter, R.D. (2020). Dissociable control of unconditioned responses and associative fear learning by parabrachial CGRP neurons. *Elife* *9*, e59799.

Campos, C.A., Bowen, A.J., Roman, C.W., and Palmiter, R.D. (2018). Encoding of danger by parabrachial CGRP neurons. *Nature* *555*, 617–622.

Canteras, N.S. (2002). The medial hypothalamic defensive system: hodological organization and functional implications. *Pharmacol. Biochem. Behav.* *71*, 481–491.

Carter, M.E., Soden, M.E., Zweifel, L.S., and Palmiter, R.D. (2013). Genetic identification of a neural circuit that suppresses appetite. *Nature* *503*, 111–114.

Carter, M.E., Han, S., and Palmiter, R.D. (2015). Parabrachial calcitonin gene-related peptide neurons mediate conditioned taste aversion. *J. Neurosci.* *35*, 4582–4586.

Chen, J.Y., Campos, C.A., Jarvie, B.C., and Palmiter, R.D. (2018). Parabrachial CGRP neurons establish and sustain aversive taste memories. *Neuron* *100*, 891–899.e5.

Chiang, M.C., Nguyen, E.K., Canto-Bustos, M., Papale, A.E., Oswald, A.-M.M., and Ross, S.E. (2020). Divergent neural pathways emanating from the

lateral parabrachial nucleus mediate distinct components of the pain response. *Neuron* *106*, 927–939.e5.

Choi, S., Hachisuka, J., and Ginty, D. (2020). Parallel ascending spinal pathways for affective touch and pain. *Nature* *587*, 258–263.

Comeras, L.B., Herzog, H., and Tasan, R.O. (2019). Neuropeptides at the crossroad of fear and hunger: a special focus on neuropeptide Y. *Ann. N. Y. Acad. Sci.* *1455*, 59–80.

Coolen, L.M., Veening, J.G., Wells, A.B., and Shipley, M.T. (2003a). Afferent connections of the parvocellular subparafascicular thalamic nucleus in the rat: evidence for functional subdivisions. *J. Comp. Neurol.* *463*, 132–156.

Coolen, L.M., Veening, J.G., Petersen, D.W., and Shipley, M.T. (2003b). Parvocellular subparafascicular thalamic nucleus in the rat: anatomical and functional compartmentalization. *J. Comp. Neurol.* *463*, 117–131.

Copits, B.A., Gowrishankar, R., O’Neill, P.R., Li, J.-N., Girven, K.S., Yoo, J.J., Meshik, X., Parker, K.E., Spangler, S.M., Elerding, A.J., et al. (2021). A photo-switchable GPCR-based opsin for presynaptic inhibition. *Neuron* *109*, 1791–1809.e11.

Craig, A.D., Bushnell, M.C., Zhang, E.-T., and Blomqvist, A. (1994). A thalamic nucleus specific for pain and temperature sensation. *Nature* *372*, 770–773.

Craig, A.D., Chen, K., Bandy, D., and Reiman, E.M. (2000). Thermosensory activation of insular cortex. *Nat. Neurosci.* *3*, 184–190.

Dal Monte, O., Costa, V.D., Noble, P.L., Murray, E.A., and Averbeck, B.B. (2015). Amygdala lesions in rhesus macaques decrease attention to threat. *Nat. Commun.* *6*, 10161.

Demarquay, G., and Mauguière, F. (2016). Central nervous system underpinnings of sensory hypersensitivity in migraine: insights from neuroimaging and electrophysiological studies. *Headache* *56*, 1418–1438.

Deyle, E.R., and Sugihara, G. (2011). Generalized theorems for nonlinear state space reconstruction. *PLoS One* *6*, e18295.

D’Hanis, W., Linke, R., and Yilmazer-Hanke, D.M. (2007). Topography of thalamic and parabrachial calcitonin gene-related peptide (CGRP) immunoreactive neurons projecting to subnuclei of the amygdala and extended amygdala. *J. Comp. Neurol.* *505*, 268–291.

Dobolyi, A., Irwin, S., Makara, G., Usdin, T.B., and Palkovits, M. (2005). Calcitonin gene-related peptide-containing pathways in the rat forebrain. *J. Comp. Neurol.* *489*, 92–119.

Feinstein, J.S., Adolphs, R., Damasio, A., and Tranel, D. (2011). The human amygdala and the induction and experience of fear. *Curr. Biol.* *21*, 34–38.

Ferreira, A., Pereira, M., Agrati, D., Uriarte, N., and Fernández-Guasti, A. (2002). Role of maternal behavior on aggression, fear and anxiety. *Physiol. Behav.* *77*, 197–204.

Fraser, A.M., and Swinney, H.L. (1986). Independent coordinates for strange attractors from mutual information. *Phys. Rev. A Gen. Phys.* *33*, 1134–1140.

Fu, J.-Y., Yu, X.-D., Zhu, Y., Xie, S.-Z., Tang, M.-Y., Yu, B., and Li, X.-M. (2020). Whole-brain map of long-range monosynaptic inputs to different cell types in the amygdala of the mouse. *Neurosci. Bull.* *36*, 1381–1394.

García-Díaz, D.E., Jiménez-Montufar, L.L., Guevara-Aguilar, R., Wayner, M.J., and Armstrong, D.L. (1988). Olfactory and visceral projections to the nucleus of the solitary tract. *Physiol. Behav.* *44*, 619–624.

Gauriau, C., and Bernard, J.-F. (2004). Posterior triangular thalamic neurons convey nociceptive messages to the secondary somatosensory and insular cortices in the rat. *J. Neurosci.* *24*, 752–761.

Gross, C.T., and Canteras, N.S. (2012). The many paths to fear. *Nat. Rev. Neurosci.* *13*, 651–658.

Han, S., Soleiman, M.T., Soden, M.E., Zweifel, L.S., and Palmiter, R.D. (2015). Elucidating an affective pain circuit that creates a threat memory. *Cell* *162*, 363–374.

