Central Amygdala Somatostatin Neurons Gate Passive and Active Defensive Behaviors

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The central amygdala (CeA) has a key role in learning and expression of defensive responses. Recent studies indicate that somatostatin-expressing (SOM+) neurons in the lateral division of the CeA (CeL) are essential for the acquisition and recall of conditioned freezing behavior, which has been used as an index of defensive response in laboratory animals during Pavlovian fear conditioning. However, exactly how these neurons participate in fear conditioning and whether they contribute to the generation of defensive responses other than freezing remain unknown. Here, using fiber-optic photometry combined with optogenetic and molecular techniques in behaving mice, we show that SOM+ CeL neurons are activated by threat-predicting sensory cues after fear conditioning and that activation of these neurons suppresses ongoing actions and converts an active defensive behavior to a passive response. Furthermore, inhibition of these neurons using optogenetic or molecular methods promotes active defensive behaviors. Our results provide the first in vivo evidence that SOM+ neurons represent a CeL population that acquires learning-dependent sensory responsiveness during fear conditioning and furthermore reveal an important role of these neurons in gating passive versus active defensive behaviors in animals confronted with threat.

Key words: active avoidance; central amygdala; fear; fiber-optic photometry; optogenetics; somatostatin neurons

Significance Statement

The ability to develop adaptive behavioral responses to threat is fundamental for survival. Recent studies indicate that the central lateral amygdala (CeL), in particular its somatostatin-expressing neurons, is crucial for both learning and the expression of defensive response. However, how exactly these neurons participate in such processes remains unclear. Here we show for the first time in behaving mice that the somatostatin-expressing neurons in the CeL acquire learning-dependent responsiveness to sensory cues predicting a threat. Furthermore, our results indicate that these neurons gate the behavioral output of an animal: whereas high activity in these neurons biases toward passive defensive responses, low activity in these neurons allows the expression of active defensive responses.

Introduction

In the face of threat, animals typically show passive reactions such as freezing or suppression of ongoing activity or active defensive behaviors such as flight or fight (Bouton and Bolles, 1980; Blanchard et al., 2001). The ability to select an appropriate action in such situations is essential for survival. Freezing behavior and suppression of ongoing activity are commonly measured as indices of fear in Pavlovian fear conditioning (LeDoux et al., 1988; Killcross et al., 1997; Amorapanth et al., 1999; Johansen et al., 2011). Extensive studies based on these measurements have established the amygdala, including its basolateral (BLA) and central (CeA) nuclei, as the centerpiece of fear regulation circuits (LeDoux et al., 1988; Duvarci and Pare, 2014; Herry and Johansen, 2014). In particular, the BLA is known as the amygdala nucleus where inputs of different modalities converge and is thought to be critical for the formation of associations between conditioned stimulus (CS) and unconditioned stimulus (US), and thus the formation of memory, during conditioning (for recent reviews, see Duvarci and Pare, 2014; Herry and Johansen, 2014; Gründemann and Lüthi, 2015).

Recent studies indicate that a major cell population in the lateral division of the CeA (CeL), the somatostatin-expressing (SOM+) neurons, is essential for both the acquisition and the
recall of aversive memories, which are measured as the expression of conditioned freezing (Li et al., 2013; Penzo et al., 2014; Penzo et al., 2015). In general, neurons involved in the formation, storage, or recall of memories, such as those in the BLA, should represent the memory-associated sensory cues (Quirk et al., 1995). However, whether SOM Cre cells neurons have such capacity is unknown because, to date, their activity has not been recorded in vivo in behaving animals. In addition, the role of these neurons in defensive responses other than freezing has yet to be examined. The present study was devised to address these questions.

