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https://doi.org/10.1523/JNEUROSCI.0752-19.2019

Cite as: J. Neurosci 2019; 10.1523/JNEUROSCI.0752-19.2019

Received: 3 April 2019
Revised: 12 September 2019
Accepted: 17 September 2019

This Early Release article has been peer-reviewed and accepted, but has not been through the composition and copyediting processes. The final version may differ slightly in style or formatting and will contain links to any extended data.

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Activity of Insula to Basolateral Amygdala Projecting Neurons is Necessary and Sufficient for Taste Valence Representation

Abbreviated title: Cortical taste valence neurons

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Keywords: valence encoding, taste learning, memory, brain circuits, insula, amygdala

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Number of pages: 43

Number of figures (6), tables (0), multimedia (0), and 3D models (0)

Number of words for abstract (221), introduction (529), and discussion (1500)

Conflict of Interest Statement

The authors declare no conflict of interest.
Acknowledgments

This research was supported by a grant from the Canadian Institutes of Health Research (CIHR), the International Development Research Centre (IDRC), the Israel Science Foundation (ISF) and the Azrieli Foundation (ISF-IDRC 2395/2015); ISF 946/17; ISF-UGC 2311/15; and ChromISyn ERANET Neuron II supported by the Israel Ministry of Health Grant 3-12492 to K.R. H.K. is a recipient of the Edmond de Rothschild’s scholarship. Finally, yet importantly, we thank the members of the K.R. laboratory, specifically Dr. Shunit Gal Ben-Ari, for critical reading of this manuscript.
Abstract

Conditioned taste aversion (CTA) is an associative learning paradigm, wherein consumption of an appetitive tastant (e.g., saccharin) is paired to the administration of a malaise-inducing agent, such as intraperitoneal injection of LiCl. Aversive taste learning and retrieval require neuronal activity within the anterior insula (aIC) and the basolateral amygdala (BLA). Here, we labeled neurons of the aIC projecting to the BLA in adult male mice using a retro-AAV construct and assessed their necessity in aversive and appetitive taste learning. By restricting the expression of chemogenetic receptors in aIC-to-BLA neurons, we demonstrate that activity within the aIC-to-BLA projection is necessary for both aversive taste memory acquisition and retrieval, but not for its maintenance, nor its extinction. Moreover, inhibition of the projection did not affect incidental taste learning per se, but effectively suppressed aversive taste memory retrieval when applied either during or prior to the encoding of the unconditioned stimulus for CTA (i.e. malaise). Remarkably, activation of the projection following novel taste consumption, without experiencing any internal discomfort, was sufficient to form an artificial aversive taste memory, resulting in strong aversive behavior upon retrieval. Our results indicate that aIC-to-BLA projecting neurons are an essential component in the ability of the brain to associate taste sensory stimuli with body states of negative valence and guide the expression of valence-specific behavior upon taste memory retrieval.

Significance statement

In the present study we subjected mice to the conditioned taste aversion paradigm, where animals learn to associate novel taste with malaise (i.e., assign it negative valence). We show that activation of neurons in the anterior insular cortex (aIC) that project into the basolateral amygdala (BLA) in response to conditioned taste aversion is necessary to form a memory for
a taste of negative valence. Moreover, artificial activation of this pathway (without any feeling of pain) following the sampling of a taste can also lead to such associative memory. Thus, activation of aIC-to-BLA projecting neurons is necessary and sufficient to form and retrieve aversive taste memory.
The anterior insular cortex (aIC) plays a crucial role in taste learning (Rosenblum, 2008) and self-referential processes shaping conscious awareness in humans (Craig, 2009). Furthermore, it likely promotes circuit-wide dysfunction in neuropsychiatric disorders (Caria et al., 2010; Kurth et al., 2010; Critchley and Seth, 2012; Pais-Vieira et al., 2016). Nonetheless, the robust nature of taste learning paradigms, as well as their proven dependence on activity at the region, have made the aIC a particularly reliable target in studies of the molecular and cellular mechanisms underlying taste learning and memory (Bures et al., 1998; Belelovsky et al., 2005; Merhav et al., 2006; Yefet et al., 2006; Bermudez-Rattoni, 2014). Administration of a malaise-inducing agent (e.g. LiCl) following the consumption of appetitive tastants, results in conditioned taste aversion (CTA), through the association of the conditioned stimulus (CS) with the negative consequences of the unconditioned stimulus (US) (Garcia et al., 1955; Rosenblum et al., 1993).

CTA learning and retrieval is subserved by the gustatory cortex, located within the aIC (Lin, et al., 2015), composed of granular (GI), dysgranular (DI), and agranular (AI) subregions, arranged in the dorso-ventral plane (Kosar et al., 1986; Yiannakas and Rosenblum, 2017). Integration of sensory and reward stimuli is thought to be facilitated by reciprocal interactions with the limbic system (Krushel and van Der Kooy, 1988), the thalamus (Cechetto and Saper, 1987), as well as the basolateral and central amygdala (Grossman et al., 2008; Moraga-Amaro and Stehberg, 2012). The basolateral amygdala in particular is known to be required for stimulus salience encoding (Fontanini and Katz, 2009), as well as generating and relaying palatability signals to the IC (Piette et al., 2012). Correlative studies have suggested that connectivity between the aIC and the basolateral amygdala (BLA) facilitates the encoding and retrieval of sensory information in relation to body states, shaping their perceived valence (Gogolla, 2017; Haley and Maffei, 2018; Tye, 2018). In agreement...
with the above, we have shown recently that recruitment of IC neurons projecting to the BLA is increased during the retrieval of aversive taste memories, in a valence-, but not stimulus-dependent manner (Lavi et al., 2018). Despite providing evidence of high temporal precision in vivo, the methodology involved is correlative by its nature. We thus attempted here to address whether activation of the IC-to-BLA projection is necessary and/or sufficient for the expression of learned aversive taste experiences. Towards this aim, first, wild type (WT) were injected bilaterally with a retrograde-adeno-associated virus 2 (retroAAV) construct at the BLA, labeling monosynaptic anterograde connections of the IC with the BLA (Tervo et al., 2016). We analyzed the connectivity of aIC to BLA comparing dorsal-ventral, anterior-posterior and inner versus outer cortical layers (Fig. 1). To test the necessity of this projection in aversive taste memory learning, retrieval and maintenance (Fig. 3-6), we stereotactically injected mice with appropriate AAV constructs resulting in chemogenetic silencing or activation of IC-to-BLA projecting neurons (Fig. 2), using administration of the synthetic ligand Clozapine-N-oxide (CNO) locally or peripherally (Gomez et al., 2017). Our results indicate that recruitment of IC-to-BLA projecting neurons is an essential component in the ability of the brain to associate taste sensory stimuli with body states of negative valence and retrieve such aversive memories.
Methods

Animals

Wild type (WT) adult male mice (8-12 weeks) were used in all experiments described (Envigo, Israel). All mice used for the purposes of these studies were housed in the local Animal Resource Unit with water and standard chow pellet available ad libitum, under a 12-hour dark/light cycle, except where stated. All procedures were approved by the institutional animal care and use committee in accordance with the University of Haifa regulations and National Institutes of Health Guidelines, under Ethical License 428/16.