Han, X., Chow, B.Y., Zhou, H., Klapoetke, N.C., Chuong, A., Rajimehr, R., Yang, A., Baratta, M.V., Winkle, J., Desimone, R., and Boyden, E.S. (2011). A high-light sensitivity optical neural silencer: development and application to optogenetic control of non-human primate cortex. *Front. Syst. Neurosci.* *5*, 18.

- Harriott, A.M., and Schwedt, T.J. (2014). Migraine is associated with altered processing of sensory stimuli. *Curr. Pain Headache Rep.* *18*, 458.
- Harte, S.E., Ichesco, E., Hampson, J.P., Peltier, S.J., Schmidt-Wilcke, T., Clauw, D.J., and Harris, R.E. (2016). Pharmacologic attenuation of cross-modal sensory augmentation within the chronic pain insula. *Pain* *157*, 1933–1945.
- Janak, P.H., and Tye, K.M. (2015). From circuits to behaviour in the amygdala. *Nature* *517*, 284–292.
- Jarvie, B.C., Chen, J.Y., King, H.O., and Palmiter, R.D. (2021). *Satb2* neurons in the parabrachial nucleus mediate taste perception. *Nat. Commun.* *12*, 224.
- Kim, J.J., and Fanselow, M.S. (1992). Modality-specific retrograde amnesia of fear. *Science* *256*, 675–677.
- Kim, E.J., Jacobs, M.W., Ito-Cole, T., and Callaway, E.M. (2016). Improved monosynaptic neural circuit tracing using engineered rabies virus glycoproteins. *Cell Rep.* *15*, 692–699.
- Kim, J., Zhang, X., Muralidhar, S., LeBlanc, S.A., and Tonegawa, S. (2017). Basolateral to central amygdala neural circuits for appetitive behaviors. *Neuron* *93*, 1464–1479.e5.
- Kocorowski, L.H., and Helmstetter, F.J. (2001). Calcitonin gene-related peptide released within the amygdala is involved in pavlovian auditory fear conditioning. *Neurobiol. Learn. Mem.* *75*, 149–163.
- Kruger, L., Sternini, C., Brecha, N.C., and Mantyh, P.W. (1988). Distribution of calcitonin gene-related peptide immunoreactivity in relation to the rat central somatosensory projection. *J. Comp. Neurol.* *273*, 149–162.
- Kunwar, P.S., Zelikowsky, M., Remedios, R., Cai, H., Yilmaz, M., Meister, M., and Anderson, D.J. (2015). Ventromedial hypothalamic neurons control a defensive emotion state. *Elife* *4*, e06633.
- de Lacalle, S., and Saper, C.B. (2000). Calcitonin gene-related peptide-like immunoreactivity marks putative visceral sensory pathways in human brain. *Neuroscience* *100*, 115–130.
- LeDoux, J. (2012). Rethinking the emotional brain. *Neuron* *73*, 653–676.
- LeDoux, J.E. (2000). Emotion circuits in the brain. *Annu. Rev. Neurosci.* *23*, 155–184.
- LeDoux, J.E., Ruggiero, D.A., and Reis, D.J. (1985). Projections to the subcortical forebrain from anatomically defined regions of the medial geniculate body in the rat. *J. Comp. Neurol.* *242*, 182–213.
- Li, Z., Wei, J.-X., Zhang, G.-W., Huang, J.J., Zingg, B., Wang, X., Tao, H.W., and Zhang, L.I. (2021). Corticostriatal control of defense behavior in mice induced by auditory looming cues. *Nat. Commun.* *12*, 1040.
- Liu, S., Ye, M., Pao, G.M., Song, S.M., Jhang, J., Jiang, H., Kim, J.-H., Kang, S.J., Kim, D.-I., and Han, S. (2022). Divergent brainstem opioidergic pathways that coordinate breathing with pain and emotions. *Neuron* *110*, 857–873.e9.
- López-Solà, M., Woo, C.-W., Pujol, J., Deus, J., Harrison, B.J., Monfort, J., and Wager, T.D. (2017). Towards a neurophysiological signature for fibromyalgia. *Pain* *158*, 34–47.
- Nagy, V., Bozdagi, O., Matynia, A., Balcerzyk, M., Okulski, P., Dzwonek, J., Costa, R.M., Silva, A.J., Kaczmarek, L., and Huntley, G.W. (2006). Matrix metalloproteinase-9 is required for hippocampal late-phase long-term potentiation and memory. *J. Neurosci.* *26*, 1923–1934.
- Nakamura, K., and Morrison, S.F. (2008). A thermosensory pathway that controls body temperature. *Nat. Neurosci.* *11*, 62–71.
- Nelson, K.B., Grether, J.K., Croen, L.A., Dambrosia, J.M., Dickens, B.F., Jelliffe, L.L., Hansen, R.L., and Phillips, T.M. (2001). Neuropeptides and neurotrophins in neonatal blood of children with autism or mental retardation. *Ann. Neurol.* *49*, 597–606.
- Osakada, F., Mori, T., Cetin, A.H., Marshel, J.H., Virgen, B., and Callaway, E.M. (2011). New rabies virus variants for monitoring and manipulating activity and gene expression in defined neural circuits. *Neuron* *71*, 617–631.
- Packard, N.H., Crutchfield, J.P., Farmer, J.D., and Shaw, R.S. (1980). Geometry from a time series. *Phys. Rev. Lett.* *45*, 712–716.
- Palmiter, R.D. (2018). The Parabrachial nucleus: CGRP neurons function as a general alarm. *Trends Neurosci.* *41*, 280–293.
- Price, D.D. (2002). Central neural mechanisms that interrelate sensory and affective dimensions of pain. *Mol. Interv.* *2*, 392–403.e9.
- Ren, W., and Neugebauer, V. (2010). Pain-related increase of excitatory transmission and decrease of inhibitory transmission in the central nucleus of the amygdala are mediated by mGluR1. *Mol. Pain* *6*, 93.
- Rizvi, T.A., Ennis, M., Behbehani, M.M., and Shipley, M.T. (1991). Connections between the central nucleus of the amygdala and the midbrain periaqueductal gray: topography and reciprocity. *J. Comp. Neurol.* *303*, 121–131.
- Rosen, J.B., Asok, A., and Chakraborty, T. (2015). The smell of fear: innate threat of 2, 5-dihydro-2, 4, 5-trimethylthiazoline, a single molecule component of a predator odor. *Front. Neurosci.* *9*, 292.
- Russell, F.A., King, R., Smillie, S.-J., Kodji, X., and Brain, S.D. (2014). Calcitonin gene-related peptide: physiology and pathophysiology. *Physiol. Rev.* *94*, 1099–1142.
- Russo, A.F. (2015). Calcitonin Gene-Related Peptide (CGRP): a new target for migraine. *Annu. Rev. Pharmacol. Toxicol.* *55*, 533–552.
- Salay, L.D., Ishiko, N., and Huberman, A.D. (2018). A midline thalamic circuit determines reactions to visual threat. *Nature* *557*, 183–189.
- Sato, M., Ito, M., Nagase, M., Sugimura, Y.K., Takahashi, Y., Watabe, A.M., and Kato, F. (2015). The lateral parabrachial nucleus is actively involved in the acquisition of fear memory in mice. *Mol. Brain* *8*, 22.
- Scott, S.K., Young, A.W., Calder, A.J., Hellawell, D.J., Aggleton, J.P., and Johnson, M. (1997). Impaired auditory recognition of fear and anger following bilateral amygdala lesions. *Nature* *385*, 254–257.
- Shaw, R. (1984). *The Dripping Faucet as a Model Chaotic System* (Aerial Press).
- Shinohara, K., Watabe, A.M., Nagase, M., Okutsu, Y., Takahashi, Y., Kurihara, H., and Kato, F. (2017). Essential role of endogenous calcitonin gene-related peptide in pain-associated plasticity in the central amygdala. *Eur. J. Neurosci.* *46*, 2149–2160.
- Silva, B.A., Mattucci, C., Krzykowski, P., Murana, E., Illarionova, A., Grinevich, V., Canteras, N.S., Ragozzino, D., and Gross, C.T. (2013). Independent hypothalamic circuits for social and predator fear. *Nat. Neurosci.* *16*, 1731–1733.
- Silva, B.A., Gross, C.T., and Gräff, J. (2016). The neural circuits of innate fear: detection, integration, action, and memorization. *Learn. Mem.* *23*, 544–555.
- Simons, L.E., Moulton, E.A., Linnman, C., Carpino, E., Becerra, L., and Borsook, D. (2014). The human amygdala and pain: evidence from neuroimaging. *Hum. Brain Mapp.* *35*, 527–538.
- Sugihara, G., May, R., Ye, H., Hsieh, C.-h., Deyle, E., Fogarty, M., and Munch, S. (2012). Detecting causality in complex ecosystems. *Science* *338*, 496–500.
- Takens, F. (1981). Detecting strange attractors in turbulence. In *Dynamical Systems and Turbulence*, Warwick 1980, D. Rand and L.-S. Young, eds. (Springer), pp. 366–381.
- Tong, W.H., Abdulai-Saiku, S., and Vyas, A. (2020). Medial amygdala arginine vasopressin neurons regulate innate aversion to cat odors in male mice. *Neuroendocrinology* *111*, 505–520.
- Usunoff, K.G., Itzev, D.E., Rolfs, A., Schmitt, O., and Wree, A. (2006). Brain stem afferent connections of the amygdala in the rat with special references to a projection from the parabrachial nucleus: a fluorescent retrograde tracing study. *Anat. Embryol.* *217*, 475–496.
- Veinante, P., Yalcin, I., and Barrot, M. (2013). The amygdala between sensation and affect: a role in pain. *J. Mol. Psychiatry* *1*, 9.
- Viviani, D., Charlet, A., van den Burg, E., Robinet, C., Humi, N., Abatis, M., Magara, F., and Stoop, R. (2011). Oxytocin selectively gates fear responses through distinct outputs from the Central Amygdala. *Science* *333*, 104–107.
- Wang, L., Gillis-Smith, S., Peng, Y., Zhang, J., Chen, X., Salzman, C.D., Ryba, N.J.P., and Zuker, C.S. (2018). The coding of valence and identity in the mammalian taste system. *Nature* *558*, 127–131.
- Watanakesuntorn, W., Takahashi, K., Ichikawa, K., Park, J., Sugihara, G., Takano, R., Haga, J., and Pao, G.M. (2020). Massively parallel causal inference of

