LTD at amygdalocortical synapses as a novel mechanism for hedonic learning

Running title: Synaptic mechanisms for CTA learning

Melissa S. Haley, Stephen Bruno, Alfredo Fontanini, and Arianna Maffei

Affiliations. Dept. of Neurobiology and Behavior, SUNY – Stony Brook, Stony Brook, NY

Abstract: 146 words;
Introduction: 484 words;
Discussion: 2004 words
6 Figures; 7 figure supplements
Methods: 2651 words;

Keywords: insular cortex, LTD, learning, plasticity, taste, conditioned taste aversion

Contributions. All authors were involved in writing and commenting the manuscript.
M.S. Haley designed and performed the experiments, analyzed the data and wrote the manuscript.
S. Bruno performed immediate early genes immunohistochemistry and cell counting.
A. Fontanini obtained funding, designed the study and wrote the manuscript.
A. Maffei obtained funding, designed the study, supervised experiments and analysis, and wrote the manuscript.

Send correspondence to: Arianna Maffei, Dept. of Neurobiology and Behavior, Life Science Building Rm 536, SUNY – Stony Brook, Stony Brook, NY 11794-5230. E-mail: Arianna.maffei@stonybrook.edu

Authors’ disclosure: Alfredo Fontanini is a member of the scientific advisory board of Sage Therapeutics, Boston, MA

Acknowledgments: This work was supported by NIH-NIDCD awards DC013770, DC015234 to A. M. and A. F. We wish to thank Dr. Antonello Bonci for useful discussions and feedback on the study.
Abstract

A novel, pleasant taste stimulus becomes aversive if associated with gastric malaise, a form of learning known as conditioned taste aversion (CTA). CTA is common to vertebrates and invertebrates and is an important survival response: eating the wrong food may be deadly. CTA depends on the gustatory portion of the insular cortex (GC) and the basolateral nucleus of the amygdala (BLA) however, its synaptic underpinnings are unknown. Here we report that CTA was associated with decreased expression of immediate early genes in rat GC of both sexes, and with reduced amplitude of BLA-GC synaptic responses, pointing to long term depression (LTD) as a mechanism for learning. Indeed, association of a novel tastant with induction of LTD at the BLA-GC input in vivo was sufficient to change the hedonic value of a taste stimulus. Our results demonstrate a direct role for amygdalocortical LTD in taste aversion learning.
Introduction

Sensory stimuli are perceived and processed according to their physicochemical and affective signatures. For example, taste perception depends on the chemical identity of a tastant, as well as on its pleasantness or aversiveness (i.e., its hedonic value). Hedonic value can be modulated by experience. Indeed, the affective value of a palatable gustatory stimulus changes if it becomes associated with gastric malaise, a process known as conditioned taste aversion (Garcia et al., 1955). In laboratory settings, CTA can be induced in rats by association of sucrose consumption with a gastric malaise-inducing intraperitoneal (ip) injection of lithium chloride. In naïve rats, this association reliably produces a persistent hedonic shift capable of driving avoidance even after a single conditioning trial. Inactivation studies point to the involvement of the gustatory portion of the insular cortex (Braun et al., 1972; Flynn et al., 1991; Yamamoto et al., 1995; Cubero et al., 1999) and the basolateral amygdala (Morris et al., 1999) as key regions necessary for CTA. Evidence from these studies is further corroborated by experiments investigating the interaction between these two regions. In naïve animals, analysis of GC neurons’ activity following inactivation of BLA indicates that amygdalar inputs to GC provide information about the affective dimension of taste stimuli (Piette et al., 2012). Recent studies suggest that the expression of aversive behaviors related to taste depend on the output of GC to the amygdala (Lavi et al., 2018; Schiff et al., 2018; Kayyal et al., 2019).

While the involvement of GC, BLA, and their connections in CTA is well-established, the synaptic changes underlying this form of learning remain unclear. Different experimental approaches produced results that are hard to reconcile. In GC, analysis of molecular markers for synaptic plasticity was brought as evidence that CTA induced long term potentiation (LTP) of GC synapses (Rosenblum et al., 1993; Escobar and Bermudez-Rattoni, 2000; Berman and Dudai,
2001; Escobar et al., 2002; Gal-Ben-Ari and Rosenblum, 2011; Rodriguez-Duran et al., 2011; Guzman-Ramos and Bermudez-Rattoni, 2012). Nevertheless, analysis of GC and BLA neurons’ spiking activity in awake rodents reported a decrease in firing rate in both regions following CTA (Grossman et al., 2008) – a finding that is inconsistent with the hypothesized induction of LTP.

Here we use a combination of immediate early gene immunohistochemistry, patch clamp electrophysiology, and optogenetic stimulation to study plasticity of amygdalocortical circuits in CTA. We demonstrate that CTA reduces activity in GC, and that this decrease is accompanied by long term depression at BLA synapses onto pyramidal neurons in layer 2/3 of GC. Finally, we identify patterns of activity that induce LTD at the BLA-GC input and show that substituting the gastric malaise with this pattern at BLA-GC terminal fields in vivo following sucrose consumption is sufficient to reduce the preference for this tastant. Our results provide the first direct evidence that CTA depends on LTD at amygdalocortical synapses, and that this plasticity alters the affective dimension of a sensory stimulus.

**Results**

In order to investigate whether learning induces synaptic plasticity at amygdalocortical synapses in GC, we trained male and female rats in a conditioned taste aversion paradigm where novel, palatable sucrose (0.1M in water) was paired with gastric malaise induced by an intraperitoneal (ip) injection of lithium chloride (LiCl, 0.15M, 7.5mL/kg body weight; Figure 1A). Animals were placed on a water restriction schedule where they received 15 min access to water from a drinking spout in an experimental chamber in the morning, followed by one-hour access to water in their home cages 4 hours later. After four days of habituation, on day 5 and 7, the morning
water was replaced with 0.1M sucrose, and the animals received an ip injection of LiCl immediately after consumption. Two conditioning sessions were used to ensure learning and consolidation of the aversive memory. In the control group (Figure 1B), pseudo-conditioned animals were exposed to both the US and CS, but they received the LiCl injection the night before sucrose access, and therefore did not form an association between the gastric malaise and tastant. Learning was assessed on day 8 using a 2-bottle test. Average fluid consumption across habituation days for each group was comparable, and animals in both groups initially found the sucrose solution to be palatable, consuming more on the first conditioning day compared to baseline water intake (Figure 1A-B, Figure 1-figure supplement 1). However, this preference shifts in the CTA group following the LiCl injection, as indicated by decreased consumption of sucrose on the second conditioning day, and a strong preference for water at testing (Figure 1A). A significant difference in the aversion index (AI), calculated as \((\text{water} - \text{sucrose})/\text{total volume}\) (Figure 1C; water preferred for \(0 < \text{AI} < 1\); sucrose preferred if \(-1 < \text{AI} < 0\)), could be seen in animals that received an ip injection of LiCl following tastant exposure compared to animals that received LiCl dissociated from the tastant exposure. Following the 2-bottle test, brain tissue was harvested for either immunohistological processing or acute slice recording experiments. To ensure that the 2-bottle test did not initiate CTA extinction, in a subset of rats we repeated the 2-bottle test on two consecutive days (Figure 1-figure supplement 2A). Results from the 2-bottle tests (Figure 1-figure supplement 2C) and aversion indices on test day and 24 hours later were comparable (Figure 1-figure supplement 2D). These results indicate that in our paradigm, CTA was reliably induced, and the 2-bottle test did not initiate an extinction process.

Expression of CTA learning results in decreased recruitment of L2/3 EXC neurons.
To begin assessing the effects of CTA on GC, we quantified neurons expressing the activity-dependent immediate early genes (IEGs) c-Fos and EGR1 one hour after the 2-bottle test (Figure 1D-F). We focused our analysis on pyramidal neurons in layers 2/3 of GC, as our previous work showed that they receive a powerful input from BLA (Haley et al., 2016). We used c-Fos and EGR1 on the same groups of control and CTA rats to cross validate our results and ensure consistency of effect across animals and conditions. There was a significant decrease in the number of c-Fos and EGR1 positive neurons in the CTA group versus control (Figure 1F), indicating that expression of the learned aversion to sucrose decreased GC neuronal activation. This effect is consistent with previous reports showing a reduction in GC spiking activity (Grossman et al., 2008).

**CTA learning does not affect the intrinsic excitability of L2/3 pyramidal neurons in GC but decreases spontaneous synaptic drive.**

The decrease in the number of IEG-labeled neurons following CTA suggests that the associative learning between taste and malaise induced plastic changes in this region of GC, but does not indicate which form of plasticity these changes represent (Inberg et al., 2013; Gandolfi et al., 2017). Indeed, the effect could result from changes to the intrinsic properties of GC neurons, affecting their overall excitability and input/output function. Alternatively, the decrease in expression could be associated with reduced synaptic efficacy, either at specific inputs, such as BLA, or with altered overall synaptic drive onto these neurons.

We first assessed the effect of CTA on the input/output function of pyramidal neurons in L2/3 of GC. Rats were trained on the control or the CTA paradigm (Figure 1A-B) starting at postnatal day 28 (P28). Following the 2-bottle test, acute slice preparations were obtained for
whole-cell patch clamp recordings (Haley et al., 2016). Excitatory neurons were identified online based on their firing response to steady-state current injection, as well as their pyramidal morphology under DIC optics. Recorded neurons were filled with biocytin for post-hoc confirmation of morphology and location, and lack of immunostaining for the GABAergic neuronal marker, GAD-67. All recorded neurons included in the analysis were in L2/3, showed pyramidal morphology, and were negative for GAD67 (Figure 2A). To test for possible changes in input/output function, we quantified the frequency of action potentials as a function of the amplitude of injected steady state current steps (Figure 2B, C; FI curve, increments of 50pA). CTA did not affect the input resistance (IR) of GC neurons and left the input/output curve unaltered, indicating a lack of changes in intrinsic properties (Figure 2C). These data indicate that following CTA expression, the decrease in the number of GC neurons expressing IEGs is not related to learning-dependent plasticity of intrinsic membrane properties.