Surgery and viral injection

Adult (8-12weeks old) male WT mice were used as specified. Animals were administered an intraperitoneal (i.p.) injection of the analgesic norocarp (0.5mg/kg), 30 min prior to being anesthetized using a suitable M3000® anesthetic machine (NBT Israel®/Scivena Scientific®) using isoflurane (5%). Following the induction of anesthesia, animals were quickly adjusted to a Model 963 Kopf® stereotaxic injection system, where anesthesia was similarly maintained using isoflurane (2%). Following exposure of the skull and relevant alignment, mice were injected with appropriate AAV constructs at the anterior agranular insula (AP +0.86; ML ±3.4; DV -4.00) and/or the basolateral amygdala (AP -1.6; ML ±3.375; DV -4.80) as specified in each experiment (Tervo et al., 2016). In a separate cohort of mice, following bilateral stereotactic injection of AAV constructs at both regions, guide steel cannulas (23gauge) were inserted 1mm above the aIC, to allow local delivery of CNO upon recovery. Cannula placement was secured using dental cement as previously described (Alapin et al., 2018). Following viral delivery and appropriate cleaning and suturing (Vetbond®) of the exposed skull, animals were administered an additional i.p. dose of

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0.5mg/kg norocarp, as well as 0.5mg/kg of Baytril® (enrofloxacin). Animals were allowed an initial 2 hours to recover in an appropriately clean and heat-adjusted cage, and were then housed in larger cages with similarly treated cage-mates. Similar weight adjusted doses of the said analgesic and antibiotic agents were administered for an additional 3 days following recovery. All animals were allowed 3 weeks of recovery in their home cages. During the 4th week of recovery, animals were transferred to individual cages for 5 days, in preparation for water restriction training as described below.

All AAV constructs used in this study were obtained from the Viral Vector Facility of the University of Zurich (http://www.vvf.uzh.ch). All mice used in our studies were injected with 0.25μl/site/hemisphere of said AAV constructs (Physical Titer = 4.5 x 10E12 vg/ml) as defined in each experiment.

**Immunohistochemistry and quantification (Figure 1)**

Brain tissue was incubated for 24 hours in 4% formaldehyde solution, followed by 48 hours in 30% sucrose in phosphate buffered saline (PBS, MFCD00131855 - Sigma Aldrich®). Tissue was subsequently frozen at -80°C and processed for slicing using a cryostat (Leica CM 1950®). Twenty-four 40μm-thick brain slices were collected between Bregma 1.18 and 0.26, and treated for fluorescent immunohistochemistry. Immediately following slicing, tissue was briefly washed in PBS, prior to blocking and permeabilization for 1 hour using a 0.3% bovine serum albumin (10775835001 - Sigma Aldrich®)/0.3% Triton X-100 (MFCD00128254 - Sigma-Aldrich®)/10% fetal bovine serum solution (MFCD00132239 - Sigma-Aldrich®) in PBS (blocking solution). Slices were washed and mounted on glass slides using Vectashield® Mounting Medium with DAPI (H-1200) or without (H-1000), depending on whether EBFP-expressing AAVs were used (Vector Laboratories Cat# H-1200 and H-1000,
RRID: AB_2336790 and AB_2336789. Slides were visualized using a vertical light microscope (Olympus CellSens Dimension ©) at 10x and 20x magnification. Images were processed using Image-Pro © Plus V-7, Media Cybernetics © and manually quantified in terms of total numbers retroAAV* neurons in the respective subregions and layers of the aIC. Data were subsequently analyzed in terms of the bilateral number of positive cells/slice using 2-way ANOVA (Graphpad Prism ®).

Behavioral procedures (Figures 3-6):

In experiments involving chemogenetic inhibition of the IC-BLA projection, we bilaterally injected WT male mice into the IC with a Cre-dependent AAV construct expressing the inhibitory DREADD receptor in neurons, AAV8_hEF1a-dlox-hM4D (Gi) _mCherry (rev)-dlox-WPRE-hGHp (A) (Addgene plasmid # 50461; http://n2t.net/addgene:50461; RRID: Addgene_50461). In order to direct expression of the DREADD receptor into IC-BLA projecting neurons, retroAAV-hSyn1-chI-EBFP2_2A_iCre-WPRE-SV40p (A) or retroAAV-hSyn1-chI-EGFP2_2A_iCre-WPRE-SV40p (A) was injected at the BLA (Addgene plasmid # 81070; http://n2t.net/addgene:81070; RRID: Addgene_81070).

Similarly, in experiments involving chemogenetic activation of the IC-BLA projection, we bilaterally injected WT male mice with a Cre-dependent AAV construct expressing excitatory DREADD receptors in neurons of the IC, through AAV8_hEF1a-dlox-hM3D (Gq) _mCherry (rev)-dlox-WPRE-hGHp (A) (Addgene plasmid # 44361; http://n2t.net/addgene:44361; RRID: Addgene_44361). DREADD receptor expression was restricted to the IC-BLA projecting neurons, by injecting retroAAV-hSyn1-chI-EBFP2_2A_iCre-WPRE-SV40p (A) or retroAAV-hSyn1-chI-EGFP2_2A_iCre-WPRE-
SV40p (A) at the BLA (Addgene plasmid # 81070; http://n2t.net/addgene:81070; RRID: Addgene_81070).

For CTA acquisition, mice having recovered from surgery, received 1 ml of 0.5% saccharin- or 0.5% NaCl-water following four days of water restriction training (Adaikkkan and Rosenblum, 2015). Forty minutes following the start of the 20min drinking session (inter-stimulus interval, ISI=40min), animals were i.p. injected with a 2% body weight dose of the malaise inducing agent LiCl (0.14M), the US. For the subsequent 2 days, mice were maintained under water restriction. Three days following CTA acquisition, mice were provided with a choice between the conditioned tastant and tap water. Aversion to the conditioned tastant was calculated by expressing the volume of water consumed as a percentage of the total intake (water and tastant). Doses of CNO used in all experiments were chosen based on recent publications demonstrating that higher chronic doses of the ligand are metabolized into clozapine, which can affect behavior in itself (Gomez et al., 2017).

IC-BLA projection inhibition during CTA acquisition or retrieval - Systemic CNO Administration (Figures 3A-D)

During CTA Acquisition, mice received CNO (0.5mg/kg, i.p.) (Enzo® - http://www.enzolifesciences.com/BML-NS105/clozapine-n-oxide/) 30min prior to CTA training for saccharin (0.5%). Mice were provided a choice test between the conditioned tastant and water during retrieval testing 3 days later. To examine the effect of circuit inhibition during CTA retrieval, mice were trained in CTA for NaCl (0.5%). Three days’ later mice received 0.5mg/kg CNO 40min prior to retrieval testing. In both experiments, control mice received weight matched (1% body weight) injections of saline at the same intervals.
A separate group of mice were trained in CTA for saccharin as described above, and underwent chemogenetic inhibition of the IC-BLA projection during retrieval. Following the retrieval session, mice were used to examine the effect of circuit inhibition during retrieval on the subsequent extinction of the CTA memory using 9 unreinforced choice tests between the CS and water.

IC-BLA projection inhibition during CTA acquisition or retrieval - Local CNO Administration (Figures 5E-I)

Mice underwent surgery in order to place aIC cannulas allowing for the local inhibition of the aIC-to-BLA projection. Based on the current literature, mice received 0.25ul of 10uM CNO/site, at a speed of 0.1ul/min, dissolved in artificial cerebrospinal fluid (ACSF), or a similar volume of the vehicle (Burnett and Krashes, 2016). Delivery cannulas were secured as to allow mice to freely move during delivery without affecting the procedure. Mice treated during CTA acquisition received CNO or ACSF immediately prior to being presented with 1 ml of 0.5% NaCl for 20min. Immediately following the drinking session, mice received a 2% body weight dose of LiCl and were returned to their cages. Three days later, mice were tested using a 20min choice test between NaCl and water. Mice were then allowed an additional 3 days of recovery and the groups were reversed, so that the same mice did not receive CNO twice. Mice treated through cannulas during CTA retrieval, acquired CTA for 0.5% Saccharin and were similarly tested 72 hours following conditioning, receiving CNO or ACSF 15minutes prior (Adaikkan and Rosenblum, 2015).