- whole brain dynamics at single neuron resolution. In 2020 IEEE 26th International Conference on Parallel and Distributed Systems (ICPADS), pp. 196–205.
- Wei, P., Liu, N., Zhang, Z., Liu, X., Tang, Y., He, X., Wu, B., Zhou, Z., Liu, Y., Li, J., et al. (2015). Processing of visually evoked innate fear by a non-canonical thalamic pathway. *Nat. Commun.* 6, 8228.
- Yasui, Y., Saper, C.B., and Cechetto, D.F. (1991). Calcitonin gene-related peptide (CGRP) immunoreactive projections from the thalamus to the striatum and amygdala in the rat. *J. Comp. Neurol.* 308, 293–310.
- Yasui, Y., Nakano, K., and Mizuno, N. (1992). Descending projections from the subparafascicular thalamic nucleus to the lower brain stem in the rat. *Exp. Brain Res.* 90, 508–518.
- Yu, L.-C., Hou, J.-F., Fu, F.-H., and Zhang, Y.-X. (2009). Roles of calcitonin gene-related peptide and its receptors in pain-related behavioral responses in the central nervous system. *Neurosci. Biobehav. Rev.* 33, 1185–1191.
- Zhang, G.-W., Sun, W.-J., Zingg, B., Shen, L., He, J., Xiong, Y., Tao, H.W., and Zhang, L.I. (2018). A non-canonical reticular-limbic central auditory pathway via medial septum contributes to fear conditioning. *Neuron* 97, 406–417.e4.
- Zhang, Y., Rózsa, M., Bushey, D., Zheng, J., Reep, D., Liang, Y., Broussard, G.J., Tsang, A., Tsegaye, G., Patel, R., et al. (2020). jGCaMP8 Fast Genetically Encoded Calcium Indicators. Online resource (Janelia Research Campus). https://janelia.figshare.com/articles/online_resource/jGCaMP8_Fast_Genetically_Encoded_Calcium_Indicators/13148243.
- Zhou, Z., Liu, X., Chen, S., Zhang, Z., Liu, Y., Montardy, Q., Tang, Y., Wei, P., Liu, N., Li, L., et al. (2019). A VTA GABAergic neural circuit mediates visually evoked innate defensive responses. *Neuron* 103, 473–488.e6.

STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Bacterial and virus strains		
AAV8-hSyn-FLEX-TVA-P2A-GFP-2A-oG	Salk Institute viral vector core	RRID:Addgene_85225
EnvA-ΔG-rabies-mCherry	Osakada et al. (2011), Salk Institute viral vector core	RRID:Addgene_32635
AAV1-DIO-GFP:TetTox	Carter et al. (2015); Han et al. (2015), Laboratory of Dr. Richard Palmiter	N/A
AAV1-DIO-EYFP	Bowen et al. (2020); Han et al. (2015), Laboratory of Dr. Richard Palmiter	RRID:Addgene_27056
AAV-DIO-jGCaMP8m	Zhang et al. (2020), Janelia Viral Vector Core	RRID:Addgene_162378
AAVDJ-EF1a-DIO-hChr2(H134R)-EYFP-WPRE-pA	Karl Deisseroth, Salk Institute Viral Vector Core	RRID:Addgene_20298
AAV5-EF1a-DIO-PPO-Venus	Copits et al. (2021), Laboratory of Dr. Michael Bruchas	RRID:Addgene_139505
AAVDJ-DIO-ArchT-GFP	Han et al. (2011), Salk Institute viral vector core	RRID:Addgene_28307
Deposited data		
Custom MATLAB scripts for data analysis	This paper	Zenodo: https://doi.org/10.5281/zenodo.6667324
Experimental models: Organisms/strains		
Mouse: <i>Calca</i> -Cre:GFP	Laboratory of Dr. Richard Palmiter	N/A
Software and algorithms		
MATLAB	MathWorks	R2019a
Ethovision XT	Noldus Information Technology	Version 12
Inscopix Data Processing Software	Inscopix	Version 1.6.0.3225
pCLAMP	Molecular Devices	Version 10
BZ-W viewer	Keyence	N/A
Fluoview	Olympus	N/A
Sigmaplot	Systat Software Inc.	Version 14.0
Prism	GraphPad Software	Version 6
Illustrator	Adobe	Version CS6 and CC2018
Other		
200um 0.22 NA Fiber-optic Cannula	Laboratory of Dr. Sung Han	N/A
ProViewO Integrated Lens 0.6 mm × 7.3 mm	Inscopix	Cat# 1050-004413

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Sung Han (sunghan@salk.edu).

Materials availability

This study did not generate new unique reagents.

Data and code availability

- All data reported in this paper will be shared by the [lead contact](#) upon request.

- All original code has been deposited at Zenodo and is publicly available as of the date of publication. DOIs are listed in the [key resources table](#).
- Any additional information required to reanalyze the data reported in this paper is available from the [lead contact](#) upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mouse lines

All protocols for animal experiments were approved by the IACUC of the Salk Institute for Biological Studies according to NIH guidelines for animal experimentation. The *Calca^{Cre}*, transgenic mouse line (expressing Cre-GFP fusion protein in CGRP positive neurons) used in this study was generated from the Richard Palmiter's laboratory (Carter et al., 2013) and backcrossed with C57Bl/6J for >6 generations. GFP expression from the Cre-GFP fusion protein is undetectable in perfused slices, in sharp contrast with the fluorescence from viral expression (see Figure S1 for example slices of AAV-DIO-jGCaMP8m unilateral expression). 8–12-week-old male and female mice were used in all studies. Animals were randomized to experimental groups and no sex differences were noted. Mice were maintained on a normal 12-hour light/dark cycle and provided with food and water *ad libitum*.

METHOD DETAILS

Stereotaxic surgery for virus injection and optic fiber implantation

Mice were anesthetized by isoflurane gas anesthesia (induction at 3.5%, maintenance at 1.5–2%; Dräger Vapor® 2000, Draeger). Mice were then placed on a stereotaxic frame (David Kopf Instruments). Holes were drilled with a micromotor handpiece drill (Fore-dom) after exposure of the skull. Viral injection was performed using a syringe (65458-01, Hamilton) connected to an ultra-micropump (UMP-3, World Precision Instruments). Coordinates for unilateral (right side) and bilateral injections into SPFP and PBel were: SPFP (AP, –3.1 mm; ML, 2.0 mm; DV, –3.6 mm from bregma); PBel (AP, –5.1 mm; ML, 1.35 mm; DV, –3.5 mm). Viruses were injected at a rate of 0.08 μ L/min (total volume of 0.75 μ L for optogenetic projection studies and 0.5 μ L for all others), and syringe needles were slowly removed from the injection site 7 min after injection. To determine the inputs to CGRP^{SPFP} and CGRP^{PBel} neurons, 0.5 μ L AAV8-hSyn-FLEX-TVA-P2A-GFP-2A-oG (3.82E+12 GC/mL; Salk Institute viral vector core) (Liu et al., 2022) was injected into the SPFP or PBel of *Calca^{Cre}* transgenic mice. After three weeks, 0.5 μ L of EnvA- Δ G-rabies-mCherry (3.95E+08 GC/mL; Salk Institute viral vector core) (Liu et al., 2022; Osakada et al., 2011) was injected into the same regions. Mice were sacrificed five days after the final injection. To silence the CGRP^{SPFP} and CGRP^{PBel} neurons, AAV1-DIO-GFP:TetTox (2.51E+12 GC/mL; reported validity in *Calca^{Cre}* transgenic mice) (Carter et al., 2015; Han et al., 2015) or AAV1-DIO-EYFP (2.12E+12 GC/mL; reported validity in *Calca^{Cre}* transgenic mice) (Bowen et al., 2020; Han et al., 2015) was injected into the SPFP (0.5 μ L) or PBel (0.3 μ L), respectively, of *Calca^{Cre}* transgenic mice, and experiments were performed two weeks after injection. For Miniscope single-cell calcium imaging experiments, 0.5 μ L AAV-DIO-jGCaMP8m (5E+12 GC/mL) (Liu et al., 2022; Zhang et al., 2020) was injected into the SPFP or PBel of *Calca^{Cre}* mice and an integrated microendoscope lens (0.6 mm \times 7.3 mm, Inscopix) was implanted. Coordinates for microendoscope lens implantation for SPFP and PBel were: SPFP (AP, –3.1 mm; ML, 2.0 mm; DV, –3.6 mm); PBel (AP, –4.95 mm; ML, 1.5 mm; DV, –3.6 mm).

