

TITLE: Dense cortical input to the rostromedial tegmental nucleus mediates aversive signaling

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ABSTRACT

The rostromedial tegmental nucleus (RMTg) encodes negative reward prediction error (RPE) and plays an important role in guiding behavioral responding to aversive stimuli. While initial studies describing the RMTg revealed the presence of cortical afferents, the density and distribution of this input has not been explored in detail. In addition, the functional consequences of cortical modulation of RMTg signaling are only just beginning to be investigated. The current study anatomically and functionally characterizes cortical input to the RMTg in rats. Findings from this work reveal dense input spanning the entire medial prefrontal cortex (PFC) as well as the orbitofrontal cortex and anterior insular cortex. Afferents were most dense in the dorsomedial subregion of the PFC (dmPFC), an area which has also been implicated in both RPE signaling and aversive responding. RMTg-projecting dmPFC neurons originate in layer V and collateralize extensively throughout the brain. In-situ mRNA hybridization further revealed that neurons in this circuit are predominantly D1 receptor-expressing with a high degree of D2 receptor colocalization. Optogenetic stimulation of dmPFC terminals in the RMTg drives avoidance, and cFos expression is enhanced in this neural circuit during exposure to aversive stimuli. Exposure to such aversive stimuli results in significant physiological and structural plasticity suggestive of a loss of top-down modulation of RMTg-mediated signaling. Altogether, these data reveal the presence of a prominent cortico-subcortical projection involved in adaptive behavioral responding and provide a foundation for future work aimed at exploring alterations in circuit function in diseases characterized by deficits in cognitive control over the balance between reward and aversion.

KEYWORDS: Prefrontal cortex, Insular cortex, aversion, conditioned fear

INTRODUCTION

Adaptive responding, in which past outcomes shape decision-making and future behaviors, is critical for survival. Impaired decision-making and maladaptive behaviors are common to a number of neuropsychiatric diseases and contribute significantly to harm and illness severity. Consequently, substantial effort has been put forth to identify the neural mechanisms governing such motivated behavior.

In the early 1980s, Corbett & Wise (1980) found that intracranial self-stimulation was most robust in rats with electrodes implanted in the ventral tegmental area (VTA) thereby uncovering a role for midbrain dopamine neurons in reward processing. Building upon this work, subsequent studies revealed that the activity of midbrain dopamine neurons encode a reward prediction error (RPE) with outcomes that are greater than expected producing a positive RPE associated with increased VTA dopamine neuron activity, and outcomes that are worse than expected producing a negative RPE associated with decreased VTA dopamine neuron activity (Schultz, 1986; Schultz et al., 1997). The neural circuitry driving RPE calculations within the VTA remains an area of intense investigation. To date, research suggests that positive RPE signals in the VTA are driven, at least in part, by excitatory input arising from the pedunculo pontine tegmental nucleus (PPTg) (Mena-Segovia and Bolam, 2017) with possible additional involvement from neurochemically heterogeneous afferents originating in the lateral hypothalamus (Nieh et al., 2015; Sharpe et al., 2017). Non-human primate studies suggested that the lateral habenula (LHb) played an important role in negative RPE signaling (Matsumoto and Hikosaka, 2007). However, this work presented somewhat of a paradox as previous studies had demonstrated that the LHb provides monosynaptic glutamatergic input to VTA dopamine neurons (Omelchenko et al., 2009) making it difficult to reconcile how this projection could facilitate a depression in phasic dopamine activity during presentation of aversive stimuli.

The solution to this paradox presented itself when two independent laboratories identified a previously unknown brain region called the rostromedial tegmental nucleus (RMTg), or tail of the VTA (Kaufling et al., 2009; Jhou et al., 2009b). Located immediately posterior to the VTA, both groups showed that the RMTg is primarily comprised of GABAergic neurons that receive dense input from the LHB and exert inhibitory control over monoaminergic and cholinergic midbrain nuclei including VTA dopamine neurons. Subsequent work found that RMTg activity increases in response to aversive stimuli of various sensory modalities (Jhou et al., 2009a; Li et al., 2019a) and that loss of RMTg function enhances active (e.g., escape) while reducing passive (e.g., freezing) responding in tests measuring fear and learned helplessness (Jhou et al., 2009a; Elmer et al., 2019). Circuit-specific approaches have further revealed that stimulation of dopamine-projecting RMTg neurons produces avoidance in a real-time place preference test and increases immobility in the forced swim test (St Laurent et al., 2020; Sun et al., 2020). In addition, using in vivo electrophysiology, Hong et al. (2011) demonstrated that inhibition of VTA dopamine neurons in response to aversive stimuli is driven via activation of a disynaptic circuit comprised of glutamatergic LHB neurons that provide input to VTA-projecting GABAergic neurons in the RMTg. Collectively, these studies suggest that the LHB-RMTg-VTA circuit plays a critical role in encoding negative RPE thereby guiding behavioral responding to aversive stimuli.