Materials and Methods

Animals
Before surgery, mice were group housed under a 12 h light/dark cycle (7:00 A.M. to 7:00 P.M. light) with food and water available ad libitum. Animals with optical fiber implants were housed singly. The Som-cre (Taniguchi et al., 2011; Li et al., 2013), Ai14 (Madsen et al., 2010), Ai35 (Madsen et al., 2012), and Ai32 (Madisen et al., 2012) mice were described previously and were purchased from The Jackson Laboratory. All mice were bred onto C57BL/6J genetic background. The Som-cre; Ai14 mice, which were heterozygous for both the Cre allele and the Lox-Stop-Lox-ttdTomato allele, were bred by crossing homozygous Som-cre mice with homozygous Ai14 reporter mice. The Som-cre Ai35 mice, which were heterozygous for both the Cre allele and the Lox-Stop-Lox-Arch-GFP allele, were bred by crossing homozygous Som-cre mice with homozygous Ai35 mice. The Som-cre Ai32 mice, which were heterozygous for both the Cre allele and the Lox-Stop-Lox-ChR2-EYFP allele, were bred by crossing homozygous Som-cre mice with homozygous Ai32 mice. Male and female mice 40–60 days of age were used for all experiments. All behavioral experiments were performed during the first 4 h of the dark cycle. All procedures involving animals were approved by the Institutional Animal Care and Use Committees of Cold Spring Harbor Laborator and were performed in accordance with National Institutes of Health standards.

Stereotoxic surgery
The AAV1.Syn.Flex.GCaMP6f.WPRE.SV40 and AAV9.EF1a.DIO.eYFP.WPRE.hGH viruses, which express GCaMP6f and eYFP, respectively, in a Cre-dependent manner, were purchased from Penn Vector Core (Philadelphia, PA). The AAV2.CAG.Flex.TelC-eGFP.WPRE.bGH virus, which expresses TelC in a Cre-dependent manner, was custom made by the vector core at the University of North Carolina.

Standard surgical procedures were followed for stereotoxic injection (Li et al., 2013; Penzo et al., 2015). Briefly, mice were anesthetized with ketamine (100 mg per kg of body weight) supplemented with dexmedetomidine hydrochloride (0.4 mg per kg) and positioned in a stereotaxic frame (David Kopf Instruments). The animals were anesthetized with ketamine (100 mg per kg of body weight) supplemented with dexmedetomidine hydrochloride (0.4 mg per kg) and positioned in a stereotaxic frame. The mice were placed on the stereotaxic device, and the scalp was incised and reflected. A craniotomy was performed, and the exposed skull was carefully cleaned with a sterile cloth. A 26-gauge needle was used to inject a small amount of saline (0.3 μL) around the dura mater to separate it from the underlying brain tissue. The needle was then inserted at the desired coordinates, and the solution was slowly injected (5–20 μL) over a period of 5–20 s. The needle was then withdrawn, and the incision was closed with sutures. The mice were then allowed to recover for 1–2 h before being returned to their home cages.

For the in vivo photostimulation experiments, Som-CreAi32 or Som-CreH2B (as the control mice) were bilaterally implanted with optical fibers that were placed above CeL (coordinates of the fiber tip: −1.22 mm from bregma; 2.9 mm lateral from the midline, and 4.3 mm vertical from pial surface). Following the above procedures, a small piece of metal bar was mounted on the skull, which was used to hold the mouse in the head fixation frame during experiments.

Behavioral tasks

Licking behavior. Water deprivation started 23 h before training. Mice were trained in the head fixation frame for 30 min on the first day and for 10 min daily in the following days. A metal spout was placed in front of the animal’s mouth for water delivery. The spout also served as part of a custom “lickometer” circuit, which registered a lick event each time a mouse completed the circuit by licking the spout while standing on a metal floor. The lick events were recorded by a computer through a custom software written in LabView (National Instruments). Each lick triggered a single opening of a water valve calibrated to deliver 0.3 μL of water.

It took mice 4–7 d to achieve stable licking, the criterion for which was 10 min of continuous licking without any gap longer than 10 s.

Fear conditioning. Mice with stable licking behavior were first subjected to sound habituation sessions (1 session/d for 2 d), during which an auditory stimulus was presented through a computer speaker in each trial. Each stimulus was composed of 5 pips of pure tone (8 kHz, 70 dB). Pip duration was 5 ms and the inter-pip interval was 750 ms. Each habituation session contained 10 trials with variable intertrial intervals (45–75 s). After habituation, mice were conditioned by pairing the auditory stimulus (CS) with a tail shock (US: 100 μA for 500 ms), which was generated from an isolator (ISO-Flex; A.M.P. Instruments) and delivered through a pair of wires secured to the tail with silicon tubing. A probabilistic reinforcement schedule (Erlich et al., 2012) was used, in which mice were exposed to 1 session of 10 trials daily for 4 consecutive days and in which the CS was paired with the US in only 30% of trials. An advantage of this reinforcement schedule is that it leads to the formation of fear memories resistant to extinction, which can potentially affect imaging experiments requiring repeated presentations of CS to conditioned mice.