IC-BLA projection inhibition during CTA memory maintenance (Figures 2E-F)

Following recovery and water restriction training, adult male mice were randomly assigned to CNO- and Saline-treated groups and underwent CTA training for saccharin (0.5%). Mice in the treatment group were administered CNO (0.5mg/kg, i.p.) 24hours following CTA training.
(2% body weight LiCl (0.14M, i.p.), ISI=40min) using 1ml of the saccharin solution, while the control group received an i.p. injection of saline (1% body weight). Animals were maintained on water restriction for an additional 48 hours, followed by a choice test between the CS and tap water.

**CNO control studies (Figures 5A-D)**

Mice were injected with retroAAV2_hSyn1-chi-EGFP_2A_iCre-WPRE-SV40P (A) at the BLA and AAV8_hEF1a-dlox-mCherry (rev)-dlox-WPRE-hGHp (A) at the aIC. Following 4 weeks of recovery mice were administered 1mg/kg CNO 30 minutes prior to CTA training, as described above. Mice were similarly examined using a choice test between the CS (0.5% NaCl) and water, 3 days later. Following a week of recovery, mice acquired CTA for a second tastant (0.5% Saccharin). Mice received CNO 40 minutes prior to a choice test between the CS and water, as described above. In order to confirm the validity of our approach using CNO in different phases of CTA acquisition (Figures 3-4), an additional cohort of 8 animals was treated with retroAAV2_hSyn1-chi-EGFP_2A_iCre-WPRE-SV40P (A) at the BLA and AAV8_hEF1a-dlox-mCherry (rev)-dlox-WPRE-hGHp (A) at the aIC. Following 4 weeks of recovery mice were administered CNO (0.5mg/kg, i.p.) 1 hour prior to the start of a 20min exposure to 1ml of 0.5% saccharin, while the control group received an i.p. injection of saline (1% body weight). Three hours later, both groups were administered a 2% body weight injection of LiCl. Mice were provided with a choice test between the conditioned tastant and tap water, 3 days later. Following an additional a week of recovery, the groups were reversed, so that the same mice did not receive CNO twice. Two hours following to the start of a 20min exposure to 1ml of the saccharin water, half of the cohort received a similar dose of CNO, while the control group received an i.p. injection of saline.
(1% body weight). Three hours from the start of drinking, both groups were administered a 2% body weight injection of LiCl.

Additional control studies were conducted in un-injected adult (8-12 weeks) WT male mice. These mice were water restricted for 5 days prior to CTA training for saccharin using 0.14M LiCl at ISI=40, as described in previous sections. Three days later, mice were given a choice test between pipettes containing the conditioned tastant and water, for 20min. Forty-five minutes prior to retrieval testing, half of the conditioned animals received CNO (1 mg/kg, i.p.), while the rest received a 1% body weight injection of saline.

IC-BLA projection inhibition prior to novel taste learning and attenuation of neophobia (Figures 4A-B)

Following surgery recovery, we separated mice into two groups (CNO and Saline). Mice in the treatment group were administered CNO (1 mg/kg, i.p.), while the control group received an i.p. injection of saline (1% body weight), 1 hour prior to a choice test between 0.5% saccharin and tap water. The above choice test was repeated for another 2 days, without any i.p. injections. Drinking volumes were recorded and relevant behavioral measures were calculated and analyzed.

IC-BLA projection inhibition prior to novel innately aversive taste exposure

We randomly assigned mice into two groups (CNO and Saline) following surgery recovery. Following 4 days of water restriction training, mice in the treatment group were administered CNO (1 mg/kg, i.p.) 1 hour prior to a choice test between 0.04% quinine and water. Conversely the control group received an i.p. injection of saline (1% body weight), prior to
the choice test. Drinking volumes were recorded and relevant behavioral measures were calculated and analyzed.

IC-BLA projection inhibition during CS encoding (Figures 3C-D)

Adult male mice randomly assigned to the treatment group were administered CNO (0.5mg/kg, i.p.) 1 hour prior to the start of a 20min exposure to 1ml of the saccharin water, while the control group received an i.p. injection of saline (1% body weight). Three hours later, both groups were administered a 2% body weight injection of LiCl (0.14M, i.p.). Mice were provided with a choice test between the conditioned tastant and tap water, 3 days later.

IC-BLA projection inhibition during US encoding (Figures 3E-F)

Mice in the treatment group were administered CNO (0.5mg/kg, i.p.) 2 hours following to the start of a 20min exposure to 1ml of the saccharin water, while the control group received an i.p. injection of saline (1% body weight). Three hours from the start of drinking, both groups were administered a 2% body weight injection of LiCl (0.14M, i.p.). Following an additional 48 hours on water restriction, mice were provided with a choice test between the conditioned tastant and tap water. Drinking volumes were recorded, and relevant behavioral measures were calculated and analyzed.

IC-BLA projection inhibition during the retrieval of trace fear conditioning (Figures 4G-H)

Mice were allowed 4 weeks of recovery and were subsequently randomly allocated into two groups (CNO and saline), without being isolated from their cage-mates. Animals were tested
in the subsequent 2 days were trained using trace fear conditioning (Curzon et al., 2014).

Twenty-four hours following training, mice were tested for contextual fear conditioning, being returned to the training chamber, where their activity was recorded for 180s without the tone. On the following day, mice in the treatment group were administered CNO (0.5mg/kg, i.p.) 45min prior to the start of the retrieval session, while the control group received an i.p. injection of saline (1% body weight). Behavior was recorded, and relevant measures were calculated and analyzed (Freeze Frame ® Coulbourn Instruments ®).

IC-BLA projection activation during weak CTA conditioning (Figures 6A-B)

Following recovery and isolation, mice were allocated into two groups, which following 4 days of water restriction training were provided 1ml of 0.5% NaCl solution. Mice in the treatment group were administered CNO (1 mg/kg, i.p.) 30 min prior to weak CTA conditioning (0.07M LiCl) while the control group received an i.p. injection of saline (1% body weight). Three days later, mice in both groups were provided with a choice test between the conditioned tastant and tap water.

IC-BLA projection activation following novel taste learning (Figures 6C-D)

Mice were allocated into two groups, which following 4 days of water restriction training were provided 1ml of 0.5% saccharin solution. Mice in the treatment group were administered CNO i.p. (1 mg/kg) 10 min following the end of drinking session, while the control group received an i.p. injection of saline (1% body weight). Mice in both groups were provided with a choice test between 0.5% saccharin-water and tap water, following an additional 48 hours of water restriction. Drinking volumes were recorded, and relevant behavioral measures were calculated and analyzed.
Mice in the treatment group were administered CNO (1 mg/kg, i.p.), 24 hours following weak CTA training (2% body weight LiCl (0.075M, i.p.), ISI=40min) using 1ml of the saccharin water, while the control group received an i.p. injection of saline (2% body weight). Animals were maintained on water restriction for an additional 48 hours, prior to a choice test. Drinking volumes were recorded, and relevant behavioral measures were calculated and analyzed.