Next, we examined if the decreased IEG activation following CTA expression could result from changes in synaptic drive. We performed patch clamp recordings in voltage clamp using a cesium based internal solution that allowed for the isolation of spontaneous excitatory (sEPSCs) currents by holding neurons at the reversal potential for chloride (-50mV in our experimental conditions) (Figure 2D-E). Analysis of spontaneous synaptic events unveiled a significant decrease in the cumulative amplitude distribution of spontaneous excitatory synaptic currents, but no change in their frequency (Figure 2E). The decrease in amplitude of spontaneous events was more pronounced for larger events and was sufficient to result in a significant reduction in spontaneous excitatory synaptic charge (Figure 2D). A change in synaptic drive onto these neurons could potentially mask a change in intrinsic membrane properties, so we repeated the experiments in Figure 2C in the presence of ionotropic receptor blockers (20μm DNQX, 50μm...
AP5, 20μm picrotoxin). No differences in intrinsic membrane properties were unveiled following bath application of the synaptic blockers (Figure 2-figure supplement 1). Our results exclude the possibility that CTA altered intrinsic excitability and point to decreased excitatory synaptic transmission as a possible plasticity mechanism for this learning paradigm. Furthermore, given that CTA-induced effects on synaptic drive were detectable in L2/3 pyramidal neurons sampled across the extent of GC, it is highly unlikely for them to depend on selective plasticity on a clustered subset of neurons (Wang et al., 2018b).

CTA learning decreases BLA input onto L2/3 pyramidal neurons in aGC

The reduction in the distribution of large events we observed is consistent with the possibility that CTA may affect afferent inputs onto GC neurons, as these typically have larger amplitudes than recurrent events (Wang et al., 2013; Wang et al., 2018a). Previous work demonstrated that CTA induction is impaired by inactivation of either GC or BLA (Braun et al., 1972; Bermudez-Rattoni and McGaugh, 1991; Morris et al., 1999), suggesting that the connection between these regions may be a fundamental player in this form of learning. We therefore asked whether the BLA-GC input onto pyramidal neurons may be affected by CTA. To do that, we used an optogenetic approach to selectively activate BLA terminal fields while recording pyramidal neurons in the superficial layers of GC. As extensively characterized in our previous study (Haley et al., 2016), we injected AAV9.CAG.ChR2-Venus.WPRE.SV40 (Petreanu et al., 2007) in the BLA (2.1 mm posterior to bregma; 4.7 mm lateral to midline; 7.0 mm below the pia) of P14 rats. CTA training began 14 days after surgery, at P28. The position of the injection site, and expression of the construct in BLA terminal fields within GC were verified histologically (Figure 3A). Lack of retrograde transport of the construct was verified by the lack of backfilled somata.
in GC. Consistency of expression across preparations was assessed as indicated in the Methods (Figure 3B), and in our previous work (Wang et al., 2013; Kloc and Maffei, 2014; Haley et al., 2016; Wang et al., 2018a). Following the 2-bottle test, acute slices were prepared (1.5mm to bregma) and patch clamp recordings were obtained from visually identified L2/3 pyramidal neurons. Brief (5ms) pulses of blue light (470nm) were delivered with an LED mounted on the fluorescence light path of an upright microscope and light-evoked excitatory postsynaptic currents (BLA-EPSCs) were recorded. The monosynaptic nature of BLA-EPSCs was verified as described in our previous work (Haley et al., 2016), and an input/output curve for BLA-EPSCs was obtained using light stimuli of increasing intensity (0.2 – 4.4mW, measured at the output of the objective). As shown in Figure 3C, BLA-EPSC amplitude was reduced following CTA, consistent with the reduction in spontaneous excitatory charge. Recordings in current clamp unveiled that the decreased amplitude of BLA-EPSCs did not reduce the average BLA-evoked EPSP (Figure 3D). However, in both groups, there was a strong correlation with the amplitude of BLA-EPSCs and BLA-EPSPs onto individual cells (Figure 3E). We observed no differences between the control and CTA groups in the proportion of cells that received input from BLA, or in the paired-pulse ratio, coefficient of variation, or decay tau of the evoked BLA-EPSC (Figure 3-figure supplement 1). Since BLA provides a powerful glutamatergic input to L2/3 neurons (Haley et al., 2016), the decrease in BLA-EPSC amplitude strongly suggests that GC neurons are activated less effectively by incoming inputs. This effect may explain the reduced expression of IEGs in Figure 1D, and is consistent with reports of decreased GC neurons’ activity in vivo (Grossman et al., 2008).

**Induction of LTD at BLA-GC synapses is occluded following CTA learning.**
We hypothesized that the decrease in BLA-EPSC amplitude may be evidence for induction of long term depression (LTD) at BLA-GC synapses onto L2/3 neurons. We first probed the capacity for plasticity of these synapses by identifying a pattern of activity that could induce LTD at this input in control animals. We designed induction paradigms that allowed for stimulation of BLA terminal fields with a phasic or a tonic pattern of activity. Both patterns of BLA neuron activity have been recorded in vivo in awake rodents (Fontanini et al., 2009; Parsana et al., 2012) and are thought to be associated with different behavioral states (Parsana et al., 2012). The effect of such activity patterns on spontaneous GC activity was reported in our previous study (Haley et al., 2016). Patch clamp recordings were obtained from L2/3 neurons in GC of slices obtained from control and CTA rats, prepared right after the 2-bottle test. Following acquisition of a 10min baseline recorded in voltage clamp at -70mV, we paired presynaptic 20Hz bursts of light pulses to activate BLA afferents with depolarization of the postsynaptic GC neuron in current clamp, then continued to record neurons in voltage clamp at -70mV for at least 40min post-induction. This phasic induction pattern reliably induced LTD at BLA-GC synapses in 56.52% of the recordings from control rats (Figure 4B). The remaining 43.48% of recordings showed either no change (21.74%) or potentiation (21.74%), indicating that BLA-GC synapses are capable of expressing both synaptic depression and potentiation. LTD induced by phasic BLA terminal field activation did not show changes in paired pulse ratio (PPR) nor in coefficient of variation (CV; Figure 4C), indicative of a postsynaptic site of expression (Malinow and Malenka, 2002; Kessels and Malinow, 2009), and consistent with the decrease in amplitude, but not frequency, of sEPSCs shown in Figure 2E.

We used this induction paradigm to perform an occlusion experiment, testing the possibility that if CTA had saturated LTD at BLA-GC synapses, further LTD induction would result in no
additional depression (Rioult-Pedotti et al., 2000; Maffei et al., 2006; Crozier et al., 2007). LTD induction by phasic patterns of BLA terminal field activity in GC was indeed occluded by CTA (Figure 4E-F): the proportion of recordings showing LTD in response to phasic BLA activity was reduced to 21.74% (from 56.52% in control, Figure 4B), while that of recordings showing LTP increased to 47.83% (from 21.74% in control, Figure 4B). The proportion of recordings in which induction produced no change in synaptic efficacy was not different from control (30.43%, Figure 4B).

LTP induced with the phasic plasticity protocol in the CTA group had no effect on either paired pulse ratio or coefficient of variation, consistent with a postsynaptic site of plasticity expression (Figure 4E). A similar number of cells in the control and CTA groups fired action potentials during the induction protocol, however neither spiking activity during induction, nor the initial PPR, CV, or decay tau, were predictive of the sign of plasticity induced in either group (Figure 4-figure supplement 1B, C). The results of the occlusion experiment strongly suggest that LTD at the BLA-GC synapse is associated with CTA learning.

To assess the specificity of the occlusion, we tested the effect of CTA on LTD induction with a different paradigm designed to mimic tonic BLA activity (Fontanini et al., 2009; Parsana et al., 2012). Here, we paired presynaptic ramping light stimuli to tonically activate BLA afferents (Haley et al., 2016), with depolarization of the postsynaptic GC neuron. LTD induction with the tonic BLA activity regime was not occluded by CTA, resulting in LTD in 100% of the recordings in both control and CTA rats (Figure 5A-F). This form of LTD was accompanied by a significant reduction in paired-pulse ratio in the control group (Figure 5C), and a significant change in both paired-pulse ratio and coefficient of variation in the CTA group (Figure 5E).
This decrease in PPR is consistent with a presynaptic site of plasticity expression, indicating that tonic BLA activity engages a distinct set of plasticity mechanisms that are not consistent with the changes in sEPSC amplitude we observed following CTA. Similar to the 20Hz protocol, neither spiking during induction, nor initial PPR, CV, or decay tau, were correlated with the degree of plasticity (Figure 5-figure supplement 1B, C). These results strongly suggest that successful CTA learning is associated with postsynaptically expressed LTD induced by phasic activation of BLA afferents in GC.

Phasic activation of BLA terminal fields in GC in vivo eliminated sucrose preference.

In view of the selective occlusion of LTD induced by phasic BLA terminal field activation, we asked whether association of novel sucrose and phasic activation of BLA terminal fields in GC would be sufficient to change the hedonic value of the tastant. To do that, we injected the AAV9.CAG.ChR2-Venus.WPRE.SV40 construct (Petreanu et al., 2007) in BLA and implanted an optic fiber (400 μm) in GC (paired 20Hz opto; Figure 6A). As control, a group of animals was injected with an AAV9.hSyn.eGFP.WPRE.bGH construct (University of Pennsylvania Vector Core) and implanted with the optic fiber in GC (GFP control). These two rat groups were trained on a modified CTA paradigm in which the LiCl-induced malaise was substituted with phasic activation of BLA terminal fields (Figure 6C, D). A third group of rats was injected with ChR2 and received non-contingent phasic activation of BLA terminal fields the evening before sucrose exposure (non-paired 20Hz opto, Figure 6E). Finally, a fourth group of rats was injected with ChR2 and the LiCl-induced malaise was replaced by tonic activation of BLA terminal fields with the ramping light stimulus (Figure 6F). All groups showed a comparable initial preference for sucrose over water, including the non-paired 20Hz opto group, indicating that 20Hz stimulation
alone did not affect sucrose detection or palatability (Figure 6-supplemental figure 1). However, as shown in Figure 6D, only the rat group that received phasic light activation of BLA terminal fields in GC paired with sucrose showed a reduced preference for sucrose. This effect is further supported by the change in AI (Figure 6G). Thus, while both the phasic 20Hz and the ramp tonic paradigm induced LTD in acute slice preparations, CTA was selectively induced by pairing the phasic 20Hz pattern of activation of BLA afferents in GC with sucrose consumption.