We used a lick suppression index to quantify animals’ performance in this task. The lick suppression index was calculated as follows: lick suppression index = (Lpre − Lcs)/(Lpre + Lcs), where Lpre is the number of licks in the 5 s period before CS onset and Lcs is the number of licks in the 5 s CS period (Erlich et al., 2012).

In vivo fiber-optic photometry

The custom fiber-optic photometry system used a diode laser (λ = 473 nm; OSEM Laser Systems) that was first passed through a bandpass filter (ET480/20s; Chroma Technology), subsequently reflected off of a dichroic mirror (FF495; Semrock), and finally routed into a patch cord. The laser beam was gated by a shutter (Uniblitz Electronic) or an acousto-optic modulator (MTS110-A3-VIS; AA Opto-Electronic) placed in the light path. To reduce photo bleaching, laser illumination started 3 or 4 s before CS onset and ended 5 s after CS offset.

GCaMP6f fluorescence was routed and collected by the same optical fiber and objective and then passed through the dichroic mirror followed by a long-pass filter (510LP; Chroma Technology), and subsequently fed onto a photomultiplier tube (PMT) (R9110; Hamamatsu) by a focusing lens. Photo current signals were amplified using a 700 MHz pulse amplifier (ZPUL-30P; MiniCircuits), the output of which was digitized through a computer (PRL-350TTL; Pulse Research Lab) (Driscoll et al., 2011). The threshold for the comparator was calibrated slightly above the dark current elicited by the PMT. Its output was acquired through a 100 MHz counter (PCIe6321; National Instruments), further queried at 1 kHz rate, returning binned photon counts using custom software written in LabView (National Instruments). To correct for baseline drifting caused by photo-bleaching, a time-dependent baseline B(t) was computed as described previously (Jia et al., 2011). The percentage ∆F/F was calculated as 100 × (F(t) − F(0))/F(0), where F(t) is the raw fluorescence signal at time t. Average ∆F/F values during the pre-CS (2 s) and CS (5 s) periods in each trial were used for statistical analysis.
In vivo optogenetics
For bilateral optogenetic stimulation in the CeL, a branched patch-cord (BFP(2)_{105/125/900-0.22_1m_FC-2xZF1.25; Doric Lenses) for light delivery was connected at one end to a laser source (/H9261/H11005 473 nm; OEM Laser Systems) and, at the other end, which was composed of two terminals, to two CeL-implanted optical fibers through sleeves (Thorlabs). For photostimulation-induced lick suppression, the stimuli were 5 ms 30 Hz light pulses delivered for 10 s. For photostimulation during active avoidance task, 5 ms 30 Hz light pulses were delivered 1 s before and coterminal with a 6 s CS (sound). Laser intensity was 10 mW measured at the end of optical fiber.

Active avoidance task
Mice were habituated on a custom running wheel under the head fixation frame for 1 h the day before training. The running wheel (diameter, 14 cm; width 8 cm) was made using a 3D printer (MakerBot Replicator 2; MakerBot). A rotary encoder (YUMO-E6B2-CW23E-1024; SparkFun Electronics) was attached to the wheel and connected with a microcontroller (Arduino UNO R3; SparkFun Electronics). The microcontroller converted the digital inputs from the rotary encoder into analog signals reflecting movement velocity, which was in turn recorded by custom software written in Labview (National Instruments).

During training, a CS consisting of 6 pips of 500 ms white noise was presented, indicating that an air puff to the face would be delivered in 1.5 s. The air puff could be prevented or aborted if mice ran in response to the CS and reached a threshold speed (5 cm/s). Mice were trained daily for one session of 300 trials. Once mice reached 75% performance (calculated as the percentage of successful avoidance trials), they were subjected to a testing session the following day. During testing, mice were exposed to 200 trials of CS presentations, 50% of which (randomly distributed) were delivered together with photostimulation in the CeL. Running speed and time stamps for CS presentations were recorded.