Electrophysiology (Figure 2)

Insula slice preparations

Mice were injected with retroAAV2_hSyn1-chi-EGFP_2A_iCre-WPRE-SV40P(A) at the BLA and AAV8_hSyn1-dlox-hM3D(Gq)_mCherry(rev)-dlox-WPRE-hGHp(A) or AAV8_hEF1a-dlox-hM4D(Gi)_mCherry(rev)-dlox-WPRE-hGHp(A) at the aIC. To obtain brain slices containing insula, mice were deeply anesthetized with 5% isoflurane and transcardially perfused with 40 ml of ice cold oxygenated cutting solution containing (in mM): 25 NaHCO3, 105 Choline-Chloride, 2.5 KCl, 7 MgCl2, 0.5 CaCl2, 1.25 NaH2PO4, 25 D-Glucose, 1 Na-Ascorbate and 3 Na-Pyruvate. All reagents were commercially obtained from Sigma-Aldrich® Israel, except where stated. 300 μm thick coronal brain slices were cut with a Campden-1000® Vibrotome using the same cutting solution. The slices were incubated for at least 60 min at 34°C in artificial cerebrospinal fluid (ACSF) containing (in mM): 125 NaCl, 25 NaHCO3, 2.5 KCl, 1.25 NaH2PO4, 2 CaCl2, 1 MgCl2 and 25 D-Glucose, before transferring them to the electrophysiological setup. For electrophysiological recordings, slices were placed in an ACSF-perfused recording chamber (2 ml/min, 32-34°C). All solutions were constantly carbogenated with carbogen (95% O2 + 5% CO2).
Whole-cell recording

The slices were illuminated with infrared light, and pyramidal cells were visualized under differential interference contrast microscope (DIC) with ×40, water-immersion objective mounted on a fixed-stage microscope (BX51-WI; Olympus®). BLA-projecting neurons of the IC expressing the chemogenetic hM4DGi or hM3DGq receptors in a Cre-dependent manner were identified by the presence of mCherry. Cells projecting to the BLA that did not express the chemogenetic receptors were identified using GFP. Whole cell recordings from double fluorescence-labeled cells were performed using an Axopatch 200B amplifier and digitized by Digidata 1440 (Molecular Devices®). The recording electrode was pulled from a borosilicate glass pipette (3–5 MΩ) using an electrode puller (P-1000; Sutter Instruments®) and filled with a K-gluconate-based internal solution (in mM): 130 K-Gluconate, 5 KCl, 10 HEPES, 2.5 MgCl₂, 0.6 EGTA, 4 Mg-ATP, 0.4 Na₃GTP and 10 Phosphocreatine (Na salt). The osmolarity was 290 mOsm, and pH was 7.3. The recording glass pipettes were patched onto the soma region of pyramidal cells (Sharma et al., 2018). Voltages for liquid junction potential (+10 mV) were not corrected online. Current-clamp recordings were low-pass filtered at 10 kHz and sampled at 20 kHz. Series resistance was compensated and only series resistance <20 MΩ was included in the data set. Pipette capacitance was ~80% compensated. Following 3 to 5-minute stable baseline recording, CNO (2-10 μM) was added to the ASCF solution containing antagonist cocktail of (DNQX (20μM); APV (50μM); bicuculline methiodide (20μM), CAS40709-69-1 - Tocris®), was applied through the bath to the brain slice to isolate post-synaptic effects. This was followed by normal ACSF application for 5 to 30 min or until significant recovery in membrane activity was observed. The changes in resting membrane potential were measured 1min following CNO application (Nakajima et al., 2016).
**Results**

The majority of IC-to-BLA projecting neurons reside in the AI and DI subregions. To examine the role of IC-BLA projecting neurons in taste valence encoding, we first measured the differential neuroanatomical connectivity between the different subregions of the aIC and the BLA using injection of an retroAAV2 construct at the BLA, allowing the labeling of BLA-projecting neurons of the aIC (Fig. 1A, n=11). Quantification of mCherry+ neurons of the aIC was used to assess the efficiency and distribution of the retroAAV2 construct to label BLA-projecting neurons.

Approximately 5% of all the cells of the superficial or deep layers of the aIC are BLA projecting neurons. Approximately 62% of the BLA projecting neurons identified across the aIC reside within deep layers IV-VI rather than the superficial layers (Fig. 1B – F (1, 168) = 27.14, p<0.0001). This effect was prominent in the posterior part across the Bregma axis (Fig. 1B – F (3, 168) = 2.825, p=0.04). The majority of BLA-projecting neurons in the superficial (Fig. 1C – F (2, 252) = 173.2, p<0.0001) and deep layers IV-VI (Fig. 1D – F (6, 252) = 6.303, p<0.0001) of the aIC reside in the AI (48%) and DI (39%), while a smaller proportion reside at the GI (13%).

Unlike the DI and GI, BLA-projecting neurons in the superficial AI decrease in abundance from Bregma 0.86 onwards (Fig. 1C – F (3, 252) = 4.498, p=0.0043). In parallel, the number of BLA-projecting neurons is increased in the posterior part of the deep layers of the DI (Fig. 1D – F (3, 252) = 4.213, p=0.0063). In the deep layers of the AI, the number of BLA-projecting neurons rapidly declines from Bregma 0.86 onwards (Fig. 1D – F (6, 252) = 6.303, p<0.0001).
Activity in BLA-projecting neurons of the aIC can be chemogenetically manipulated using the retroAAV system. In order to test the functionality of this projection, we injected both retroAAV2_hSyn1_EGFP_iCre at the BLA, and Cre-dependent, AAV8_hEF1α-driven constructs at the IC to restrict the expression of chemogenetic receptors (hM4DGi or hM3DGq) in aIC-to-BLA neurons (see Methods). Whole-cell patch clamp recordings in slices from BLA-projecting neurons of the aIC confirmed that bath application of CNO (10 μM) rapidly hyperpolarized the resting membrane potentials (Wilcoxon test, Z=-21.00, p=0.0313) and inhibited the activity of neurons expressing hM4DGi-mCherry+ (Fig. 2, A, E, and F), but not in the hM4DGi-mCherry- or GFP+ expressing control cells (Fig. 2, C, I, and J, Wilcoxon test, Z=-7.00, p=0.3750). Similarly, in slices from mice treated to express hM3DGq in aIC-to-BLA neurons, bath application of CNO (2 μM) rapidly depolarized the resting membrane potentials (Wilcoxon test, Z=21.00, p=0.0313) and increased action potential firing in neurons expressing hM3DGq-mCherry+ (Fig. 2, B, G, and H), but not in the hM3DGq-mCherry- or GFP+ controls (Fig. 2, D, K, and L, Wilcoxon test, Z=0.00, p<0.9999).

IC-to-BLA projecting neurons are necessary for the acquisition and retrieval but not for the extinction or maintenance of learned aversive memories. Inhibition of aIC-to-BLA projecting neurons using CNO during CTA acquisition suppressed the aversive response (76.07±5.42%, n=8) compared to control animals (Fig. 3, B-C, 93.00±2.76%, n=6, Unpaired t-test, p=0.0276, t=2.507, DF=12). Significant suppression of the aversive response was also observed in comparing animals that acquired CTA normally (81.90±6.64, n=6), to ones that experienced inhibition (47.74±11.06%, n=7) of the projection during retrieval (Fig. 3, D and E, Unpaired t-test, p=0.0278, t=2.533, DF=11).

This latter treatment did not affect the subsequent extinction of this aversive memory (Fig. 3, H-I, p=0.9327, F (1, 24) =0.007284). Treated mice (88.04±3.996%, n=7) exhibited similar
aversion to control mice (93.04±5.329%, n=7) one day following the retrieval. Following 14 unreinforced choice tests between water and saccharin, control (57.30±11.69%, p=0.0166) and IC-BLA inhibited (63.70±9.389%, p=0.0344) mice exhibited similar extinction over time (Repeated Measures ANOVA - F (13, 156) =8.514, p<0.0001), but no significant effect of treatment was observed (F (1, 13) =0.3314, p=0.5747). Similarly, inhibition of the projection during intervals associated with memory maintenance following conditioning (Fig. 3, F-G) resulted in similar aversion (Unpaired t-test, p=0.8693, t=0.16 81, DF=12) to the CS upon retrieval between treated (92.01±3.924%, n=8) and control animals (91.12±3.124%, n=6).