Discussion

Our data demonstrate that CTA learning leads to reduced activation of the taste responsive portion of the insular cortex and to an LTD-related reduction in strength of the BLA-GC connection. The results also demonstrate that induction of LTD at amygdalocortical synapses onto L2/3 pyramidal neurons with a phasic 20Hz pattern of stimulation of BLA terminal fields in GC is sufficient for changing the hedonic value of a taste stimulus. These findings confirm the central role of the projection from BLA to GC in CTA, and also reveal that circuit changes following CTA learning do not depend, as previously thought, on strengthening of synaptic connections, but rely on a significant decrease in synaptic input. Previous work reported a substantial change in the cortical representation of the conditioned tastant in GC following aversive learning (Yasoshima and Yamamoto, 1998; Accolla et al., 2007; Grossman et al., 2008). Our work provides novel evidence regarding the circuit mechanisms underlying this reorganization process and identifies long term plastic changes associated with CTA.

CTA and immediate early gene expression
The behavioral paradigm we used consisted of two rounds of conditioning with taste-malaise pairing to ensure reliable expression of aversion learning. As a readout of the aversive memory, we used a 2-bottle test to determine volume consumption of the conditioned stimulus following learning, and to compute the aversion index between control and CTA animals. The 2-bottle test did not initiate an extinction process, as we show that the aversion index remained stable when animals were tested on two consecutive days. Quantification of IEGs, as a proxy for GC neurons’ activity, showed reduced expression of both c-Fos and EGR1 following testing in CTA animals compared to control. The decrease in IEG expression provides additional support as to the lack of extinction initiation, as extinction has been shown to increase IEG expression in GC (Hadamitzky et al., 2015).

In our paradigm, all animals, including controls, are exposed to sucrose twice. While for CTA animals sucrose is always associated with a gastric malaise, in control animals sucrose is a safe, palatable stimulus and animals consume it more than water. In control animals, the consumption of sucrose on the second exposure is higher than the first, consistent with an attenuation of neophobia (Lin and Reilly, 2012), an effect that is associated with a reduction in c-Fos expression in GC (Lin et al., 2012). In the CTA group, the consumption of sucrose at the second exposure was significantly lower than at the first. The differences in IEG expression we report may reflect a blunted GC activation to sucrose in CTA animals compared to control. Consistent with this possibility, previous studies showed increased IEG expression in the insular cortex following consumption of a novel tastant alone (saccharine), but not in response to a LiCl injection alone (Wilkins and Bernstein 2006; Bernstein and Koh, 2007). Thus, our data support the interpretation that the differences in IEG expression are due to animals in the CTA group expressing an aversion to sucrose, rather than the control group learning that sucrose is “safe”.
As IEG expression is considered a proxy for activity, our results are consistent with previously published work showing reduced spiking activity of GC neurons in the palatability epoch following CTA (Grossman et al., 2008; Piette et al., 2012). Thus, the aversive memory induced by our behavioral paradigm was reliable, allowing us to assess the effects of CTA on the BLA-GC circuit.

### CTA reduced excitatory synaptic drive onto GC neurons

Our analysis of synaptic and intrinsic properties of GC neurons allowed us to parse apart which of these components was responsible for the reduction in GC activity seen with CTA. We show that excitatory spontaneous synaptic charge onto GC pyramidal neurons is reduced following CTA, an effect that points to widespread modulation of synaptic transmission in GC, indicating that CTA affects the circuit in GC broadly. CTA altered synaptic drive and capacity for plasticity at BLA-GC synapses, effects consistent with changes in the activation of GC. These results strongly suggest that decreased synaptic drive onto GC neurons underlies the CTA-dependent circuit reorganization reported in previous work (Accolla and Carleton, 2008) and explains the reduced firing rates of GC neurons following CTA (Grossman et al., 2008).

Analysis of spontaneous excitatory synaptic events revealed a significant decrease in amplitude, and no changes in frequency. These effects are consistent with a postsynaptic site of expression of CTA-dependent modifications of synaptic efficacy (Malinow and Malenka, 2002; Rumpel et al., 2005). Further confirmation of the postsynaptic site of expression comes from analysis of BLA-evoked responses onto GC pyramidal neurons. Following CTA, we observed a significant decrease in the amplitude of BLA-evoked responses, but no change in their paired pulse ratio. Furthermore, the lack of changes in intrinsic excitability indicate that the profound effects taste
aversion learning has on GC pyramidal neurons is via modulations at specific synaptic inputs and not an alteration in intrinsic membrane properties. Together, this set of findings confirms the central role of the BLA-GC connection in CTA learning, and demonstrates a role for plasticity of BLA-evoked synaptic responses onto L2/3 pyramidal neurons in hedonic learning.

**Postsynaptically expressed LTD at amygdalocortical synapses as a mechanism for CTA**

When we compared the effect of CTA on two different forms of LTD, induced by patterns of stimulation designed to mimic BLA neurons’ activity in different behavioral states (Fontanini et al., 2009; Parsana et al., 2012), we found a selective occlusion of plasticity induced by a phasic pattern of activity. Phasic stimulation induced a postsynaptically expressed form of LTD in control rats, but LTP in the CTA group, suggesting that phasic LTD and CTA share common mechanisms. The occlusion was specific to the phasic induction pattern, as LTD induced with a tonic pattern of BLA terminal field activation, a form of LTD with a presynaptic site of expression, was not affected by CTA. These results point to postsynaptic LTD at BLA-GC synapses as a mechanism for taste aversion learning.

The underlying mechanisms of postsynaptically expressed LTP and LTD have been studied extensively. Signaling cascades leading to the regulation of AMPA receptors in the postsynaptic membrane are thought to mediate changes in synaptic efficacy (Malinow and Malenka, 2002; Kessels and Malinow, 2009). Current thinking associates learning with an increase in synaptic efficacy (Bliss and Collingridge, 1993; Rogan et al., 1997; Dityatev and Bolshakov, 2005), and changes in molecular markers for plasticity following CTA have been interpreted as indication of LTP expression (Shema et al., 2007; Gal-Ben-Ari and Rosenblum, 2011). However, many of the molecular mechanisms involved in CTA (Berman and Dudai, 2001; Gal-Ben-Ari and
Rosenblum, 2011; Guzman-Ramos and Bermudez-Rattoni, 2012) can be engaged by both LTP and LTD (Yilmaz-Rastoder et al., 2011; Gandolfi et al., 2017; Li and Pozzo-Miller, 2019). Additional evidence in support of the interpretation that CTA should potentiate BLA input to GC came from studies based on BLA activation with drug infusions or tetanic electrical stimulation during CTA learning. Local field potential (LFP) recordings in GC of anesthetized rodents showed that the induction of LTP of the population signal evoked by tetanic stimulation of the amygdala was occluded by CTA (Escobar and Bermudez-Rattoni, 2000; Escobar et al., 2002), suggesting that plasticity at the BLA input to GC may play a role in this learning paradigm. However, this experimental design does not assess whether the baseline response has been affected by CTA, nor which component of the population signal has been modified by CTA. Furthermore, these data do not indicate whether LTP at BLA-GC synapses may be sufficient for learning.

A key feature of our study that distinguishes it from previous work is the use of optogenetic techniques to selectively activate BLA terminal fields within GC. This approach prevented confounds due to recruitment BLA collaterals to other areas which inevitably occurs with somatic activation of BLA neurons (Klavir et al., 2017). Furthermore, our study is the first to examine baseline synaptic transmission, and therefore may yield different results from field potential recordings, which lack synaptic resolution, and are difficult to interpret as the circuit components that contribute to the LFP trace in GC have not been described (Escobar and Bermudez-Rattoni, 2000; Escobar et al., 2002; Rodriguez-Duran et al., 2011). No analysis of changes in synaptic efficacy has been reported up to the present work. In view of our results demonstrating a reduction in the efficacy of BLA glutamatergic synapses onto GC neurons...
following CTA, it is likely that in L2/3 pyramidal neurons, signaling mechanisms for LTD are
recruited during CTA.

LTD at amygdalocortical synapses in GC is sufficient for the induction of CTA

Association of sucrose and phasic stimulation of BLA terminal fields in GC *in vivo* was
sufficient to induce a weak CTA, unveiled as a loss of preferences for sucrose over water during
the 2-bottle test. In these experiments, the phasic pattern of stimulation was delivered after
sucrose in place of LiCl, thus selectively activating the BLA-GC input in the absence of a gastric
malaise. Our results indicate that the pattern of activity for phasic 20Hz LTD at BLA-GC
synapses was sufficient to change the hedonic value of sucrose. The effect was specific to the
pairing of sucrose and phasic 20Hz stimulation of BLA terminal fields in GC, as neither non-
contingent stimulation, nor the tonic ramping stimulus affected sucrose preference. While phasic
20Hz activation of BLA-GC terminal fields *in vivo* was sufficient for changing the valence of
sucrose, it did not induce as strong an aversion as LiCl. This effect was expected, as LiCl induces
a gastric malaise that is not recruited in animals that received light stimulation through the optic
fiber during CTA training. GC receives extensive visceral input (Cechetto and Saper, 1987)
which is not activated during BLA terminal field stimulation. Furthermore, previous studies
have shown no change in IEG expression in the insular cortex following LiCl-induced illness,
suggesting that processing of information about the US primarily occurs in other parts of the
central gustatory system (Wilkins and Bernstein 2006; Bernstein and Koh, 2007). Our findings
therefore suggest that the plasticity of BLA-GC input primarily modulates the valence of a
tastant, and that integration of other sources of information about the US may be necessary for
eliciting a strong aversion.
Recent work has identified a role for corticoamygdala projecting neurons in CTA (Lavi et al., 2018; Kayyal et al., 2019), suggesting that this population of GC neurons is involved in the acquisition and retrieval of the aversive taste memory. Whether the population of amygdala-projecting and amygdala-recipient neurons in GC overlap is currently unknown, but this study raises interesting questions about how and when reciprocal connections between BLA and GC are recruited in CTA. It is also unclear whether amygdalocortical neurons in GC project exclusively to BLA, or also interact with other nodes of the CTA circuit.