The prefrontal cortex (PFC) integrates incoming multisensory information with previous experience to provide top-down inhibitory control over behavior and guide goal-directed responding. Interestingly, subregions spanning the dorsomedial PFC (dmPFC), which includes the anterior cingulate cortex (ACC) and prelimbic (PL) PFC in rats, corresponding to Broadman's area 24 and 32 in humans, respectively, exhibit a number of functional similarities with the RMTg. For example, similar to the RMTg, neuronal activity in the PL PFC increases during exposure to aversive stimuli (Burgos-Robles et al., 2009) and loss of PL function reduces passive fear responding (Corcoran and Quirk, 2007). In addition, the activity in the dmPFC, and ACC in

particular, has been heavily implicated in RPE signaling (Alexander and Brown, 2019). As with the dmPFC, the rodent ventromedial PFC (vmPFC) is also comprised of two subregions. The more ventral rodent infralimbic (IL) PFC, which is thought to be homologous with Broadman's area 25 in humans, is well-characterized as frequently exerting opposing functions to the more dorsal PL mPFC (Peters et al., 2009; Gourley and Taylor, 2016) with activity in this region facilitating extinction of responses driven by PL PFC activity (Do-Monte et al., 2015). The dorsal peduncular cortex (DP) makes up the ventral-most portion of the vmPFC. While some have suggested the DP exhibits functional overlap with the IL PFC (Peters et al., 2009), very few studies have examined the DP directly and the human homolog of this region is, to date, unknown.

It is of interest that initial anatomical characterization of the RMTg revealed the presence of cortical afferents to the RMTg, including some that arose from the mPFC (Kaufling et al., 2009; Jhou et al., 2009b). However, the anatomy and function of these inputs have not been well-characterized as much of the research aimed at investigating the role of RMTg-associated neural circuits in aversive signaling has focused on the LHb-RMTg-VTA projection. The present study begins to fill this gap by anatomically and functionally characterizing cortical inputs to the RMTg with particular focus on those arising from the dmPFC given the functional overlap it shares with the RMTg. Our findings reveal the presence of dense input spanning a number of functionally distinct regions of the prefrontal and insular cortices and uncover a role for RMTg-projecting dmPFC neurons in top-down control over RMTg-mediated aversive signaling.

MATERIALS & METHODS

Animals

For all experiments, adult male Long-Evans rats (P60 upon arrival, Envigo Laboratories, Indianapolis, IN) were individually housed in standard polycarbonate cages. The vivarium was maintained on a 12:12 reverse light-dark cycle with lights off at 09:00. Rats were habituated to

the vivarium for at least one week before beginning experiments. All rats were provided with Teklad 2918 (Envigo) standard chow and water *ad libitum*. All experiments were approved by the Institutional Animal Care and Use Committees at the Medical University of South Carolina and University of Illinois at Chicago and adhered to the guidelines put forth by the NIH (National Research Council, 2011).