For electrophysiological experiments, AAVDJ-EF1a-DIO-hChR2(H134R)-EYFP-WPRE-pA (4.5E+12 GC/mL) (Liu et al., 2022) was injected into the SPFP (0.5 μ L) or PBel (0.3 μ L) of *Calca^{Cre}* mice. Experiments were performed two weeks after viral injection for recording SPFP and PBel neurons or four weeks after injection for recording cells in terminal regions. For optogenetics experiments, AAVDJ-EF1a-DIO-hChR2(H134R)-EYFP-WPRE-pA, AAV5-EF1a-DIO-PPO-Venus (3E+13 GC/mL) (Copits et al., 2021), AAVDJ-DIO-ArchT-GFP (1.26E+12 GC/mL) (Han et al., 2011; Liu et al., 2022) or AAV1-DIO-EYFP was injected into the SPFP (0.5 μ L) or PBel (0.3 μ L) of *Calca^{Cre}* mice, and custom-made mono fiberoptic cannulae (200 μ m diameter, 0.22 NA) were implanted above SPFP (0.5 mm above the injection site), PBel (0.5 mm above the injection site), LA (AP, –1.8 mm; ML, 3.6 mm, DV, –4.0 mm from bregma), or CeA (AP, 1.2 mm; ML, 2.7 mm; DV, –4.2 mm from bregma). Experiments were performed two weeks after injection for cell body manipulations or four weeks for terminal manipulations.

Histology and quantification of rabies tracing experiment

Mice were intracardially perfused with 4% paraformaldehyde in PBS (phosphate buffered saline) 5 days after the rabies virus injection. Spinal cords were post-fixed at 4°C for 1 h and dehydrated with 30% sucrose at 4°C overnight. 40 μ m transverse sections were obtained with a cryostat (CM, 1950; Leica) throughout the spinal cord. Spinal cord slices were directly dry mounted on superfrost plus microscope slide glasses (12-550-15, Fisher Scientific). The labeled neurons were counted manually by dividing the transverse spinal sections into four groups (cervical, thoracic, lumbar, and sacral) or different dorsal horn layers. Brains were kept in 4% PFA overnight for post-fixation and dehydrated in 30% sucrose for 1–2 days before sectioning. Frozen brains were cut into 50 μ m coronal slices with a cryostat and stored in PBS before mounting. Both spinal cord and brain tissues were mounted on a slide glass with a DAPI containing mounting solution (0100-20, SouthernBiotech). Half of the slices collected throughout the brain (100 μ m apart) were mounted on slide glasses and all of them were imaged with the automatic fluorescence microscope (BZ-X710, Keyence) using included imaging software (BZ-X viewer, Keyence). Red fluorescence neurons indicating the presynaptic input neurons were counted manually. Brain regions were delineated by comparing with the Allen Brain Atlas Reference Atlases.

Multi-sensory aversive stimuli experiments

Auditory stimuli

For the Miniscope single-cell calcium imaging experiment, mice were placed in a cylindrical arena (11 cm diameter with 15 cm height) with home cage bedding and habituated for 30–120 min. The experiment was controlled to have no sudden unrelated noise or visual cues. An intense sound (85 dB, 2 sec) or a control sound (70 dB, 2 sec) was played.

For the loss-of-function experiments, mice were placed inside an open field chamber and habituated for 10 min. After 1 min baseline, an intense sound (85 dB, 2 sec) was delivered three times with an inter-stimulus interval of 28 sec. Laser was shined throughout the experiment for terminal inhibition. All trials were recorded by a USB camera (DFK 33GX236, Imagine Source) attached to a computer, and freezing behavior was analyzed using video-tracking software (Ethovision XT, Noldus). Freezing behavior were monitored starting from the first stimuli to the end of the experiment and analyzed with Ethovision (“Not moving” 1.25–1.5 cm/s, for 3s samples).

Visual stimuli

For the Miniscope single-cell calcium imaging experiment, mice were placed in the same arena as the auditory experiment. The experiment was controlled to have no sudden unexpected noise or visual cues. An expanding looming stimulus (2 sec, 25-cm diameter) was delivered three times with 10 sec inter-stimulus interval with a LED screen facing the arena from above. For the loss-of-function experiment, mice were placed in a cage with bedding, positioned under the same LED screen, and habituated for 20–30 min. When mice were in the center of the cage, the expanding looming stimulus (2 sec) was delivered three times with 10 sec inter-stimulus interval. Laser was shined throughout the experiment for terminal inhibition. All trials were recorded by a USB camera (DFK 33GX236, Imagine Source) attached to a computer, and freezing behavior was analyzed using video-tracking software (Ethovision XT, Noldus). Freezing behavior were monitored starting from the first stimuli to the end of the experiment and analyzed with Ethovision (“Not moving” 1.25–1.5 cm/s, for 3s samples).

Gustatory stimuli

For the Miniscope single-cell calcium imaging experiment, mice were placed in the same arena as used for auditory and visual stimulus experiments, but with an additional 2-cm drilled hole on the bottom. The water bottle spout was inserted into this hole, and the calcium signal was measured when mice were licking. The bottle was filled with water or quinine (0.5 mM, QU109, Spectrum Chemical).