**Conclusion**

LTD was proposed as a mechanism for learning in cerebellar circuits (Ito et al., 2014; Hirano et al., 2016) and is thought to contribute to the control of alcohol seeking behavior in the basal ganglia (Ma et al., 2018). In both circuits, LTD is induced at glutamatergic synapses onto GABAergic neurons (Hansel and Linden, 2000; Kakegawa et al., 2018), providing a form of disinhibition to the downstream synaptic targets. Our study focused on plasticity of BLA input onto excitatory neurons in L2/3 of GC, strongly supporting a role for LTD at glutamatergic synapses onto pyramidal neurons in hedonic learning.

To our knowledge, our results are the first to directly assess the effect of CTA on BLA-GC synapses, and provide the first evidence for a direct link between LTD of amygdalocortical transmission onto GC neurons and taste aversion learning. Taste circuits are highly conserved among species, and changes in taste preference can be induced similarly in humans as well as many other organisms, making CTA and the amygdalocortical system ideal models for investigating neural mechanisms of hedonic learning. Our results link a specific form of LTD at the BLA-GC input to a change in taste preference, emphasizing how modulation of synaptic
efficacy underlie adaptive shifts in hedonic value capable of influencing perception and behavior.

**Methods**

Long Evans rats of both sexes were used for this study. Animals were singly housed in a vivarium on a 12h/12h light dark cycle with ad libitum access to food and water, except where otherwise noted. Experiments were conducted during the light period. All surgical and experimental procedures were approved by the Institutional Animal Care and Use Committee of Stony Brook University and followed the guidelines of the National Institutes of Health.

*Conditioned taste aversion training.* Rats were placed on water restriction with free access to food for a total of 8 days. Rats were habituated to a behavioral chamber where they had 15min access to a drinking spout with H₂O, followed by 1hour access to H₂O in their home cage four hours later. The volume consumed was recorded daily throughout training (total volume (ml): juveniles, 12.75 ± 0.70; adults, 16.96 ± 1.17), and rats’ weight was monitored to ensure that it remained within 85% of initial weight. Four days of habituation training was sufficient to stabilize fluid intake levels (Habituation, Figure 1B, C). This was followed by two conditioning trials, with a recovery day in between identical to the habituation days. The recovery day enabled us to confirm that the conditioning procedures did not affect thirst levels. For immunohistochemistry and slice electrophysiology, conditioning consisted of 15 min access to a drinking spout with 0.1M sucrose, followed by an ip injection of 0.15M LiCl (7.5mL/kg) to induce gastric malaise. Rats in the control group received an ip LiCl injection (0.15M) in the evening on day 4 and 6 of training. The injection was delivered approximately 16h before sucrose presentation to ensure lack of association between gastric malaise and sucrose
consumption (Figure 1B, pseudo CTA). For experiments in vivo, conditioning for rats in the GFP
test group (Figure 6C) and paired 20Hz opto group (Figure 6D) consisted of 15min access to
a drinking spout with 0.1M sucrose, followed by optogenetic 20Hz stimulation of BLA terminal
fields in GC. Rats in the non-paired 20Hz opto group (Figure 6E) received 20Hz stimulation of
BLA terminal fields in GC the evening prior to sucrose exposure. Rats in the ramp opto group
(Figure 6F) had sucrose exposure followed by optogenetic ramping stimulation of BLA terminal
fields in GC. On the 8th training day, all groups engaged in a 2-bottle test to assess a preference
for H2O or sucrose. In a subset of experiments, the 2-bottle test was repeated on day 9, to assess
whether the first test had initiated an extinction process (Figure 1-figure supplement 2).

Immunohistochemistry. Detection of immediate early gene expression - One hour after the 2-
bottle test, rats were anesthetized and intracardially perfused with PBS followed by 4%
paraformaldehyde in PBS (4% PFA). The brain was dissected out and thin (50µm) coronal slices
containing GC were cut with a fixed tissue vibratome (Leica VT1000). Sections were washed in
PBS (3 x 10min rinse), permeabilized and blocked in a solution containing 0.5% Triton X and
10% normal goat serum in PBS for 1h, then incubated overnight at 4°C in a solution containing
0.1% Triton X and 3% normal goat serum in PBS, mouse anti-GAD67 (1:500, MilliporeSigma,
MAB5406, monoclonal), and either rabbit anti-c-Fos (1:500, Cell Signaling, 2250S, monoclonal)
or rabbit anti-EGR1 (1:500, Cell Signaling, 4153S, monoclonal). Sections were then rinsed in
PBS (3 x 10min) and incubated at 25°C for 2h in a solution containing 0.1% Triton X and 3%
normal goat serum in PBS, goat anti-mouse Alexa Fluor-647 (1:500, Invitrogen, A21235), goat
anti-rabbit Alexa Fluor-568 (1:500, Invitrogen, A11011), and NeuroTrace 435/455 (1:1000,
Invitrogen, N21479). After a final wash in PBS (3 x 10min), sections were mounted with
Fluoromount-G (Southern Biotech). Sections were imaged with a confocal microscope (Olympus
Fluoview) at 20x magnification. Four sections, spaced at 200µm, were counted per animal for each IEG (c-Fos = 4, EGR1 = 4) using ImageJ software by a person blind to experimental condition. Cell counts represent the number of IEG-positive, GAD67-negative neurons, normalized by the counting area (L2/3 aGC) for each section.

Recovery of recorded neurons in acute slice preparation - Recorded slices were fixed in 4% PFA for 1 week. They were then washed in PBS (3 x 10min rinse), permeabilized and blocked in a solution containing 1% Triton X and 10% normal goat serum in PBS for 2h, then slices were incubated overnight at 4°C in a solution containing 0.1% Triton X and 3% normal goat serum in PBS, streptavidin Alexa Fluor-568 conjugate (1:2000, Invitrogen, S11226), mouse anti-GAD67 (1:500, MilliporeSigma, MAB5406, monoclonal), and chicken anti-GFP (1:1000, Abcam, ab13970, polyclonal). GC slices were then rinsed in PBS (3 x 10 min) and incubated at 25°C for 2h in a solution containing 0.1% Triton X and 3% normal goat serum in PBS, goat anti-mouse Alexa Fluor-647 (1:500, Invitrogen, A21235), goat anti-chicken Alexa Fluor-488 (1:500, Abcam, ab150173), and Hoechst 33342 (1:5000, Invitrogen, H3570). BLA slices were incubated overnight at 4°C in a solution containing 0.1% Triton X and 3% normal goat serum in PBS and chicken anti-GFP (1:1000, Abcam, ab13970, polyclonal). They were then rinsed in PBS (3 x 10 min) and incubated at 25°C for 2h in a solution containing 0.1% Triton X and 3% normal goat serum in PBS, goat anti-chicken Alexa Fluor-488 (1:500, Abcam, ab150173), and Hoechst 33342 (1:5000, Invitrogen, H3570). Validation of injection sites and positioning of fiber optics - Following behavioral training, animals were anesthetized and perfused intracardially with PBS followed by 4% PFA. Coronal slices containing BLA and GC were cut at 100µm with a fixed tissue vibratome (Leica VT1000). All sections were washed in PBS (3 x 10 min rinse) and incubated at 25°C for 1h in a solution containing 0.1% Triton X and Hoechst 33342 (1:5000,
Invitrogen, H3570). Sections were mounted with Fluoromount-G (Southern Biotech) and the accuracy of viral injections and fiber placements was assessed.

**Electrophysiology.** Acute coronal slices containing GC were obtained immediately following the 2-bottle test. Slice preparation was as described in our previous study (Haley et al., 2016). Briefly, rats were anesthetized with isoflurane (Bell jar to effect) and rapidly decapitated. The brain was dissected in ice cold, oxygenated artificial cerebrospinal fluid (ACSF), and 300µm coronal slices containing GC were prepared using a fresh tissue vibratome (Leica VT1000) starting at 1.5mm to bregma. Patch clamp recordings were obtained from visually identified L2/3 pyramidal neurons under DIC optics. Their identity was tested online with the injection of square current pulses (700ms) to assess regular firing patterns, and post-hoc with immunohistochemistry aimed at reconstructing neuron morphology, assessing location, and confirming lack of expression of the GABA neuron marker GAD67. To assess possible changes in intrinsic excitability induced by CTA, recordings were obtained in current clamp and square current pulses of increasing amplitude were injected to examine input resistance, action potential threshold, and frequency/intensity curves. Input resistance was calculated from the linear portion of the voltage response to a -50pA current injection. Action potential threshold was calculated as the membrane voltage when the first derivative of the voltage trace $dV/dt = 20V/s$ (Bekkers and Delaney, 2001). In a subset of experiments intrinsic properties were assessed in current clamp in the presence of fast synaptic receptor blockers (in µM: DNQX, 20; APV, 50; picrotoxin, 20). In a different set of experiments, spontaneous synaptic activity was recorded in voltage clamp. For these experiments, a cesium sulfate-based internal solution containing the sodium channel blocker QX314 (2mM, Tocris Bioscience) was used to stabilize recordings during prolonged depolarization. Spontaneous excitatory synaptic currents were recorded at three different holding
potentials around the reversal potential for chloride (to record sEPSCs, in mV: -55, -50, -45). Current versus voltage functions were used to identify the voltage that better isolated the current of interest, which were used for analysis of charge and spontaneous events’ amplitude and frequency (Maffei et al., 2004; Haley et al., 2016). Total charge was calculated by integrating 5s sections of the current trace. Evoked EPSCs/EPSPs from stimulation of BLA terminal fields were obtained in voltage clamp and current clamp using a potassium gluconate solution and holding neurons at -70mV. The monosynaptic nature of BLA-EPSCs was verified as in our previous study (Haley et al., 2016). Event-triggered average of light-evoked BLA-EPSCs and BLA-EPSPs was used to align BLA-EPSC and BLA-EPSP onset and calculate the average amplitude of the light-evoked response. Latencies of BLA-evoked responses were calculated from the onset of the phasic 5ms light pulse. For plasticity experiments, 2 LED pulses (5ms, 10Hz) were used to elicit BLA-EPSCs every 30s. A brief (10ms) 1mV depolarization step was used to monitor series resistance (R_s). After a 10min baseline, either the phasic or tonic induction paradigm was applied in current clamp, after which BLA-EPSCs were recorded for an additional 40min. Neurons with R_s > 20 MΩ or that changed >15% during recording were excluded from analysis.