Inhibition of IC-BLA-projecting neurons of the aIC disrupts specifically CS-US association during CTA acquisition, but does not affect innate and appetitive taste behaviors. To test the role of the projection in innate taste behaviors, we examined the effect of inhibition during appetitive novel taste learning and how this subsequently affects the attenuation of neophobia (Fig. 4A). Mice were treated with CNO before exposure to a novel tastant and were choice-tested for three consecutive days. Responses indicated a significant effect of taste exposure (Figure 4B – Repeated Measures ANOVA, F (2,38) =68.92, p<0.0001) but not of treatment (F (1,19) =0.072, p=0.7915). Aversion to the tastant observed in the first session (AN1 - Saline 69.122±5.257%, n=10; CNO 64.8±6.240%, n=11) was attenuated in the second (AN2 - Saline 38.231±5.677%, n=10; CNO 42.404±6.730%, n=11) and third session (AN3 - Saline 23.442±7.914%, n=10; CNO 30.113±6.160%, n=11). Aversion was significantly suppressed in AN2 compared to AN1, in both the CNO (p=0.0001) and Saline (p<0.0001) groups. Aversion was also significantly suppressed in AN3 compared to AN2, in both the CNO (p=0.0434) and Saline (p<0.0169) groups. We then tested whether inhibition of the projection affects aversion of innately aversive quinine by administering CNO or saline 1 hour prior to a choice test between quinine and water (not shown). Mice experiencing inhibition of the projection (100±0%, n=6) and mice receiving
saline (99.02±2.405%, n=6), exhibited no significant differences in aversion to quinine (Unpaired t-test, p=0.3409, t=1.000, DF=10).

To further examine whether the observed differences were due to taste recognition, we inhibited the projection prior to presenting the taste, and confined US association within the IC-dependent 3-hour margin (Adaikkan and Rosenblum, 2015). Projection inhibition during intervals associated with CS encoding affected the expression of aversion upon CTA retrieval when comparing saline (87.18±4.432%, n=9) and CNO (65.63±7.167%, n=8) treated animals (Fig. 4, C-D, Unpaired t-test, p=0.019, t=2.623, DF=15). Using the same rationale (Adaikkan and Rosenblum, 2015), we expanded the inter-stimulus interval and inhibited the projection prior to US administration (Fig. 4E). Similarly (Fig. 4F – Unpaired t-test, p=0.0016, t=3.686, DF=19), inhibition during US encoding for CTA suppressed aversion for the CS upon retrieval in CNO treated mice (46.01±8.611%, n=11), but not in saline (84.0±5.179%, n=10) treated mice.

We further show that inhibition of the projection prior to the retrieval of trace-fear conditioning does not affect freezing behavior (Fig. 4G-H), indicative of the specificity of this pathway for taste-malaise associations. Context testing resulted in similar freezing (F (1, 10) = 0.2873, p=0.6037) between the two groups. During the pre-tone, Saline (12.135±5.077%, n=5) and CNO (13.693±1.834%, n=5) treated mice showed similar freezing (p=0.4740). Tone (Saline – 35.722±6.842%; CNO – 37.910±8.565%; p=0.7879), and post-tone (Saline – 19.065±3.879%; CNO – 17.713±6.208%; p=0.6753) intervals was similar between the two groups (2-way ANOVA, F (1, 8) = 0.004139, p=0.9503).

In order to account for any non-specific effects of CNO administration (Gomez et al., 2017), we injected animals with AAV and retroAAV constructs at the IC and BLA, resulting in the expression of mCherry, without DREADD receptors. Behavioural experiments were
conducted (Fig 5, A-C) as previously described (see Methods). Control virus treated mice receiving CNO (n=5) and mice receiving saline (n=6) prior to CTA acquisition or retrieval, exhibit no significant effect differences due to treatment (Fig. 5B, 2-way ANOVA, F (1, 18) = 0.03366, p=0.8565), or as a consequence of the retrieval and acquisition protocols (Fig. 5B – 2-way ANOVA (1, 18) = 1.887, p=0.1864). Furthermore, towards assuring that the association in CS-and US experiments (Fig. 5, C and D) was to the saccharin and not to the injection of CNO, we repeated the experiments using control virus-injected mice. Our results indicate that CNO (94.00±1.78%, n=4) and saline (87.25±6.25%, n=4) treated mice show similar aversion to saccharin (Fig. 5, C-D, p=0.339, t=1.039, DF=6), meaning that the aversion was not altered by the consecutive intraperitoneal injections or CNO administration itself. Similar result was observed in the protocol described for CS association (Fig. 5C), where CNO injected (91.25±5.154%, n=4) and saline injected mice (82.25±5.154%, n=4) showed similar aversion to saccharin (p=0.4296, t=0.8468, DF=6), which further strengthens our suggestion of specific association to gastric malaise caused by the LiCl.

To rule out the possibility that additional collaterals from the BLA mediate the effect we observed, we infused CNO locally via cannulas during CTA acquisition or retrieval (Fig. 5E). Steel guide cannulas where placed at the aIC in addition to the relevant injection of AAV and retroAAV at the aIC and BLA (see Methods). CNO or saline was infused prior to the conditioning or retrieval of CTA for salty or sweet taste (Fig. 5, F-G). Following the intervention during conditioning, CNO (62.58±12.525%, n=4) treated mice showed a significant impairment in aversion in comparison to saline (91.40±1.55%, n=5) treated mice (Fig. 5G, Unpaired t-test, p=0.0361, t=2.587, DF=7). When examining the intervention during the retrieval of the memory (Fig. 5H-I, Unpaired t-test, p=0.0165, t=3.133, DF=7), aIC-to-BLA inhibited mice showed a reduction in aversion (61.84±7.663%, n=5) in comparison to their controls (90.20±2.888%, n=4). In summary, these results indicate that IC-
BLA projections underlie the encoding of malaise (i.e., US) or the association between tastes and unwanted effects of a perceived US. We thus tested the hypothesis that activation of the pathway is sufficient to serve as an artificial US.

**Activation of IC-to-BLA projecting neurons following the sampling of a novel taste is sufficient to form an artificial aversive taste memory.** In order to interrogate the validity of our hypothesis that activation of IC-to-BLA projecting neurons is sufficient for the expression of aversive taste behavior, mice were injected with AAV constructs, resulting in expression of the activating chemogenetic receptor (hM3DGq) tethered to mCherry in the projection (Fig. 6A, see Methods). Artificial activation of BLA-projecting neurons of the aIC during weak CTA conditioning (Fig. 6B-C, Unpaired t-test, p<0.0001, t=5.869, DF=18), significantly increased aversion (86.08±2.847%, n=10) to the conditioned tastant, compared to control (44.13±6.555%, n=10) animals. To examine whether activity within the IC-to-BLA pathway is sufficient for the formation of an aversive taste memory, a separate group of mice consumed a novel tastant (saccharin) followed by activation of the projection immediately afterwards (Fig. 6D, CNO 80.78±3.064%, n=12; Saline 24.24±8.015%, n=10), without receiving a malaise-inducing US (i.e., no visceral experience). This resulted in strong aversive behavior during the retrieval test (Fig. 6E, Unpaired t-test, p<0.0001, t=7.053, DF=20). Conversely, activation of the projection 24 hours following weak conditioning (Fig. 6F) resulted in similar aversion between control (86.3±7.100%, n=8) and CNO-treated mice (85.67±4.912, n=7) upon retrieval testing 48 hours later (Fig. 6G, p=0.9444, t=0.07109, DF=13), indicating that the influence of the projection does not extend to processes facilitating memory maintenance. In summary, our results demonstrate that activation of the IC-BLA pathway minutes following novel taste consumption is sufficient and necessary for the encoding and enhancement of inputs allowing for the association of CS and US during learning and is re-activated to guide behavior during retrieval.
Reciprocal connections of the anterior and posterior IC with the amygdala complex have been heavily implicated in reward-related encoding of both innate and learned (Stone et al., 2011; Jezzini et al., 2013; Haley et al., 2016). CTA acquisition results in a concerted bidirectional interplay of activity (Escobar and Bermúdez-Rattoni, 2000), post-translation regulation (Adaikkan and Rosenblum, 2012), protein synthesis (Levitan et al., 2016; Guzmán-Ramos et al., 2018), as well as transcription (Inberg et al., 2016) and histone deacetylation (Rodríguez-Blanco et al., 2019), at the two regions. Though a number of gustatory relay stations undergo changes in activity and plasticity markers (Lamprecht and Dudai, 1995; Rosenblum et al., 1997; Swank, 1999), the aIC is unique in that its inactivation produces deficits in acquisition (Buresová, 1978; Gallo and Bures, 1991), and prominently suppresses CTA retrieval (Yasoshima et al., 2000; Yasoshima and Yamamoto, 2005; Gal-Ben-Ari and Rosenblum, 2012).