For the loss-of-function experiment, mice were water-deprived overnight. The next day, mice were placed in a standard mouse cage with a bottle containing either water or 0.5 mM quinine inserted into the water valve slot. Mice were allowed to drink for 10 min without habituation. Laser was turned on throughout the terminal inhibition experiment. All trials were recorded by a USB camera (DFK 33GX236, Imagine Source) attached to a computer, and licking behaviors were counted manually.

Olfactory stimuli

For the Miniscope single-cell calcium imaging experiment, mice were placed in the same arena as used for gustatory stimulus experiments. A cotton swab soaked with either water- or trimethylthiazoline (TMT, 97%, 5 μ L, 1G-TMT-97, BioSRQ) was inserted into the arena through the hole. Calcium signals were measured when mice smelled the cotton swab. For the loss-of-function experiment, mouse movement in a two-chamber arena (30 \times 60 \times 30 cm) was tracked using a USB camera (DFK 33GX236, Imagine Source) and video-tracking software (EthoVision XT 12, Noldus). Two Petri dishes with small holes were placed into each chamber—one at the corner of the left chamber, and the other at the corner of the right chamber. On day 1, mice were allowed to habituate and explore the arena for 10 min. The next day, a water-soaked cotton swab was placed in one dish, and a TMT-soaked cotton swab was placed in the other. Mice were first placed at the center of the arena and monitored for 10 min as they interacted with the two dishes. Laser was turned on throughout the terminal inhibition experiment.

Somatosensory stimuli

A threat conditioning chamber (26 \times 30 \times 33 cm, ENV-007CT, MED Associates) with a metal grid floor (ENV-005, MED Associates) connected to a standalone aversive electric shock stimulator (ENV-414S, MED Associates) was used for foot shock delivery. A USB camera (DFK 33GX236, Imagine Source) was connected to a computer, and video tracking software (Ethovision XT, Noldus) was used for shock delivery and behavioral analysis. The chamber was enclosed in a light- and sound-attenuating cubicle (ENV-018MD, MED Associates). The chamber was cleaned with 70% ethanol and double-distilled water after each trial.

For both Miniscope and loss-of-function experiments, mice were placed inside the chamber without habituation. After a 2 min baseline, an electric shock (2 sec, 0.6 mA) was delivered, and behavior was recorded for 2 more min. For loss-of-function experiments, freezing behavior was monitored one day before (habituation), immediately after (conditioning), and one day after the shock (post-test). For the terminal inhibition experiment, procedures were the same as for the whole-population inhibition experiment, except that the laser was turned on throughout the conditioning and post-test periods.

Cued threat conditioning

The same threat-conditioning chamber and video tracking system were used for cued threat conditioning as during somatosensory stimulus delivery. Two speakers (AX210, Dell) were placed beside the chamber to deliver auditory cues to act as conditioned stimuli (CS). The chamber was enclosed in a light- and sound-attenuating cubicle (ENV-018MD, MED Associates). The chamber was cleaned with 70% ethanol and double-distilled water between each trial. On day 1 (habituation day), mice were introduced to the chamber after connecting the optic fiber or miniature microscope. After a 2-min baseline period, CS (30 sec, 2 kHz pure tone) was delivered six times with random intervals. On day 2 (conditioning day), mice were returned to the same arena. After a 1-min baseline, mice received 5 foot-shocks (2 sec, 0.2 mA) (US) co-terminating with CS at random intervals. On day 3 (retrieval day), mice were

assessed first in a context test, then a cue test after 4 h. For the context test, mice were placed into the same chamber as on the conditioning day for 3 min; for the cue test, mice were placed into a new context (a glass cylinder wrapped with a non-transparent material), and the 30-s CS was delivered alone three times (i.e., not paired with the US). For optogenetic terminal inhibition, optic fiber was connected at all times and the laser was turned on during conditioning or retrieval periods (detailed info in the [Optogenetics](#) section). Freezing behavior was analyzed by video-tracking software (EthoVision XT 12, Noldus, “Not moving” 1.25–1.5 cm/s for 3-s), which has been shown to match the manual scoring based on the characteristic rigid posture ([Han et al., 2015](#)). “BL” in the cue test indicates freezing during baseline before the first tone was delivered, and “CS+” is the average freezing during the three tones.

Miniscope single-cell calcium imaging

Calcium activity was recorded at 10 or 20 Hz and 0.4–1.2 mW/mm² LED power using an Inscopix nVista miniature microscope. Animals were habituated with the miniature microscope at least 3 times before the actual recording. On the testing day, over-night water-restricted animals were exposed to the following stimuli 3 times each with at least 2-min inter-trial-interval: 10 μ L of 0.5 mM quinine (gustatory), cotton tip with 2 μ L of 99% TMT (olfactory), 85 dB intense sound burst (auditory; 2 sec); 1 expanding looming circle (visual; 2 sec, 25-cm diameter); 0.2 mA foot shock (somatosensory; 2 sec). All stimuli were delivered in the same miniscope-baseplate docking configuration to facilitate the registration of cells across sessions. In other words, we performed all recording sessions in one run, and during post-processing, we concatenated different videos to a single file and used it to identify cells. This design could eliminate the need for longitudinal registration of cells which would cause significant loss of cell number. The control experiments were performed during a separate session with the following stimuli delivered 3 times to water-restricted animals: 10 μ L water (gustatory), cotton tip with 2 μ L water (olfactory), 70 dB gentle sound (auditory; 2 sec); 1 small looming circle (visual; 2 sec, 5-cm diameter); no foot shock (somatosensory). Ethovision XT 12 was used to record behavior videos and trigger calcium activity recordings. The onset of behavioral responses was manually identified from the behavior video.

For threat conditioning recordings, the behaviors were performed following the previous “*Cued threat conditioning*” session. Separate video files were analyzed for each day without cross-registration of cells.