Optogenetics. Ex vivo experiments: Rats (P14) were anesthetized intraperitoneally with a mixture containing 70mg/kg ketamine, 0.7mg/kg acepromazine, and 3.5mg/kg xylazine (KXA) and mounted on a stereotaxic apparatus. Animals received an injection of the AAV9.CAG.ChR2-Venus.WPRE.SV40 construct ((Petreanu et al., 2007); University of Pennsylvania Vector Core) in the BLA using a nanoject pressure injector (Drummond Nanoject II; 100nL volume containing 5.6412 particles/mL) as described in our previous study (Haley et al., 2016). The stereotaxic coordinates for the injections were 2.1mm posterior to bregma; 4.7mm lateral to midline; 7.0mm
below the pia. Rats were allowed to recover from surgery for 2 weeks before CTA training and recordings. AAV9 was chosen because it is primarily transported anterogradely, and because ChR2 expressed via AAV9 does not alter the short-term dynamics of evoked responses (Jackman et al., 2014). The lack of backfilled somata in all of our GC preparations further confirms forward direction of transport and indicates that only BLA terminal fields in GC were activated by light pulses. Consistency of ChR2-Venus expression across preparation was as previously reported (Haley et al., 2016). Briefly, as shown in Figure 3B, the intensity of the Venus signal in GC was quantified across a 500μm wide region of interest (ROI) spanning the cortical mantle from the pia to L5/6 (included) across 10 preparations. The average and standard deviation of the fluorescent signal were plotted as a function of depth. The expression of the Venus signal was then assessed for all preparations used for electrophysiological recordings and compared to the calibration curve. Only data obtained from preparations with a fluorescence profile within 1 standard deviation of the average calibration curve were included in the analysis. BLA-EPSCs were evoked with a 5ms light pulse delivered through the 40x objective mounted on an upright microscope (Olympus BX51WI). The intensity, frequency, and timing of the blue light stimuli (470nm) were controlled with an LED driver (ThorLabs). Intensity at the output was verified with a light meter (Thor Labs, Range: 0.2 - 4.4mW). Phasic stimulation for plasticity induction was organized in 20 trains of 20-5ms light pulses at 20 Hz delivered every 250ms and paired with depolarization of the postsynaptic neuron with a 50pA current step. Tonic activation of BLA terminal fields was achieved using a ramping light stimulus (which prevented inactivation of ChR2 channels (Lin et al., 2009)). The duration of the ramping stimulus was 6s from 0 to maximum (4.4mW) light intensity and was controlled through the LED driver...
connected to an analog output of the amplifier. For plasticity induction, 10 ramping stimuli spaced 5s were paired with depolarization of the postsynaptic neuron with a 50pA current step.

*In vivo* experiments: Stimulation of BLA terminal fields *in vivo* was achieved through optical fibers (Doric lenses, 400μm) chronically implanted in GC (1.4mm anterior to bregma; 5mm lateral to midline; 4.5mm below the pia). Rats, weighing > 250g were anesthetized intraperitoneally with the KXA cocktail described above and supplemented with 40% of the induction dose as needed to maintain surgical levels of anesthesia. The scalp was exposed and holes were drilled for anchoring screws, optical fibers, and viral injection. The AAV9.CAG.ChR2-Venus.WPRE.SV40 construct (Petreanu et al., 2007) was bilaterally injected in the BLA (3.0mm posterior to bregma; 5mm lateral to midline; 7.2mm below the pia) using a nanoject pressure injector (Drummond Nanoject II; 200nL volume containing 5.6412 particles/mL) as described above. Optical fibers (Doric Lenses, 400μm) were coated with DiI for better detection of the optic fiber tract and implanted bilaterally to target GC and were secured in place with dental cement.

For these experiments, recovery from surgery was 4 weeks prior to CTA training. The injection site in BLA and the positioning of the optic fiber in GC were verified histologically for each preparation. Stimulation was controlled via the digital output (phasic protocol) or analog output (ramp protocol) of a Multi-patch clamp amplifier (HEKA) and software (Patchmaster) and delivered through the implanted optical fibers coupled to a blue laser (Shanghai Dream, 470nm). A control group of rats received a BLA injection of the AAV9.hSyn.eGFP.WPRE.bGH construct (University of Pennsylvania Vector Core) to ensure that possible changes in sucrose preference were due to BLA terminal field stimulation and not to unspecific effects of the light stimulation. Light intensity at the tip of the optic fiber was measured with a light meter (6.5mW, ThorLabs)
The induction paradigm \textit{in vivo} had the same structure as the plasticity induction pattern used in slices (Phasic: Trains of 20 pulses, in this case 10ms long, at 20Hz delivered every 250ms, repeated 100 times; Ramp: 12 ramping stimuli 6s long, spaced 5s). Animals in the GFP control group, paired 20Hz opto group, and ramp opto group received photostimulation immediately after exposure to the 0.1M sucrose solution on conditioning days 1 and 2. The non-paired 20Hz opto group received photostimulation the evening prior to sucrose exposure.

\textit{Data analysis.} Data were acquired with a HEKA 4 channels amplifier with integrated acquisition board. Data were sampled at 10kHz. Patchmaster (WaveMetrix) was used as acquisition software. Analysis was performed with custom procedures in Igor (WaveMetrix), Matlab, and Clampfit (Molecular Devices). For two-group comparisons (paired and unpaired), we used estimation statistics (Ho et al., 2019) to report the effect size and calculate the 95% confidence interval of the mean difference via bootstrap resampling (5000 resamples). Graphic visualization of the data includes the group mean and 95% confidence interval, swarm plots of individual data values, and histograms of the resampling distribution of the difference in the means. Multiple comparisons were assessed with One-Way ANOVA followed by post-hoc Tukey HSD post hoc tests. Cumulative distributions were compared using the 2-sample Kolmogorov-Smirnov (K-S) test. Rank order correlation analysis was used to determine correlations between IEG expression and sucrose consumption (Figure 1), BLA-EPSC/BLA-EPSP amplitude (Figure 3E, input/output function), and plasticity and PPR, CV, or Decay tau (Figure 4-figure supplement 1, Figure 5-figure supplement 1). For plasticity induction experiments, differences between BLA-EPSC amplitude, PPR, and CV were assessed in the 10min baseline period and in the 10min period 30-40min post-induction. Differences in probability of LTD vs LTP induction following CTA were assessed with a $3 \times 2 \chi^2$ for contingency test. Significant differences were considered for $p < 0.05$. 
Solutions. ACSF contained the following (mM): 126 NaCl, 3 KCl, 25 NaHCO3, 1 NaHPO4, 2 MgSO4, 2 CaCl2, 14 dextrose with an osmolarity of 315-325mOsm. The internal solution for analysis of excitability and evoked synaptic responses was as follows (mM): 116 K-Glu, 4 KCl, 10 K-HEPES, 4 Mg-ATP, 0.3 Na-GTP, 10 Na-phosphocreatine, 0.4% biocytin (V_{rev} [Cl^{-}] = −81mV). The pH of the internal solution was adjusted to 7.35 with KOH and the osmolarity adjusted to 295mOsm with sucrose. The internal solution to assess spontaneous excitatory and inhibitory charge and events contained (mM): 20 KCl, 100 Cs-sulfate, 10 K-HEPES, 4 Mg-ATP, 0.3 Na-GTP, 10 Na-phosphocreatine, 3 QX-314 (Tocris Bioscience), 0.2% biocytin (V_{rev} [Cl^{-}] = −49.3mV). Drugs. To assess the monosynaptic nature of BLA-EPSCs recordings were obtained in the presence of (in µM): 1 TTX (Tocris Bioscience), 100 4-aminopyridine (4-AP) (Tocris Bioscience). To determine possible changes in intrinsic excitability, the following blockers were bath applied (in uM): 20 DNQX (Tocris Bioscience), and 50 APV (Tocris Bioscience), 20 picrotoxin (Tocris Bioscience).

Figure Legends

Figure 1: Conditioned taste aversion learning differentially recruits EXC neurons in L2/3 aGC. (A) Behavioral paradigm schematic. Animals in the control group (N = 30) received ip LiCl (7.5mL/kg of 0.15M LiCl) the evening prior to exposure to a sucrose solution. Animals in the CTA group (N = 28) received ip LiCl after sucrose exposure. (B) Fluid consumption across training days for each group. Rats in both groups showed an initial preference for sucrose over water which shifts after conditioning (Control: C1 sucrose 6.66 ± 0.63mL vs. C2 sucrose 7.75 ± 0.70mL, p < 10^{-3}; CTA: C1 sucrose 6.93 ± 0.61mL vs. C2 sucrose 5.06 ± 0.53mL, p < 10^{-7}) and at testing (2-bottle test, Control: sucrose 7.72 ± 0.64 mL vs. H2O 1.52 ± 0.45 mL, p < 10^{-14}; CTA: sucrose: 0.66 ± 0.22 mL vs. H2O 4.79 ± 0.38 mL, p < 10^{-16}). (C) Aversion index on test day (Control -0.68 ± 0.08 vs. CTA 0.76 ± 0.07, p < 10^{-32}). (D) Representative image depicting region of interest (ROI, L2/3 aGC, white box). Scale bar = 500 µm. Red: immediate early gene;
blue: Nissl counterstain. For details on imaging procedures see Methods. (E) Representative images of c-Fos expression (top) and EGR1 expression (bottom) following 2-bottle testing from control group (left) and CTA group (right). Scale bar = 100 μm. (F) Average counts of c-Fos expressing and EGR1 expressing neurons normalized to the area of the ROI. (c-Fos [N = 6]: Control 128.13 ± 9.46 positive nuclei/0.1mm² vs. CTA 82.21 ± 11.73, p < 0.01; EGR1 [N = 6], control: 175.90 ± 15.65 vs. CTA: 133.03 ± 10.10, p < 0.05). IEG expression was not significantly correlated with sucrose consumption for any group (Control: c-Fos, r_s = 0.60, p = 0.21; EGR1, r_s = 0.43, p = 0.40; CTA: c-Fos, r_s = -0.37, p = 0.47; EGR1, r_s = 0.60, p = 0.21). * indicates p ≤ 0.05. Error bars ± 95% CI.