Studies using CTA and other outcome devaluation models have since demonstrated that excitatory aIC neurons are necessary for the acquisition of learned aversive memories, but also the expression of non-homeostatic, valence-driven choice behaviors (Adaikkan and Rosenblum, 2015; Parkes et al., 2015; Baldo et al., 2016; Rogers-Carter et al., 2018). The DI and AI subregions integrate chemosensory, somatosensory, visceral and limbic information into a complex temporal code (Katz et al., 2001; Jones et al., 2006; Yokota et al., 2011), which provides the encoding framework for associative relationships between experience and outcome (Gardner and Fontanini, 2014; Haley and Maffei, 2018). The aIC is thus increasingly viewed as a center dedicated to the fine-tuning of behavior to salient sensory-reward/aversive associations, rather than a chemosensory center in the broader sense (Stehberg and Simon, 2011; Baldo et al., 2016; Fletcher et al., 2017; Fonseca et al., 2018; Inui et al., 2019b).
Presentation of the CS for CTA induces marked ERK activation in both the IC and BLA (Berman, 2003; Lin et al., 2010) and activation of BLA projections to reward centers (Inui et al., 2013). However, electrophysiological activity is inhibited in the majority of recorded BLA units in vivo (Uwano et al., 1995; Kim et al., 2010). Nonetheless, electrophysiological and molecular changes within excitatory BLA neurons have indeed been reported in a number of studies examining CTA acquisition and retrieval (Yasoshima and Yamamoto, 2005; Barot et al., 2008; Guzman-Ramos and Bermudez-Rattoni, 2012; Osorio-Gómez et al., 2017). Perhaps, despite their activation during acquisition, only a small proportion of BLA neurons is re-activated during retrieval, while the contribution of fibers projecting through or away from the BLA is indeed significant (Dunn and Everitt, 1988; Bahar et al., 2004; Inui et al., 2019a).

We have previously demonstrated using in vivo two-photon Ca\(^{2+}\) imaging of the aIC in mice, that CTA retrieval increases recruitment of the IC-BLA pathway, in a valence- and not stimulus-specific manner (Lavi et al., 2018). To further address the role of this projection in the expression of valence-specific behaviors, we first attempted to better define its spatial distribution (Fig 1.), by imaging the extended aIC in mice injected with retroAAV at the BLA (Tervo et al., 2016). We found that consistent with the literature (Krushel and van Der Kooy, 1988; Kobayashi, 2011; Gogolla, 2017), BLA-projecting neurons primarily reside within deep layers of the AI and DI (Fig. 1B-D). We then confirmed that infected cells at injection sites were limited to the BLA, and that projecting neurons can be chemogenetically manipulated through the Cre-dependent expression of DREADDs at the aIC (Fig. 2).

The bi-directional connectivity of the aIC with the BLA has been suggested to be key in shaping the perceived valence of sensory experiences through integrating the CS with subjective visceral and emotional experiences (Shi and Cassell, 1998; Maffei et al., 2012; Avery et al., 2017). We therefore hypothesized that recruitment of the projection subserves...
the encoding of negative internal body states (e.g., malaise induced by an US), but also facilitates their association with taste stimuli (e.g., saccharin), guiding the expression of appropriate behavioral responses upon retrieval (Höistad and Barbas, 2008; Craig, 2009). We tested our hypothesis by inhibiting or activating the projection during two different behavioral paradigms: the associative, negative learning paradigm of CTA and the incidental, positive learning paradigm of novel taste learning (Fig. 3-6). Indeed, inhibition of the projection during CTA acquisition and retrieval using intraperitoneal or local CNO injections (Fig. 3 & 5), resulted in significantly suppressed aversion to the CS upon choice testing, in support of previous findings (Lavi et al., 2018).

Furthermore, chemogenetic activation of the aIC-to-BLA pathway during weak conditioning enhanced aversive memory retrieval, and was surprisingly sufficient to drive the expression aversion to the CS in the absence of a real US sensory experience (Fig. 6B-E). In agreement with the above, inhibition during the CS- or US-encoding phase of CTA acquisition significantly suppressed learned aversive behavior, but did not affect innate taste behaviors (Fig. 4A-B). Importantly, inhibition during retrieval did not lead to faster extinction (Fig. 3H-I), while neither inhibition nor activation of the projection affected memory maintenance (Fig. 3F-G & 6F-G), suggesting that the manipulation itself does not permanently change or damage the neurons involved. We thus propose that during CTA acquisition, activation of the aIC-to-BLA projection facilitates CS-US associations, shapes the valence encoding to learned taste experiences, and is re-activated to enable the retrieval of past learned associations (Small et al., 2003; Fontanini et al., 2009; Piette et al., 2012). Inhibition of aIC-to-BLA neurons using local CNO delivery at the aIC further confirmed the necessity of the projection for CTA memory acquisition and expression (Fig. 5E-I), while suggesting that its recruitment acts upstream of BLA collaterals to reward centers (Dunn and Everitt, 1988; Juárez-Muñoz et al., 2017; Inui et al., 2019a). Nonetheless, it is possible that aIC collaterals
of the projection to other regions might also contribute to the effects elicited by chemogenetic
interventions (Wright and Groenewegen, 1996; Reynolds, 2005; Stachniak et al., 2014).