For data post-processing, Inscopix data processing software was used. The main steps and related parameters are listed as follows: “Preprocessing” (2x spatial downsampling, no temporal downsampling, determine crop area, fix defective pixels); “Spatial Filter” (low cut-off: 0.005 pixel⁻¹; high cut-off: 0.5 pixel⁻¹); “Motion Correct” (using the first frame as the global reference frame); “Identify Cells” (CNMFe, starting with cell diameter = 15 μ m, min pixel correlation = 0.9, min pixel-to-noise ratio = 10, then adjusting parameters based on cell detection quality compared to manual identification); “Additional Tools” (deconvolve traces); “Export” (export countour map and cell traces). The output trace unit is change in fluorescence over noise ($\Delta F/F$).

To classify cells that respond to each stimulus, we calculated z-scored calcium activity for each trial, including pre-stimulation and post-stimulation episodes. The pre-stimulation baseline period is defined as –5 to –1 sec (total 4 sec) before stimulus onset, and post-stimulation period is defined as 0 to 10 sec (total 10 sec) after stimulus onset. Since both SPFP and PBeI neurons displayed baseline activity, to better visualize evoked calcium responses, we normalized the pre-stimulus baseline activity to zero by subtracting the averaged baseline activity from the whole trace. A given cell was classified as stimulus-responsive if the maximum z-score value during the post-stimulation period was larger than 2.5 in all 3 trials.

Heatmap visualization of individual neuronal activity was generated from 15-sec peri-stimulus traces (5 sec before and 10 sec after stimulus delivery) for all cells with each row indicating the same neuron across stimuli, grouped by the maximum responsive stimuli. Averaged traces of neuronal activities during the multi-sensory threat and control trials were also plotted, where representative neurons are the ones showing maximum responses to each individual stimulus. For the cued threat conditioning experiments, area under the curve (AUC) was calculated from the activities 0 to 5 sec after the stimulus onset (the stimulus refers to the tone during the habituation and cue tests, and shock during the conditioning test).

Convergent cross-mapping

We measured the network relationships between neurons using convergent cross-mapping (CCM) ([Sugihara et al., 2012](#)), a nonlinear time series embedding method based on the Takens theorem ([Packard et al., 1980](#); [Takens, 1981](#)) and its generalized form ([Deyle and Sugihara, 2011](#)). CCM builds low-dimensional manifolds from time series and makes predictions across variables to determine causation between variables. However, in the cases where there is synchrony due to strong coupling, this causal determination cannot be made.

Here, the inputs of CCM were single-cell calcium traces pooled from multiple animals. Neurons from the same animal were highly correlated due to the stimulation protocol. Therefore, our prediction score does not measure causality within a neuron pair, but it does measure the similarity across two embedded manifolds. This non-traditional use of CCM is conceptually similar to using the prediction skills of reservoir computing models to assess the similarity between systems. This CCM measure of classification contains the information of cross-correlation, dynamic time warping, and cases for which individual local dynamical events are out of order. The main difference between CCM and the traditional cross-correlation is that CCM correlations are in phase space and are based on topological similarities between the embedded time series. CCM correlations are measured on the manifolds of the embedded time series, taking into account the influences of latent unobserved variables that are nevertheless observable as influences on the neuronal time series. This allows CCM-based comparisons to detect similarities of dynamics that are not temporally coincident, but that occur out of sequence or with delays.

For all five sensory stimuli, we performed CCM on the peri-stimulus activity (5 sec before and 5 sec after the stimulus) of each neuron pair. The output of the CCM is a prediction score describing the similarity of manifold (neuronal dynamics and information content) within each neuron pair (Shaw, 1984). The manifold similarities were then used as inputs to compare the population recruited by each stimulus. Specifically, the prediction score matrices for each stimulus were used as inputs for hierarchical clustering using Pearson correlation as the distance metric (MATLAB “clustergram”). The clusters in the clustergram denoted the group of neurons that shared the same “broadcasters” and “receivers,” and could thus be approximated as the functional network recruited by a particular stimulus. The exact order of neurons in each clustergram (i.e., seed) was used to plot the remaining four heatmaps corresponding to the four other stimuli. We reasoned that if the same clusters could be detected in the other heatmaps, the same functional networks could be reused to respond to different stimuli. The similarity between each heatmap and its seed heatmap was quantified by their Pearson correlation coefficient for all entries. The similarities across 25 matrices were presented as a 5x5 matrix.

CCM analyses and predictions were calculated using the R package rEDM 0.7.4 (<https://ha0ye.github.io/rEDM/>) in the RStudio environment or with mpEDM (Watanakeesuntorn et al., 2020). These packages were run on a dual Intel Xeon Gold 6148 Server with 384GB RAM, a two-socket AMD EPYC Rome 7742 with 1TB RAM and an NVIDIA A100 40GB GPU, and an Intel Core i9 2.4 GHz MacBook Pro with 32 GB RAM. Key parameters were determined individually by lagged coordinate embedding using the simple function implementation in rEDM to optimize predictive skill as measured by the prediction score $\rho(\Sigma_{\text{obs}(1,2,\dots,n)}/\Sigma_{\text{pred}(1,2,\dots,n)})$. Parameters included the delay τ , which calculates the characteristic timescale of the series, and the embedding dimensionality E , which estimates the number of variables driving the system. The choice of τ was informed by minimizing mutual information (Fraser and Swinney, 1986), approximately corresponding to an autocorrelation of ~ 0.3 . To prevent data contamination, an exclusion radius larger than the smoothing window of five timesteps was applied.

Optogenetics

A 470 nm laser (LRD-0470-PFFD-00100-05, LaserGlow Tech) was used for ChR2 (4 ms pulse, 20 or 40 Hz, 8–9 mW) and PPO (10 ms pulse, 10 Hz, 8–9 mW) experiments and a 589 nm laser (LRS-0589-GFM-00050-05, continuous, 6–7 mW) was used for ArchT experiments. Optic fibers were bilaterally connected to pre-implanted optic ferrules on the mice.