Statistics and data presentation
All data are presented as mean ± SEM. All statistics are indicated where used. Data were analyzed with Origin8 Software (OriginLab). Normality was tested by D’Agostino–Pearson or Shapiro–Wilk normality test. Behavioral tests and imaging data acquisition were performed by an investigator with knowledge of the identity of the experimental groups. All behavior experiments were controlled by computer systems and data were collected and analyzed in an automated and unbiased way. No single data points were excluded. Virus-injected animals in which the injection site was incorrect were excluded.

Results
To investigate how SOM^{+} CeL neurons contribute to the generation of defensive behaviors, we set out to monitor the activity of these neurons in vivo in mice performing a conditioned lick suppression task (Erlich et al., 2012) in which a CS predicting a noxious US suppresses licking for water in water-deprived mice (Fig. 1A–E). Mice in this task had stable and high baseline lick rate (Fig. 1D,E). After conditioning by pairing the CS (pips of sound) with the US (tail shocks) (Fig. 1A–C), sound presentation alone caused marked reduction in lick rate, indicating the formation of CS–US association (Fig. 1D,E).
To enable selective recording of SOM⁺ CeL neuron activity, we used Som-Cre mice, in which expression of the Cre recombinase is driven by the endogenous SOM promoter (Taniguchi et al., 2011), in the conditioned lick suppression task. In addition, before the behavioral training, we injected the CeL of these mice with an adeno-associated virus (AAV) expressing GCaMP6f, the genetically encoded Ca²⁺ indicator (Chen et al., 2013), in a Cre-dependent manner (AAV.Flex.GCaMP6f; Fig. 2A–F), followed by implanting an optical fiber above the CeL (Fig. 2A, D). The GCaMP6f signals from SOM⁺ CeL neurons in these mice during fear conditioning were recorded and analyzed. The results showed that SOM⁺ CeL neuron activity increased during the CS period, with a significant rise in fluorescence intensity compared to the baseline (Fig. 2B). This activity was further confirmed histologically, where the expression of GCaMP6f in SOM⁺ CeL neurons was detected based on its intrinsic fluorescent signal (green), and SOM⁺ neurons were identified based on their expression of tdTomato (middle). The GCaMP6f was predominately (>96%) expressed in SOM⁺ neurons (right).

Figure 2. Imaging the activity of SOM⁺ CeL neurons during fear conditioning. A, Schematic of the experimental configuration. B, Quantification of GCaMP6f fluorescence signals recorded from SOM⁺ CeL neurons in a representative mouse, showing average (top) and trial-by-trial (bottom) ΔF/Δt over time. C, Average GCaMP6f fluorescence signals from 8 mice (habituation, baseline vs CS: t(79) = −2.7, p = 0.01, paired t test; recall, baseline vs CS: t(79) = −7.4, p = 2.8E − 11, paired t test; habituation vs recall during CS: t(79) = −6.47, p = 1.17E − 9, t test). Data are represented as mean ± SEM and the shaded lines indicate SEM. D, Drawing of optical fiber placement. Each black dot denotes where the tip of the optical fiber was located in each mouse (n = 8). E, Histology showing expression of GCaMP6f in SOM⁺ CeL neurons in the representative mouse in B. F, Images of a brain section from a Som-Cre;Ai14 mouse in which the CeL was injected with the AAV.Flex.GCaMP6f. The expression of GCaMP6f (left) in CeL neurons was detected based on its intrinsic fluorescent signal (green). SOM⁺ neurons (red) were identified based on their expression of tdTomato (middle). The GCaMP6f was predominately (>96%) expressed in SOM⁺ neurons (right).
the lick suppression task could therefore be recorded using a fiber-optic photometry strategy (Cui et al., 2013; Gunaydin et al., 2014; see Materials and Methods; Fig. 2). To facilitate recording and minimize any motion artifacts, experiments were performed in a head-fixation configuration (Fig. 2A).

Notably, in the mice that had been trained in the conditioned lick suppression task, CS presentations not only induced suppression of the licking behavior (Fig. 1D, E), but also evoked a transient increase in GCaMP6f signals, indicating the activation of SOM⁺ CeL neurons (Fig. 2B, C). These results, together with our previous finding that excitatory synaptic transmission from the BLA onto SOM⁺ CeL neurons is potentiated after fear conditioning (Li et al., 2013), indicate that SOM⁺ CeL neurons become responsive to the threat-predicting sensory stimulus in a learning-dependent manner.