Projections of the posterior IC to the central amygdala (CeA) have been reported to shape and
reverse, upon manipulation, the valence of innate stimulus-specific responses to bitter tastants
(Schiff et al., 2018; Wang et al., 2018), while the anterior rostral IC-to-BLA projection has
been implicated in innate appetitive behaviors to sweet taste (Wang et al., 2018). Our
findings do not preclude these circuits being involved in innate taste responses, as we focused
on learned aversive behaviors and BLA-projecting neurons in portions of the region that were
not examined in the aforementioned studies (Schiff et al., 2018; Wang et al., 2018).
Moreover, the apparent discrepancy (Wang et al., 2018), might be further explained by
reports demonstrating a shift of the spatial arrangement of aIC responses across the posterior
axis following CTA acquisition in a valence-specific manner, resembling innate aversive
responses (Accolla and Carleton, 2008; Carleton et al., 2010; Lavi et al., 2018). However,
even though manipulation of the aIC-to-BLA projection does not affect innate taste
responses, other aIC circuits engaged in such innate responses, might contribute to CTA
learning and retrieval (Bales et al., 2015; Schiff et al., 2018; Inui et al., 2019b). On the other
hand, given the multimodal role of the insula, perhaps spatial overlapping between circuits
subserving innate and learned behaviors of opposing valence, might not necessarily constitute
a discrepancy (Ohla et al., 2019). For example, a recent study characterized a distinct Nos1-
expressing IC-to-CeA projection that is necessary for learned food overconsumption, but not
for homeostatic feeding (Stern et al., 2019). Activity and connectivity within complex
neocortical structures such as the insula, might not operate merely under spatial segregation
rules, and perhaps the role of certain circuits should be more carefully considered in relation
to their distinct molecular characteristics, among others (Grant, 2019).
To examine whether aIC-to-BLA neurons are also involved in the retrieval of other types of memory, we inhibited the projection during fear memory retrieval. However, chemogenetic inhibition (Gomez et al., 2017) failed to affect freezing behavior in conditioned mice (Fig. 4G-H). As other recent studies indicate, distinct amygdala-projecting neurons in the posterior IC (pIC) are more prominently involved (Grewe et al., 2017; Berret et al., 2019). Interestingly, even though this pIC projection participates in multiple stages during memory formation and retrieval (Berret et al., 2019), distinct CeA-projecting neurons drive acute freezing behavior, without leaving a memory trace (Grewe et al., 2017). Even so, we cannot rule out the possibility that the aIC-to-BLA projection participates in fear memory acquisition, or other types of learning and memory (Parkes and Balleine, 2013; Rogers-Carter et al., 2018). The aIC has been implicated in self-referential processes, and our results further suggest learned aversive behavior to arise, at least in part, through changes in activity and connectivity of the region in relation to internal, subjective states (Critchley and Seth, 2012; Pais-Vieira et al., 2016). The aIC-to-BLA projection is necessary for CS-US association during CTA acquisition, and is subsequently re-activated to guide retrieval, likely consequent to brain-wide adaptations necessary for memory maintenance (Smolen et al., 2019). Future studies should dissect the role of aIC connectivity and distinct cell types in taste memory encoding, maintenance and retrieval more comprehensively.
Author Contributions

HK and AY conducted surgeries, behavioral and immunohistochemical experiments, imaging and quantification, data sorting and analysis. SKC conducted all electrophysiological experiments. Early pilot studies were carried by VS. Mouse colony maintenance, welfare and genotyping by MK. AY, with HK and KR, planned the research wrote and edited the manuscript. KR supervised the research.

References


Baldo BA, Spencer RC, Sadeghian K, Mena JD (2016) GABA-Mediated Inactivation of


Burnett CJ, Krashes MJ (2016) Resolving Behavioral Output via Chemogenetic Designer


from the basolateral amygdala during the retrieval of conditioned taste aversion.


Inui T, Sugishita T, Inui-yamamoto C, Yasoshima Y (2019a) The basolateral nucleus of the amygdala executes the parallel processes of avoidance and palatability in the retrieval of conditioned taste aversion in male rats. Title: The basolateral nucleus of the amygdala executes the parallel processes of avoidance.


Yasoshima Y, Yamamoto T (2005) Effects of midazolam on the expression of conditioned


Figure 1. The majority of IC-to-BLA projecting neurons reside in the AI and DI subregions. BLA-projecting neurons of the aIC were labelled using a retroAAV2_hSyn_mCherry construct, and their distribution was quantified in the different spatial dimensions of the aIC across 24 coronal slices of the mouse brain (A). The average number of BLA-projecting neurons/slice in superficial layers I-III and deep layers IV-VI of the aIC, were plotted in relation to the Bregma axis and as percentage of number of DAPI+ cells (B). Analysis of the distribution using two-way ANOVA, indicated that a higher proportion of BLA-projecting neurons resided in deep layers (B) than in superficial layers, an effect that was observed at Bregma 0.62 and 0.38 across the axis (B). The percentage of BLA-projecting neurons in superficial layers (C), of the AI and the DI surpassed the numbers observed in the GI, across the Bregma axis (C). Higher number of BLA-projecting neurons of the AI was observed at 1.10 of the Bregma axis in comparison to the DI (C). Similarly, in deep layers (D), the percentages of BLA-projecting neurons in the AI and DI (D) were higher compared to the GI. In the DI, the number of BLA-projecting neurons tended to increase across the rostro-caudal axis (D), while it was decreased in the AI.

Figure 2. Activity in BLA-projecting neurons of the aIC can be chemogenetically manipulated using the retroAAV system. Using dual viral injections at the two regions (See Methods), we restricted the expression of chemogenetic receptors in IC-to-BLA projecting neurons in a Cre-dependent fashion (A-L). Whole cell current clamp recordings of mCherry⁺hM4DGi⁺ IC neurons projecting to BLA (F), showed rapid hyperpolarization of membrane potentials and reduced firing rates after the application of 10μM CNO for 60 seconds in the bath (A). Example traces from 5 similar recordings (n=6 cells from 5 mice)
from mCherry+hM4DGi+ neurons projecting from the IC to the BLA (A). Unlike responses to 10 μM CNO application in mCherry+ hM4DGi+ cells, non-hM4DGi expressing (mCherry-) EGFP+ neurons (J), did not show any change in membrane potential (C - n=4 from 4 mice).

Whole cell current clamp recording of mCherry+ hM3DG+ neurons in the insula projecting to BLA (H), showed rapid depolarization of membrane potentials and increased firing rates after the application of 2μM CNO in the bath for 60secs (B - n=6 cells from 5 mice). Unlike responses to 2 μM CNO application in mCherry+ hM3DGq+ cells, non-hM3DGq expressing (mCherry-) EGFP+ neurons (L), did not show any change in membrane potential (D - n=4 from 4 mice).

Measurement of resting membrane potential changes in DREADD and non-DREADD expressing cells, before and after the application CNO, showed a significant change in the RMP of DREADD expressing cells after the application of CNO, inhibitory and activator DREADD respectively (E, G - n=6). There was no change in the RMP of non- DREADD expressing control cells before and after the application of CNO (I, K), inhibitory and activator DREADD respectively (n=4). Scale bars in the traces (A-D) represent 20mV (y-axis) and 1minute (x-axis).

Figure 3. IC-to-BLA projecting neurons are necessary for the acquisition and retrieval but not for the extinction or maintenance of learned aversive memories. Adult (8-12 weeks) WT male mice were injected with viral constructs at the BLA (blue) and aIC (red), resulting in expression of inhibitory hMD4Gi in aIC-to-BLA neurons (A). Representative schematic overlays of the Cre-dependent expression of the chemogenetic receptors using the retroAAV systems is shown, demonstrating the expression to be restricted in the BLA and aIC (n=12 slices). Animals were split into Saline (light blue) and CNO (dark red) groups, and treated prior to CTA acquisition (B). Chemogenetic inhibition during CTA acquisition (C),
significantly suppressed aversion upon retrieval compared to control animals (C). In separate experiments, similarly injected mice, were used to assess the role of the projection in CTA retrieval (D). Inhibition during CTA retrieval, significantly suppressed aversion compared to control animals (E). To test whether the projection is also involved in CTA memory maintenance, we inhibited 24 hours following acquisition, and proceeded to test the animals 48 hours later (F-G). Inhibition 24 hrs following conditioning, resulted in similar aversion upon retrieval testing in control and treated animals. Even though inhibition of the projection during the retrieval of CTA suppresses aversion to the CS (H), 24 hours later, treated mice (I) exhibited similar aversion to control mice. Following 14 unreinforced choice tests between water and saccharin, control and IC-BLA inhibited mice exhibited similar extinction of the conditioned response (I) and no significant differences were observed among the two groups due to our intervention.