Figure 1-figure supplement 1: (A) Average water consumption prior to conditioning did not differ between groups and rats in both groups showed an initial preference for sucrose over water (Control, [N = 30]: average H2O 4.42 ± 0.32mL/day vs. C1 sucrose 6.66 ± 0.63mL, p < 10^-9; CTA, [N = 28]: average H2O 4.24 ± 0.29mL/day vs. C1 sucrose 6.93 ± 0.61mL, p < 10^-10. Control H2O vs. CTA H2O, p = 0.43). * indicates p ≤ 0.05. Error bars ± 95% CI.

Figure 1-figure supplement 2: Stability of CTA learning across testing days. (A) Diagram of behavioral paradigm to assess whether the 2-bottle test represents an extinction trial. (B) Average fluid consumption across training sessions (Extinction testing, [N = 8 rats]: average H2O 4.77 ± 0.74mL/day vs. C1 sucrose 7.70 ± 0.99mL, p < 0.01; C1 sucrose 7.70 ± 0.99mL vs. C2 sucrose 4.84 ± 1.38mL, p < 0.01). (C) Results from 2-bottle tests on two consecutive days (Test A: sucrose 0.54 ± 0.29mL vs. H2O 5.41 ± 1.75mL, p < 10^-4; Test B: sucrose 0.47 ± 0.26mL vs. H2O 4.90 ± 2.00mL, p < 10^-3). (D) Aversion indices on test day and 24 hours later (AI: test A 0.77 ± 0.15 vs. test B 0.74 ± 0.16, p = 0.74). * indicates p ≤ 0.05. Error bars ± 95% CI.

Figure 2: CTA learning does not affect intrinsic excitability but reduces excitatory synaptic drive onto EXC neurons in L2/3 aGC. (A) Images of example recorded neurons from a control animal (top) and CTA animal (bottom). Left: biocytin (green), middle: GAD-67 (magenta), right: Merge showing lack of co-localization. Scale bar = 25μm. (B) Steady-state firing patterns of neurons from a control animal (top) and CTA animal (bottom). (C) FI curves from control (N = 15 rats, 3-4 cells/animal) and CTA animals (N = 14 rats, 3-4 cells/animal).
Average FI slope and input resistance did not differ between groups (FI slope: Control 0.07 ± 0.01 vs. CTA 0.09 ± 0.02, p = 0.17; Input Resistance: Control 136.47 ± 13.62MΩ vs. CTA 148.97 ± 24.59MΩ, p = 0.38). (D) Sample traces of sEPSCs (holding = -50mV) onto a control neuron and CTA neuron. Average EXC charge was larger in the control group (Control, [N = 8 rats, 3 cells/animal] 16.09 ± 0.90pC vs. CTA [N = 7 rats, 3-4 cells/animal] 13.32 ± 1.14pC, p < 0.01). (E) Cumulative probability plots showing a decrease in sEPSC amplitude (left), but not sEPSC instantaneous frequency (right) in the CTA group (sEPSC amplitude [n = 24 cells/group, α = 0.0004]: Control vs. CTA, p < 10^{-11}; sEPSC frequency [n = 24 cells/group, α = 0.0004]: Control vs. CTA, p = 0.02). Average sEPSC amplitude and frequency (insets) did not differ between groups (sEPSC amplitude: Control, [N = 8 rats, 3 cells/animal] 20.23 ± 3.12pA vs. CTA [N = 7 rats, 3-4 cells/animal] 17.94 ± 2.96pA, p = 0.32; sEPSC frequency: Control, [N = 8 rats, 3 cells/animal] 8.14 ± 5.17Hz vs. CTA [N = 7 rats, 3-4 cells/animal] 9.55 ± 5.90Hz, p = 0.75). * indicates p ≤ 0.05. Error bars ± 95% CI.

Figure 2-figure supplement 1: (A) FI curves from the control group before and after bath application of synaptic blockers (20μm DNQX, 50μm AP5, 20μm picrotoxin). Average FI slope decreased (Control [n = 18 cells from 3 rats] pre 0.055 ± 0.005 vs. post 0.051 ± 0.006, p < 0.006) and input resistance increased (Control [n = 18 cells from 3 rats] pre 128.24 ± 21.27MΩ vs. post 141.95 ± 21.99MΩ, p < 0.009) in the presence of synaptic blockers. (B) FI curves from the CTA group before and after bath application of synaptic blockers (20μm DNQX, 50μm AP5, 20μm picrotoxin). Average FI slope was unchanged (CTA [n = 19 cells from 3 rats] pre 0.062 ± 0.010 vs. post 0.059 ± 0.011, p = 0.16) but input resistance increased (CTA [n = 19 cells from 3 rats] pre 130.32 ± 18.66MΩ vs. post 139.26 ± 19.96MΩ, p < 0.05) in the presence of synaptic blockers. (C) Comparison of FI curves from the control and CTA groups after bath application of synaptic blockers (20μm DNQX, 50μm AP5, 20μm picrotoxin). FI slope (Control vs. CTA, p = 0.19) and input resistance (Control vs. CTA, p = 0.86) did not differ between groups in the presence of synaptic blockers. * indicates p ≤ 0.05. Error bars ± 95% CI.

Figure 3: CTA decreases BLA input onto L2/3 pyramidal neurons in aGC. (A) Example of histological verification of injection site and expression of opsin construct in BLA terminal fields in aGC. Left: Image of ChR2-Venus expression at BLA injection site (scale bar = 500μm; BLA,
Figure 3-figure supplement 1. (A) CTA does not affect the percentage of neurons that receive input from BLA (Control [N = 16 rats] 72.43 ± 14.19% vs. CTA [N = 13 rats] 69.74 ± 15.04%, p = 0.80), paired-pulse ratio (Control [N = 13 rats, 2-3 cells/animal] 1.45 ± 0.17 vs. CTA [N = 12 rats, 2-3 cells/animal] 1.36 ± 0.21, p = 0.55), coefficient of variation (Control [N = 13 rats, 2-3 cells/animal] 0.28 ± 0.06 vs. CTA [N = 12 rats, 2-3 cells/animal] 0.32 ± 0.09, p = 0.38), or decay tau (Control [N = 13 rats, 2-3 cells/animal] 0.008 ± 0.002 vs. CTA [N = 12 rats, 2-3 cells/animal] 0.007 ± 0.002, p = 0.66). Error bars ± 95% CI.

Figure 4. CTA learning occludes LTD induction onto L2/3 pyramidal neurons in aGC. (A) Schematic of 20 Hz plasticity induction paradigm. BLA terminal fields were activated with 20 bursts of 20 light pulses (5ms) at 20Hz delivered every 250ms, while the postsynaptic neuron was depolarized subthreshold by injection of a 50pA current step. (B) Distribution of the outcome of plasticity induction across all neurons recorded (Control LTD [n = 13 cells from 8 rats] 56.52%, no change [n = 5 cells from 5 rats] 21.74%, LTP [n = 5 cells from 2 rats] 21.74% vs. CTA LTD [n = 5 cells from 5 rats] 21.74%, no change [n = 7 cells from 4 rats] 30.43%, LTP [n = 11 cells from 6 rats] 47.83%, 3x2 χ², p < 0.05). (C) Sample traces of control group BLA-EPSC before and after 20Hz plasticity induction. 20Hz LTD induction did not affect control...
group PPR or CV (Control [13 cells from 8 rats]: PPR pre 1.66 ± 0.38 vs. post 1.36 ± 0.26, p = 0.10; CV pre 0.27 ± 0.09 vs. post 0.27 ± 0.09, p = 0.96). (D) Time course of change in BLA-EPSC amplitude (Control: % change from baseline -35.66 ± 2.65) and series resistance (Control: % change from baseline 2.57 ± 0.78) following 20Hz LTD induction in control group. Boxes indicate pre- and post-induction epochs. (E) Sample traces of CTA group BLA-EPSC before and after 20Hz plasticity induction. 20Hz LTP induction did not affect CTA group PPR or CV (CTA [11 cells from 6 rats]: PPR pre 1.39 ± 0.33 vs. post 1.22 ± 0.31, p = 0.09; CV pre 0.33 ± 0.08 vs. post 0.27 ± 0.10, p = 0.41). (F) Time course of change in BLA-EPSC amplitude (CTA: % change from baseline 62.37 ± 8.55) and series resistance (CTA: % change from baseline 0.94 ± 0.47) following 20Hz LTP induction in CTA group. Boxes indicate pre- and post-induction epochs. * indicates p ≤ 0.05. Error bars ± 95% CI.

**Figure 4-figure supplement 1.** (A) Sample traces from the control and CTA group before and after plasticity induction superimposed. (B) Summary of spiking activity during the 20Hz plasticity induction protocol. In both groups, the induction protocol elicited action potentials in a subset of neurons (Control: spikes [n = 7 cells from 4 rats] 30.43%, no spikes [n = 16 cells from 11 rats] 69.57% vs. CTA: spikes [n = 9 cells from 4 rats] 39.13%, no spikes [n = 14 cells from 7 rats] 60.87%, $\chi^2$, p = 0.75). (C) In both the control group and CTA group there was not a significant correlation between the magnitude of plasticity induced with the 20Hz protocol and the initial paired-pulse ratio (Control: $r_s = 0.00$, p = 1.00; CTA: $r_s = -0.07$, p = 0.75), coefficient of variation (Control: $r_s = 0.12$, p = 0.59; CTA: $r_s = -0.12$, p = 0.59), or decay tau (Control: $r_s = 0.07$, p = 0.74; CTA: $r_s = 0.11$, p = 0.61).