The population average response dynamics of SOM⁺ CeL neurons suggest that they might contribute to the generation of lick suppression. We therefore tested whether activation of SOM⁺ CeL neurons is sufficient to drive lick suppression in unconditioned mice. To this end, we implanted optical fibers bilaterally above the CeL of Som-Cre;Ai32 mice (Madisen et al., 2012),
in which the light-sensitive cation channel channelrhodopsin-2 (ChR2; Zhang et al., 2006) is specifically expressed in SOM+ cells (Li et al., 2013; Fig. 3A–E). Once these mice had been trained for licking, we delivered pulses of blue light into the CeL to activate the ChR2-expressing SOM+ neurons. Remarkably, photostimulation elicited an abrupt and near complete suppression of licking, which was rapidly reversible upon the cessation of light (Fig. 3B, C). Light delivery into the CeL in which SOM+ neurons expressed eYFP did not induce lick suppression (Fig. 3B, C). Therefore, these results demonstrate that SOM+ CeL neuron activation is sufficient to induce lick suppression, mimicking the effect of presenting a CS that predicts a noxious US.

Could SOM+ CeL neurons be necessary for lick suppression? To address this question, we inhibited these neurons while animals were performing the conditioned lick suppression task. For this purpose, we implanted optical fibers bilaterally above the CeL of Som-Cre:Ai35 mice (Madisen et al., 2012) in which the light-sensitive proton pump archaerhodopsin (Arch; Chow et al., 2010) is expressed specifically in SOM+ cells (Li et al., 2013; Fig. 4A–G). After training in the conditioned lick suppression task, these mice were tested in a recall session during which continuous green light was delivered into the CeL in interleaved trials to inhibit the Arch-expressing SOM+ neurons. To assess whether this manipulation could affect lick suppression as well as other forms of defensive behavior, we allowed the mice to run on a wheel throughout the conditioned lick suppression task (Fig. 4A). We found that photoinhibition of SOM+ CeL neurons failed to have an effect on conditioned lick suppression (Fig. 4B, C, D, E). We found that photoinhibition of SOM+ CeL neurons failed to have an effect on conditioned lick suppression (Fig. 4B, C, D, E).
Interestingly, these mice showed CS-induced running behavior in addition to lick suppression and this conditioned running was consistently enhanced by photoinhibition of SOM/CeL neurons (Fig. 4D,E). Light delivery into the CeL of control mice did not affect either lick suppression or running (Fig. 4C,E). Therefore, inhibition of SOM/CeL neurons facilitated CS-induced running, suggesting that the ongoing activity of SOM/CeL neurons normally may suppress active behaviors. By contrast, the same manipulation did not affect conditioned lick suppression; this could be because that the mice were engaged in a different defensive behavior—running—and thus were not able to lick in response to the CS. These results also imply that the SOM/CeL neurons may not be essential for the subjective feeling of fear; rather, they may be important for determining the type of defensive responses, passive or active, to be expressed as the behavioral output.

To further investigate whether SOM/CeL neurons influence active defensive behaviors, we sought to inhibit these neurons using a potent molecular method because optogenetic inhibition suffers from limited light coverage within the CeL that may lead to underestimation of the behavioral effect. We injected the CeL of Som-Cre mice bilaterally with an AAV expressing the tetanus toxin light chain (TeLC; Murray et al., 2011), which inhibits neuronal output by preventing synaptic vesicle fusion, or GFP (as a control) in a Cre-dependent manner (AAV.Flex.TeLC-eGFP or AAV.DIO.eYFP, respectively; Fig. 5A). We subsequently trained these mice in an active avoidance task in which a CS (sound) predicted a noxious US (air puff delivered to the face) and running above a set threshold speed would either prevent or terminate the US (Fig. 5A,B). Compared with the eYFP control mice, the TeLC group showed markedly improved performance in this task (Fig. 5C,D), indicating that the endogenous activity of SOM/CeL neurons acts to repress active defensive behaviors.