Figure 4. Inhibition of IC-BLA-projecting neurons of the aIC are disrupts CS-US association during CTA acquisition, but does not affect innate taste behaviors. Attenuation of neophobia was assessed for 3 days, in mice expressing hMD4Gi in aIC-to-BLA neurons, receiving CNO (dark red) or Saline (light blue) prior to novel saccharin consumption (A). Responses indicated a significant effect of taste exposure but not of treatment (B). Aversion to the tastant observed in the first session was attenuated in the second and third session. Aversion was significantly suppressed in AN3 compared to AN1, in both the CNO and Saline groups. Targeted inhibition of the CS for CTA (C), resulted in suppressed aversion upon retrieval in CNO treated to Saline groups (D). Correspondingly, inhibition at intervals associated with US encoding (E), resulted in significantly less aversion to the CS in CNO - treated mice compared to controls (F). A separate cohort of WT male mice treated to express hM4DGi in IC-to-BLA neurons, were trained in appropriate chambers.
to associate a tone to the administration of an electric shock using trace-fear conditioning (G).  
Mice were randomly split into two groups (CNO – dark red, Saline – light blue) and were  
tested in the subsequent 48 hours (G). The two groups exhibited similar levels of freezing  
during context-testing (H). On the following day, the CNO group received the artificial  
ligand 45min prior to being placed in a different context, where the tone was presented  
(without shock) at time points identical to the conditioning trial (G, H). Freezing recorded  
during the pre-tone, tone, and post-tone (H) intervals was similar between the two groups.

Figure 5. Chemogenetic inhibition of IC-BLA-projecting neurons of the aIC through  
local CNO administration at the aIC disrupts CTA memory acquisition and retrieval.  
Adult male mice were injected with viral constructs at the aIC and BLA, resulting in the  
expression of mCherry, without DREADD receptors (see Methods). Mice treated in this  
fashion received CNO either prior to the acquisition or prior to retrieval of CTA (B). Control  
virus treated mice receiving CNO and mice expressing chemogenetic receptors in the aIC-to-  
BLA projection but receiving saline prior to CTA acquisition or retrieval, exhibit no  
significant effect differences due to treatment or as a consequence of the retrieval and  
acquisition protocols. A separate set of mice that were treated in this fashion, and received  
CNO while expanding the inter-stimulus interval for CTA to examine CS- and US-specific  
effects (C). Aversion in CNO and Saline groups was similar (D). In a separate group of mice,  
DREADD receptors were introduced in the projection and guide cannulas were placed at the  
IC, allowing for local inhibition of the circuit, limiting the influence of collateralization (E).  
Diagrammatic representation of cannula placement at the aIC, as well as overlays of mean  
DREADD expression in cannulated mice, as indicated by mCherry+ neurons (E). Inhibition  
of the projection either during the acquisition (F, G), or during the retrieval of CTA (H, I),  
significantly suppressed aversion to the CS, regardless of stimulus identity.
Figure 6. Activation of IC-to-BLA projecting neurons following the sampling of a novel taste is sufficient to form an artificial aversive taste memory. WT male mice were injected with viral constructs at the BLA and aIC, resulting in Cre-dependent expression of activating hMD3Gq tethered to mCherry in BLA projecting neurons of the aIC (A). Representative schematic overlays of the Cre-dependent expression of the chemogenetic receptors using the retroAAV systems is shown, demonstrating the expression to be restricted in the BLA and aIC (n=12 slices). Following recovery, animals were split into groups receiving either Saline (light blue) or CNO (green) 30min prior to weak CTA training, followed by a retrieval choice test as described in (B). Chemogenetic activation of the projection using CNO (B) significantly enhanced the expressed aversion compared to control animals (C). We then tested whether chemogenetic activation of the IC-to-BLA projecting neurons following novel taste learning in mice, is sufficient to induce the expression of aversive taste behavior upon retrieval testing (D). Activation of the projection by administering CNO immediately following novel taste learning resulted in significant aversion in treated mice compared to control animals (E). To examine whether activation of aIC-to-BLA projecting neurons affects the maintenance of memory, in similarly injected animals, we activated the projection using CNO, 24hours following weak CTA training (F). Nonetheless, the mean aversion to the CS in the CNO and Saline (G) groups was similar following this intervention.
Fig. 1

A

Bregma 0.14mm
10x
Bregma 0.62mm
10x
Bregma 1.10mm
10x
B Total IC
10
5
0
1.10 0.86 0.62 0.38
retroAAV / DAPI [%]

B
Total IC

C
Layers I-III

D
Layers IV-VI

Bregma

Layer I-III
Layer IV-VI

AI
DI
GI

DI VS. AI
AI VS. GI
DI VS. GI

* DI VS. AI
v AI VS. GI
* DI VS. GI
FIG. 2

Inhibitory DREADDs
A

Cre+hM4Di  CNO 10 uM

-50mV

C

GFP+CNO 10uM

-50mV

D

Cre+hM3Di CNO 2uM

Excitatory DREADDs
B

GFP+CNO 2uM

-72mV

E

ACSF CNO[10uM]

0

-10

-20

-30

-40

-50

-60

-70

-80 RMP [mV]

F

G

H

J

I

L

M

N

O

P

Q

R

S

T

U

V

W

X

Y

Z
FIG. 3

A

B

C

D

E

F

G

H

I

Day 5
Conditioning
CNO/Saline 30min. Saccharin 40min. LiCl
Saline+Water

Day 8
Retrieval

Aversion Index [%]

Day 5
Conditioning
NaCl 40min. LiCl

Day 8
Retrieval
CNO/Saline 40min. NaCl+Water

Aversion Index [%]

Day 5
Conditioning
Saccharin 40min. LiCl

Day 6

Day 8
Retrieval
Saccharin+Water

Aversion Index [%]

Day 5
Conditioning
Saccharin 40min. LiCl

Day 8
Retrieval
CNO/Saline 40min. Saccharin+Water

Aversion Index [%]

Day 5
Conditioning
Saccharin 40min. LiCl

Day 8
Retrieval
CNO/Saline 40min. Saccharin+Water

Aversion Index [%]

Day 5
Conditioning
Saccharin 40min. LiCl

Day 8
Retrieval
CNO/Saline 40min. Saccharin+Water

Aversion Index [%]

0 10 20 30 40 50 60 70 80 90 100

Saline

CNO

Day

8 9 10 11 12 13 14 15 16 17 18 19 20

Aversion Index [%]
FIG. 5

A

Day 5
Conditioning
CNO/ Saline 30min. Saccharin 40min. LiCl

Day 8
Retrieval
Saccharin+Water

C

Day 5
Conditioning
CNO/ Saline 1 hr. Saccharin 3 hr. LiCl

Day 8
Retrieval
Saccharin+Water

D

Aversion Index [%]

0 20 40 60 80 100

Saline
CNO

US-specific
CS-specific

B

Aversion Index [%]

0 20 40 60 80 100

Saline
CNO

Acquisition
Retrieval

E

F

Day 5
Conditioning
Cannulated mice CNO/ Saline 20min. NaCl+Water

Day 8
Retrieval
NaCl+Water

G

Aversion Index [%]

0 20 40 60 80 100

Saline
CNO

* p<0.05

H

Day 5
Conditioning
Cannulated mice
Saccharin
LiCl

Day 8
Retrieval
Saccharin+Water

I

Aversion Index [%]

0 20 40 60 80 100

Saline
CNO

* p<0.05
FIG. 6

A

BLA 10x
IC 20x
20x 20x

B

Day 5 Day 8
Conditioning Retrieval
NaCl+Water
LiCl/CNO/
Saline NaCl 40min. 30min. (Weak CTA)

C

Aversion Index [%]
100
80
60
40
20
0

D

Day 5 Day 8
Conditioning Retrieval
CNO/Saline 30min. NaCl 40min. LiCl (Weak CTA) NaCl+Water

E

Aversion Index [%]
100
80
60
40
20
0

F

Day 5 Day 6 Day 8
Conditioning Retrieval
Saccharin 10min. CNO/Saline Saccharin+Water

G

Aversion Index [%]
100
80
60
40
20
0

Day 5 Day 6 Day 8
Conditioning Retrieval
Saccharin 40min. LiCl CNO/Saline Saccharin+Water