**Figure 5: Occlusion of LTD following CTA is specific to BLA activity regime.** (A) Schematic of tonic induction paradigm. BLA terminal fields were activated with 6s ramping stimuli delivered every 250ms, while the postsynaptic neuron was depolarized subthreshold by injection of a 50pA current step. (B) Distribution of the outcome of plasticity induction across all neurons recorded (Control LTD [n = 9 cells from 4 rats] 100.00%, vs. CTA LTD [n = 7 cells from 4 rats] 100.00%, 3x2 $\chi^2$, p = 1.00). (C) Sample traces of control group BLA-EPSC before and after ramp plasticity induction. Ramp LTD induction reduced control group PPR but did not affect CV (Control [9 cells from 4 rats]: PPR pre 1.22 ± 0.12 vs. post 0.92 ± 0.12, p < 10^{-3}; CV pre 0.18 ±
0.03 vs. post 0.20 ± 0.04, p = 0.33). (D) Time course of change in BLA-EPSC amplitude (Control: % change from baseline -61.87 ± 10.92) and series resistance (Control: % change from baseline 4.37 ± 4.01) following ramp LTD induction in control group. Boxes indicate pre- and post-induction epochs. (E) Sample traces of CTA group BLA-EPSC before and after ramp plasticity induction. Ramp LTD induction reduced CTA group PPR and increased CV (CTA [7 cells from 4 rats]: PPR pre 1.03 ± 0.16 vs. post 0.68 ± 0.21, p < 0.03; CV pre 0.20 ± 0.07 vs. post 0.24 ± 0.08, p < 0.03). (F) Time course of change in BLA-EPSC amplitude (CTA: % change from baseline -52.70 ± 6.48) and series resistance (CTA: % change from baseline 2.16 ± 4.79) following ramp LTD induction in CTA group. Boxes indicate pre- and post-induction epochs. * indicates p ≤ 0.05. Error bars ± 95% CI.

**Figure 5-figure supplement 1.** (A) Sample traces from the control and CTA group before and after plasticity induction superimposed. (B) Summary of spiking activity during the ramp plasticity induction protocol. The ramp induction protocol elicited action potentials in a subset of CTA neurons (CTA: spikes [n = 3 cells from 3 rats] 42.86%, no spikes [n = 4 cells from 2 rats] 57.14%) but no control neurons. (C) In the control group there was a significant correlation between the amount of plasticity induced with the ramp protocol and the initial coefficient of variation (Control: rs = -0.85, p < 0.01) but not with the paired-pulse ratio (Control: rs = -0.34, p = 0.37), or decay tau (Control: rs = 0.28, p = 0.47). There was not a significant correlation between the magnitude of plasticity induced with the ramp protocol and the initial paired-pulse ratio (CTA: rs = 0.16, p = 0.73), coefficient of variation (CTA: rs = 0.18, p = 0.70), or decay tau (CTA: rs = -0.22, p = 0.64) in the CTA group.

**Figure 6: BLA terminal field activation in vivo with the phasic 20 Hz pattern eliminated sucrose preference.** (A) Cartoon of experimental approach. AAV9.hSyn.eGFP.WPRE.bGH: (GFP control group) or AAV9.CAG.ChR2-Venus.WPRE.SV40 (Paired 20Hz group) were injected in BLA and an optic fiber (400µm diameter, coated with DiL) was implanted in aGC. (B) Correct positioning of injection site in BLA and of optic fiber in aGC were verified histologically. Left, green: injection site of ChR2-Venus; magenta: Hoechst counterstain; Right, green DiL indicating optic fiber tract; magenta: counterstain. White lines delineate anatomical landmarks indicating the location of BLA (left).
and GC (right) based on the rat brain atlas (Paxinos and Watson, 1998). Scale bar = 500µm. (C) To control for non-specific effects of light stimulation, one group of rats was injected with an AAV9 construct containing only the fluorescent tag Venus (GFP control, N = 8 rats). Diagram of behavioral paradigm for CTA in which the LiCl injection was substituted with 20Hz optogenetic stimulation of BLA terminal fields in GC following sucrose exposure. Fluid consumption across training days for GFP control group. Rats showed an initial preference for sucrose over water which persisted after conditioning (GFP control: C1 sucrose 11.60 ± 1.01mL vs. C2 sucrose 12.27 ± 0.86mL, p = 0.23) and at testing (GFP control: test day sucrose 12.03 ± 0.73mL vs. test day H2O 0.77 ± 0.12mL, p < 10⁻⁷). (D) Diagram of behavioral paradigm for CTA in which the LiCl injection was substituted with 20Hz optogenetic stimulation of BLA terminal fields in GC following sucrose exposure for rats that were injected with ChR2 (paired 20Hz opto, N = 8 rats). Fluid consumption across training days for paired opto group. Rats showed an initial preference for sucrose over water which shifted after conditioning (paired 20Hz opto: C1 sucrose 11.10 ± 2.55mL vs. C2 sucrose 12.36 ± 1.43mL, p = 0.41) at testing (paired 20Hz opto: test day sucrose 7.91 ± 2.10mL vs. test day H2O 5.35 ± 2.31mL, p = 0.24). (E) Diagram of behavioral paradigm for CTA in which rats received non-contingent 20Hz optogenetic stimulation of BLA terminal fields in GC the evening prior to sucrose exposure (non-paired 20Hz opto, [N = 5 rats]. Fluid consumption across training days for GFP control group. Rats showed an initial preference for sucrose over water which persisted after conditioning (non-paired 20Hz opto: C1 sucrose 13.11 ± 4.22mL vs. C2 sucrose 12.31 ± 1.62mL, p = 0.68) and at testing (non-paired 20Hz opto: test day sucrose 11.98 ± 3.40mL vs. test day H2O 2.86 ± 2.33mL, p = 0.07). (F) Diagram of behavioral paradigm for CTA in which the LiCl injection was substituted with optogenetic ramp stimulation of BLA terminal fields in GC following sucrose exposure for rats that were injected with ChR2 (ramp opto, [N = 6 rats]. Fluid consumption across training days for ramp opto group. Rats showed an initial preference for sucrose over water which persisted after conditioning (ramp opto: C1 sucrose 7.64 ± 1.66mL vs. C2 sucrose 9.60 ± 2.24mL, p = 0.06) and at testing (ramp opto: test day sucrose 9.81 ± 1.89mL vs. test day H2O 0.73 ± 0.43mL, p < 10⁻⁴). (G) Sucrose preference scores (C1 sucrose/average H2O) did not differ between groups (GFP Control 1.70 ± 0.28, paired 20Hz opto 1.52 ± 0.32, non-paired 20Hz opto 1.90 ± 0.74, ramp opto 1.30 ± 0.17, 1-way ANOVA p = 0.25). Aversion index of the paired 20Hz opto group differed from both the GFP control and ramp opto on test day (GFP Control 1.70 ± 0.28, paired 20Hz opto 1.52 ± 0.32,
non-paired 20Hz opto 1.90 ± 0.74, ramp opto 1.30 ± 0.17, 1-way ANOVA p < 10⁻³, GFP control vs. paired 20Hz opto, p = 0.001; paired 20Hz opto vs. ramp opto, p = 0.0017). * indicates p ≤ 0.05. Error bars ± 95% CI.

**Figure 6-figure supplement 1:** (A) Average water consumption prior to conditioning did not differ between groups (GFP control: average H₂O 6.87 ± 0.41mL/day vs. paired 20Hz opto: average H₂O 7.23 ± 0.42mL/day vs. non-paired 20Hz opto: average H₂O 7.14 ± 1.28mL/day vs. ramp opto: average H₂O 5.90 ± 1.28mL/day, 1-way ANOVA p = 0.21). (B) Rats in all four groups drank significantly more sucrose on the first conditioning day compared to baseline water intake (GFP control: average H₂O 6.87 ± 0.41mL/day vs. C1 sucrose 11.60 ± 1.01mL, p < 10⁻³; paired 20Hz opto: average H₂O 7.23 ± 0.42mL/day vs. C1 sucrose 11.10 ± 2.55mL, p = 0.0128; non-paired 20Hz opto: average H₂O 7.14 ± 1.28mL/day vs. C1 sucrose 13.11 ± 4.22mL, p < 10⁻⁵; ramp opto: average H₂O 5.90 ± 1.28mL/day vs. C1 sucrose 7.64 ± 1.66mL, p < 10⁻⁵). * indicates p ≤ 0.05. Error bars ± 95% CI.

**References**

Accolla R, Carleton A (2008) Internal body state influences topographical plasticity of sensory representations in the rat gustatory cortex. Proc Natl Acad Sci U S A 105:4010-4015.

Accolla R, Bathellier B, Petersen CC, Carleton A (2007) Differential spatial representation of taste modalities in the rat gustatory cortex. J Neurosci 27:1396-1404.

Bekkers JM, Delaney AJ (2001) Modulation of excitability by alpha-dendrotoxin-sensitive potassium channels in neocortical pyramidal neurons. J Neurosci 21:6553-6560.

Berman DE, Dudai Y (2001) Memory extinction, learning anew, and learning the new: dissociations in the molecular machinery of learning in cortex. Science 291:2417-2419.

Bermudez-Rattoni F, McGaugh JL (1991) Insular cortex and amygdala lesions differentially affect acquisition on inhibitory avoidance and conditioned taste aversion. Brain Res 549:165-170.

Bliss TV, Collingridge GL (1993) A synaptic model of memory: long-term potentiation in the hippocampus. Nature 361:31-39.

Braun JJ, Slick TB, Lorden JF (1972) Involvement of gustatory neocortex in the learning of taste aversions. Physiol Behav 9:637-641.

Cechetto DF, Saper CB (1987) Evidence for a viscerotopic sensory representation in the cortex and thalamus in the rat. J Comp Neurol 262:27-45.

Crozier RA, Wang Y, Liu CH, Bear MF (2007) Deprivation-induced synaptic depression by distinct mechanisms in different layers of mouse visual cortex. Proc Natl Acad Sci U S A 104:1383-1388.

Cubero I, Thiele TE, Bernstein IL (1999) Insular cortex lesions and taste aversion learning: effects of conditioning method and timing of lesion. Brain Res 839:323-330.
Dityatev AE, Bolshakov VY (2005) Amygdala, long-term potentiation, and fear conditioning. Neuroscientist 11:75-88.

Escobar ML, Bermudez-Rattoni F (2000) Long-term potentiation in the insular cortex enhances conditioned taste aversion retention. Brain Res 852:208-212.

Escobar ML, Alcocer I, Bermudez-Rattoni F (2002) In vivo effects of intracortical administration of NMDA and metabotropic glutamate receptors antagonists on neocortical long-term potentiation and conditioned taste aversion. Behav Brain Res 129:101-106.

Flynn FW, Grill HJ, Schwartz GJ, Norgren R (1991) Central gustatory lesions: I. Preference and taste reactivity tests. Behav Neurosci 105:933-943.

Fontanini A, Grossman SE, Figueroa JA, Katz DB (2009) Distinct subtypes of basolateral amygdala taste neurons reflect palatability and reward. J Neurosci 29:2486-2495.