Given the observations described above, we reasoned that increasing the activity of SOM/CeL neurons might be sufficient to block active defensive responses. To test this hypothesis, we implanted optical fibers bilaterally above the CeL of Som-Cre;Ai32 mice, which were subsequently trained in the active avoidance task (Figs. 6A, 5B). Once these mice mastered the task, as evidenced by their ability to reduce the frequency of US delivery effectively by running in response to the CS, they were tested for conditioned avoidance. In interleaved trials, SOM/CeL neurons...
Discussion

Recent studies have shown that fear conditioning modifies the response properties of CeL neurons to sensory stimuli. In particular, a population of neurons, the "CeLON" neurons, is activated by the CS in fear-conditioned animals (Ciocchi et al., 2010; Duvarci et al., 2011). However, the identity of these neurons has been unknown. In addition, it is also unclear whether and how the activity of these CeLON neurons is causally related to the generation of defensive behavioral responses.

Our study shows for the first time in behaving animals that SOM⁺ CeL neurons are activated by sensory cues predicting an aversive outcome. This result indicates that SOM⁺ CeL neuron activity is sufficient to convert an active defensive behavior into a passive response.

Figure 6. SOM⁺ CeL neurons inhibit active avoidance. A, Schematic of the experimental configuration. B, Quantification of the effect of photostimulation on running velocity of a representative mouse (comparing trials that had no photostimulation with trials that had photostimulation; 100 trials for each condition). C, Left, Quantification of the effect of photostimulation on running distance (comparing total running distance in response to CS presentations in trials that had no photostimulation with that in trials that had photostimulation; **p = 0.01, t = 5.1, paired t test, n = 4 mice). Right, Quantification of the effect of photostimulation on avoidance (comparing the percentage of successful avoidance in response to CS presentations in trials that had no photostimulation with that in trials that had photostimulation; ***p = 6.6E–5, t = 32.1, paired t test, n = 4 mice). D, Histology results from a representative mouse. E, Drawing of optical fiber placement. Each pair of dots of the same color denotes bilateral placement of the tips of the optical fibers in one mouse (n = 4). In B and C, data are represented as mean ± SEM. The shades along the velocity traces in B indicate the SEM.

were photostimulated with light trains that cover the entire duration of the CS (Figs. 6B, 3). CS presentation triggered robust running behavior in these mice, which was completely abolished by photostimulation of SOM⁺ CeL neurons (Fig. 6B–E). This result indicates that SOM⁺ CeL neuron activity is sufficient to convert an active defensive behavior into a passive response.

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tions (such as freezing or lick suppression), low or no activity in these neurons allows the expression of active responses (such as running or active avoidance). We propose that the activity of SOM⁺ CeL neurons needs to be adjusted flexibly for the expression of adaptive defensive behaviors. For instance, when a threat is close, these neurons need to be highly active to drive freezing or call a halt to all ongoing behavioral activities; whereas when a threat is distant, these neurons should be suppressed, thereby allowing escape or avoidance (Schiff et al., 1962; Fanselow and Lester, 1988; McNaughton and Corr, 2004).

The results showing that inhibition of SOM⁺ CeL neurons does not affect conditioned lick suppression, but rather enhances conditioned running and active avoidance, are particularly interesting, because they suggest that these neurons are likely not essential for producing the negative motivations associated with the US (e.g., the “feeling of fear”). Consistent with this interpretation, although photostimulation of SOM⁺ CeL neurons induced robust lick suppression, the recovery of licking, which was almost instantaneous after the cessation of light (Fig. 3B,C), seems too fast if these neurons had caused any negative motivations (e.g., see the recovery time course of the CS-induced lick suppression in Fig. 1).

The downstream circuitry that mediates the behavioral effects of SOM⁺ CeL neurons is currently unknown. These effects could be mediated by the local inhibitory interactions between SOM⁺ CeL neurons and other CeL populations, including the corticotropin-releasing hormone-expressing neurons (Day et al., 1999) and the protein kinase C-β-expressing neurons (Haubensak et al., 2010), or by the long-range interactions of SOM⁺ CeL neurons with extramygdaloid structures, such as the periaqueductal gray or the paramedian nucleus of the thalamus (Penzo et al., 2014).

References