Real-time place aversion (RTPA)

A two-chamber arena (30 × 60 × 30) was used for the RTPA test. Behavior was tracked with a USB camera (DFK 33GX236, Imagine Source) using video-tracking software (EthoVision XT 12, Noldus). After connecting the optic fiber, mice were placed in one side of the chamber. No stimulation was given for a 10-min baseline period. One side of the chamber was then pseudorandomly selected, and the mouse was photostimulated (20 Hz for cell body stimulation, and 40 Hz for terminal stimulation, 8–9 mW) for as long as it stayed in that chamber. The test ended 20 min after initiation of the photostimulation phase. The stimulated chamber was counter-balanced between animals. Mice showing over 15% preference to one side during baseline were excluded. The first 10 min (baseline) was used for laser “OFF” and the last 10 min (20–30 min) during the photostimulation phase was laser “ON”. Time spent in the stimulated side of the chamber (%) was used for RTPA data presentation.

Auditory cue dependent optogenetic conditioning

The experimental paradigms for auditory cue-dependent optogenetic conditioning were the same as those described in the “Cued threat conditioning” section above, except that 10-s photostimulation (20 Hz frequency for cell body and 40 Hz for terminal stimulation, 8–9 mW intensity) was used as the US instead of foot shock. The 10-s photostimulation is an artificial protocol that was used to induce threat memory in a previous study (Han et al., 2015). It does not mimic the endogenous activity.

Terminal inhibition during multi-sensory aversive stimuli

Laser (10 Hz 470 nm for PPO and continuous 589 nm for ArchT) was kept on during the whole experiment to allow thorough inhibition during the stimuli and avoid rebound activity during the conditioning trial.

Terminal inhibition during cued threat conditioning

The experimental paradigms for terminal inhibition during cued threat conditioning were the same as those described in the “Cued threat conditioning” section above, except that terminal photoinhibition was performed during conditioning or retrieval period. To inhibit during the conditioning period, laser (10 Hz 470 nm for PPO and continuous 589 nm for ArchT) was turned on from 10s after the start of the first tone until the end of the conditioning trial. This was to allow thorough inhibition during the shock episode and avoid rebound activity during the conditioning trial. To inhibit during the retrieval period, laser was kept on during the whole context and cue test.

Preparation of acute brain slices and electrophysiology with optogenetics

Calca^{Cre} mice injected with AAV-DIO-ChR2-EYFP in the SPFP or PBel were anesthetized with isoflurane and perfused via the vascular system using ice-cold cutting solution (110.0 mM choline chloride, 25.0 mM NaHCO₃, 1.25 mM NaH₂PO₄, 2.5 mM KCl, 0.5 mM CaCl₂, 7.0 mM MgCl₂, 25.0 mM glucose, 5.0 mM ascorbic acid and 3.0 mM pyruvic acid, bubbled with 95% O₂ and 5% CO₂). After decapitation, brains were quickly removed and chilled in an ice-cold cutting solution. Coronal slices containing the pLC or the amygdaloid complex (300 μm) were cut by using a Leica VT 1200S Vibratome (Leica Biosystems), and subsequently transferred to a storage chamber containing artificial cerebrospinal fluid (aCSF; 124 mM NaCl, 2.5 mM KCl, 26.2 mM NaHCO₃, 1.2 mM NaH₂PO₄, 13 mM glucose, 2 mM MgSO₄ and 2 mM CaCl₂, at 32°C, pH 7.4, bubbled with 95% O₂ and 5% CO₂). After at least 30 min recovery time, slices were maintained at room temperature (22–24°C) for at least 60 min before use. Slices were transferred into the recording chamber, perfused with aCSF (flow rate ~ 2 mL/min). The temperature of aCSF was held constant at 32°C by

TC-324C temperature controller (Warner Instruments). The neurons were visualized under trans-illumination by far-red microscope (Scientifica) and pyramidal neurons (LA, pIC) or medium spiny neurons (Astr) were patched based on the morphology. Recording glass electrode (3.0–5.0 M Ω) was back filled with internal solution: CsMeSO₃ 130 mM, CsCl 5 mM, HEPES 10 mM, MgCl₂ 2.5 mM, EGTA 0.6 mM, Sodium phosphocreatine 10 mM, Na₂ATP 4 mM and Na₃GTP 0.4 mM, pH 7.23, 285 Osm). Whole-cell patch clamp was performed with Multiclamp 700B amplifiers (Molecular Devices). Signals were digitized at 10 kHz with Digidata 1550B (Molecular Devices). 2–5 ms pulse of 470 nm LED light (TTL from Clampex to Cool Led pE-300) was illuminated through 40X NA 0.8 objective lens at 0.1 Hz to evoke optogenetically evoked postsynaptic current. The internal solution was calculated to make chloride reversal potential equal -70 mV. EPSCs were recorded at -70 mV, and IPSCs were recorded at 0 mV. CNQX (10 μ M) was perfused to verify the glutamatergic synapse. EPSCs and IPSCs were analyzed using pCLAMP 10 software (Molecular Devices). Latency was defined as the time from the initiation of the 470 nm LED light to the 5% of EPSC or IPSC peak amplitude.

Imaging

The images were taken with an automatic fluorescence microscope (BZ-X710, Keyence) using included imaging software (BZ-X viewer, Keyence) or with a scanning confocal microscope (FV 1000, Olympus) using with Fluoview software (Olympus). For quantification purposes, images were processed with the same gain, offset, and exposure time. Cell counting for retrograde tracing was done manually.

QUANTIFICATION AND STATISTICAL ANALYSIS

Quantifications of miniscope single-cell imaging, monosynaptic rabies tracing, behavioral tests, and electrophysiology are described in the corresponding sections of text and methods. All data are presented as mean \pm SEM and were analyzed using Student's t-test, one-way ANOVA with Tukey's post hoc comparison, and two-way ANOVA with Sidak's post hoc comparison when they passed the Shapiro-Wilk normality test. If not ($p < 0.05$), data were analyzed with MannWhitney test and Kruskal-Wallis one-way ANOVA with Dunn's multiple comparisons test. All statistical analyses were done using Prism 6 (GraphPad Software) and Sigmaplot 14.0 (Systat Software Inc.). NS $p > 0.05$, * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. Full details of statistical tests in individual figures are described in the figure legend and [Table S5](#).