Gal-Ben-Ari S, Rosenblum K (2011) Molecular mechanisms underlying memory consolidation of taste information in the cortex. Front Behav Neurosci 5:87.

Gandolfi D, Cerri S, Mapelli J, Polimeni M, Trito S, Fuzzati-Armentero MT, Bigiani A, Blandini F, Mapelli L, D'Angelo E (2017) Activation of the CREB/c-Fos Pathway during Long-Term Synaptic Plasticity in the Cerebellum Granular Layer. Front Cell Neurosci 11:184.

Garcia J, Kimeldorf DJ, Koelling RA (1955) Conditioned aversion to saccharin resulting from exposure to gamma radiation. Science 122:157-158.

Grossman SE, Fontanini A, Wieskopf JS, Katz DB (2008) Learning-related plasticity of temporal coding in simultaneously recorded amygdala-cortical ensembles. J Neurosci 28:2864-2873.

Guzman-Ramos K, Bermudez-Rattoni F (2012) Interplay of amygdala and insular cortex during and after associative taste aversion memory formation. Rev Neurosci 23:463-471.

Hadamitzky M, Bosche K, Engler A, Schiedowski M, Engler H (2015) Extinction of conditioned taste aversion is related to the aversion strength and associated with c-fos expression in the insular cortex. Neuroscience 303:33-41.

Haley MS, Fontanini A, Maffei A (2016) Lamina- and Target-Specific Amygdalar Inputs in Rat Primary Gustatory Cortex. J Neurosci 36:2623-2637.

Hansel C, Linden DJ (2000) Long-term depression of the cerebellar climbing fiber–Purkinje neuron synapse. Neuron 26:473-482.

Hirano T, Yamazaki Y, Nakamura Y (2016) LTD, RP, and Motor Learning. Cerebellum 15:51-53.

Ho J, Tumkaya T, Aryal S, Choi H, Claridge-Chang A (2019) Moving beyond P values: data analysis with estimation graphics. Nat Methods 16:565-566.

Inberg S, Elkobi A, Edri E, Rosenblum K (2013) Taste familiarity is inversely correlated with Arc/Arg3.1 hemispheric lateralization. J Neurosci 33:11734-11743.

Ito M, Yamaguchi K, Nagao S, Yamazaki T (2014) Long-term depression as a model of cerebellar plasticity. Prog Brain Res 210:1-30.

Jackman SL, Beneduce BM, Drew IR, Regehr WG (2014) Achieving high-frequency optical control of synaptic transmission. J Neurosci 34:704-7714.

Kakegawa W, Katoh A, Narumi S, Miura E, Motohashi J, Takahashi A, Kohda K, Fukazawa Y, Yuzaki M, Matsuda S (2018) Optogenetic Control of Synaptic AMPA Receptor Endocytosis Reveals Roles of LTD in Motor Learning. Neuron 99:985-998 e986.

Kayyal H, Yiannakas A, Kolatt Chandran S, Khamaisy M, Sharma V, Rosenblum K (2019) Activity of Insula to Basolateral Amygdala Projecting Neurons is Necessary and Sufficient for Taste Valence Representation. J Neurosci 39:9369-9382.

Kessels HW, Malinow R (2009) Synaptic AMPA receptor plasticity and behavior. Neuron 61:340-350.

Klaver O, Prigge M, Sarel A, Paz R, Yizhar O (2017) Manipulating fear associations via optogenetic modulation of amygdala inputs to prefrontal cortex. Nat Neurosci 20:836-844.
Kloc M, Maffei A (2014) Target-specific properties of thalamocortical synapses onto layer 4 of mouse primary visual cortex. J Neurosci 34:15455-15465.

Lavi K, Jacobson GA, Rosenblum K, Luthi A (2018) Encoding of Conditioned Taste Aversion in Cortico-Amygdala Circuits. Cell Rep 24:278-283.

Li W, Pozzo-Miller L (2019) Differences in GluN2B-Containing NMDA Receptors Result in Distinct Long-Term Plasticity at Ipsilateral versus Contralateral Cortico-Striatal Synapses. eNeuro 6.

Lin JY, Reilly S (2012) Amygdala-gustatory insular cortex connections and taste neophobia. Behav Brain Res 235:182-188.

Lin JY, Lin MZ, Steinbach P, Tsien RY (2009) Characterization of engineered channelrhodopsin variants with improved properties and kinetics. Biophys J 96:1803-1814.

Lin JY, Roman C, Arthurs J, Reilly S (2012) Taste neophobia and c-Fos expression in the rat brain. Brain Res 1448:82-88.

Ma T, Cheng Y, Roltsch Hellard E, Wang X, Lu J, Gao X, Huang CCY, Wei XY, Ji JY, Wang J (2018) Bidirectional and long-lasting control of alcohol-seeking behavior by corticostraitional LTP and LTD. Nat Neurosci 21:373-383.

Maffei A, Nelson SB, Turrigiano GG (2004) Selective reconfiguration of layer 4 visual cortical circuitry by visual deprivation. Nat Neurosci 7:1353-1359.

Maffei A, Nataraj K, Nelson SB, Turrigiano GG (2006) Potentiation of cortical inhibition by visual deprivation. Nature 443:81-84.

Malinow R, Malenka RC (2002) AMPA receptor trafficking and synaptic plasticity. Annu Rev Neurosci 25:103-126.

Morris R, Frey S, Kasambira T, Petrides M (1999) Ibogenic acid lesions of the basolateral, but not the central, amygdala interfere with conditioned taste aversion: evidence from a combined behavioral and anatomical tract-tracing investigation. Behav Neurosci 113:291-302.

Parsana AJ, Li N, Brown TH (2012) Positive and negative ultrasonic social signals elicit opposing firing patterns in rat amygdala. Behav Brain Res 226:77-86.

Petreanu L, Huber D, Sobczyk A, Svoboda K (2007) Channelrhodopsin-2-assisted circuit mapping of long-range callosal projections. Nat Neurosci 10:663-668.

Piette CE, Baez-Santiago MA, Reid EE, Katz DB, Moran A (2012) Inactivation of basolateral amygdala specifically eliminates palatability-related information in cortical sensory responses. J Neurosci 32:9981-9991.

Rioult-Pedotti MS, Friedman D, Donoghue JP (2000) Learning-induced LTP in neocortex. Science 290:53-56.

Rodriguez-Duran LF, Castillo DV, Moguel-Gonzalez M, Escobar ML (2011) Conditioned taste aversion modifies persistently the subsequent induction of neocortical long-term potentiation in vivo. Neurobiol Learn Mem 95:519-526.

Rogan MT, Staubli UV, LeDouc JE (1997) Fear conditioning induces associative long-term potentiation in the amygdala. Nature 390:604-607.

Rosenblum K, Meiri N, Dudai Y (1993) Taste memory: the role of protein synthesis in gustatory cortex. Behav Neural Biol 59:49-56.

Rumpel S, LeDouc J, Zador A, Malinow R (2005) Postsynaptic receptor trafficking underlying a form of associative learning. Science 308:83-88.

Schiff HC, Bouhuis AL, Yu K, Penzo MA, Li H, He M, Li B (2018) An Insula-Central Amygdala Circuit for Guiding Tastant-Reinforced Choice Behavior. J Neurosci 38:1418-1429.

Shema R, Sacktor TC, Dudai Y (2007) Rapid erasure of long-term memory associations in the cortex by an inhibitor of PKM zeta. Science 317:951-953.

Wang L, Kloc M, Gu Y, Ge S, Maffei A (2013) Layer-specific experience-dependent rewiring of thalamocortical circuits. J Neurosci 33:4181-4191.
Wang L, Kloc M, Maher E, Erisir A, Maffei A (2018a) Presynaptic GABA A Receptors Modulate Thalamocortical Inputs in Layer 4 of Rat V1. Cereb Cortex.

Wang L, Gillis-Smith S, Peng Y, Zhang J, Chen X, Salzman CD, Ryba NJP, Zuker CS (2018b) The coding of valence and identity in the mammalian taste system. Nature 558:127-131.

Yamamoto T, Fujimoto Y, Shimura T, Sakai N (1995) Conditioned taste aversion in rats with excitotoxic brain lesions. Neurosci Res 22:31-49.

Yasoshima Y, Yamamoto T (1998) Short-term and long-term excitability changes of the insular cortical neurons after the acquisition of taste aversion learning in behaving rats. Neuroscience 84:1-5.

Yilmaz-Rastoder E, Miyamae T, Braun AE, Thiels E (2011) LTP- and LTD-inducing stimulations cause opposite changes in arc/arg3.1 mRNA level in hippocampal area CA1 in vivo. Hippocampus 21:1290-1301.
### CTA paradigm

| DAY 1 - 4 | CONDITIONING x2 | TEST |
|-----------|------------------|------|
| 15min H2O (am) | sucrose | H2O | sucrose | 2-bottle | record or perfuse |
| ip LiCl | ip LiCl | ip LiCl | ip LiCl | |

### Pseudo training paradigm (control)

| DAY 1 - 4 | CONDITIONING x2 | TEST |
|-----------|------------------|------|
| 15min H2O (pm) | sucrose | H2O | sucrose | 2-bottle | record or perfuse |
| ip LiCl | ip LiCl | ip LiCl | ip LiCl | |

**Graphs:**

- **A:** Consumption (mL) over training days for H2O and sucrose.
- **B:** Aversion index over test day for H2O and sucrose.
- **C:** Paired D mean over training and test days.

**Images:**

- **D:** Immunohistochemistry for aGC L2/3.
- **E:** Control and CTA for CFOS and EGR1.
- **F:** CFOS+ cells/area and EGR1+ cells/area with mean values.
### A. Extinction testing

| HABITUATION | CONDITIONING x2 | TEST A | TEST B |
|-------------|------------------|--------|--------|
| 15min H2O (am) | sucrose | H2O | sucrose |
| DAY 1-4 | 5 | 6 | 7 |
| | ip LiCl | | ip LiCl |
| | 8 | | record or perfuse |
| | 9 | | record or perfuse |

### B

- Graph showing consumption (mL) over training days.

### C

- Graph showing consumption (mL) during test day comparison between H2O and sucrose.

### D

- Comparison of Aversion index between test days A and B.
A) control pre, control post, CTA pre, CTA post

B) control - 20Hz

C) Paired-pulse ratio

D) Coefficient of variation

E) Decay time