Stereotaxic surgery

Rats undergoing stereotaxic surgery were induced and maintained under a surgical plane of anesthesia using isoflurane (1-5%). Intracranial injections were made with back-filled custom glass pipettes connected to a nanojector (Drummond Scientific Company, Broomall, PA, USA) or a 200 μ L Hamilton syringe operated using a motorized pump (WPI, Inc, Sarasota, FL, USA). For tract tracing experiments, 100 nL 0.5% Cholera toxin B (CtB; Sigma Aldrich, St Louis, MO, USA) or 200 nL green fluorescent retrobeads (Lumafluor, Durham, NC, USA) were unilaterally injected into the RMTg (AP: -7.3; ML: +1.4; DV: -8.0 from skull; 6° lateral) at a rate of ~30 nL/s. For optogenetics experiments, rats were bilaterally injected with 500 nL of AAV2-hSyn-hChR2-(H134R)-eYFP-WPRE-pA (UNC Vector Core, Chapel Hill, NC, USA) into the dmPFC (AP: +3.2; ML: $\pm$ 0.6; DV: -3.5 from skull) or lateral habenula (LHb; AP: -3.6; ML: $\pm$ 1.2; DV: -4.1 from dura; 6° lateral) at a rate of 1-3 nL/s. During the same surgery, rats were implanted with custom-made 200 μ m optic fiber implants targeting either the RMTg (AP: -7.3; ML: $\pm$ 2.1; DV: -7.9 from skull; 10° lateral) or VTA (AP: -5.6; ML: $\pm$ 1.4; DV: -7.8 from skull; 6° lateral). Implants were secured with dental cement. An intersectional, dual-virus approach was used to investigate the extent of dmPFC-RMTg collateralization and structural plasticity in dmPFC-RMTg neurons following exposure to aversive stimuli. For these experiments, rats were unilaterally injected with 500 nL of either AAVretro-Cre (gift from Janelia Farms, Ashburn VA, US) or AAV2retro-pmSyn1-EBFP-Cre (Addgene, Watertown, MA, USA) into the RMTg (AP: -7.3; ML: +1.4; DV: -8.0 from skull; 6° lateral)

and 500 nL of AAV8.2-hEF1alpha-DIO-SYP-EYFP (Rachel Neve, MIT Vector Core) into the dmPFC (AP: +3.2; ML: +0.6; DV: -3.5 from skull) at a rate of 1-3 nL/s.

Cell density analysis

Rats unilaterally injected with CtB into the RMTg (n=9) were transcardially perfused with phosphate buffered saline (PBS) followed by 4% paraformaldehyde (PFA). Brains were immersion fixed overnight in 4% PFA, cryoprotected in 30% sucrose, and stored at -80 °C until ready for processing for microscopic analysis. Brains were sliced at 40 µm on a cryostat held at -20 °C. Slices containing the PFC and RMTg were labeled for CtB and NeuN using standard immunofluorescence procedures. In brief, slices were incubated in 50% (v/v) methanol for 30 min followed by incubation in 1% H₂O₂. Permeabilization was enhanced by incubation in 0.4% Triton-X in PBS followed by incubation in primary antibodies in PBS containing 0.2% Triton-X overnight at 4 °C (CtB 1°: 1:500, List Biological Laboratories #703; NeuN 1°: 1:500, EMD Millipore, MAB377). The tissue was then incubated with secondary antibodies for 2 h at room temperature (1:250, Jackson ImmunoResearch), rinsed in PBS, and mounted onto SuperFrost plus charged slides before being coverslipped with Fluoromount mounting medium (Sigma Aldrich). Images were acquired at 10X and tiled using an EVOS FL Auto microscope. Anatomical boundaries and rostrocaudal level of each PFC slice were determined by aligning the acquired microscopic images with atlas schematics generated using Paxinos & Watson (2007) in GIMP. ImageJ was used to apply a bandpass filter to Fourier-transformed images after which CtB+ and NeuN+ cells were automatically identified by searching for maximum intensity points.

Cell-type analysis

Adjacent tissue from that used in the cell density analysis experiments (n=3) was used to evaluate whether RMTg-projecting cortical neurons were glutamatergic or GABAergic projection neurons. Slices containing the PFC were labeled for CtB and the glutamatergic marker, CaMKIIα, or CtB

and the GABAergic marker, GAD67, using the same immunofluorescence procedures described above (CaMKII α 1°: 1:3,000, Invitrogen MA1048; GAD67 1°: 1:3,000, EMD Millipore MAB5406). Images of labeling in the dmPFC were acquired at 10X using a Zeiss AxioImager.M2 microscope. CtB-, CaMKII α -, and GAD67-labeled cell bodies were counted manually in each image using ImageJ. Cell counts were averaged across five slices spanning the rostrocaudal extent of the dmPFC for each rat and the ratio of cells labeled with each cell-type marker and CtB relative to all CtB-labeled neurons was calculated.

Collaterals analysis

A dual-virus, intersectional approach was used to label RMTg-projecting dmPFC neurons. After waiting at least eight weeks for optimal viral transduction and transgene expression, rats were euthanized and brains harvested using the same procedures described above. Brains were sliced on a cryostat at 40 μ m and eYFP signal was amplified using avidin-biotin immunohistochemistry procedures as previously published (Glover et al., 2016; GFP Abcam, ab290; 1:10,000). Brains were visually inspected from the rostral tip of the PFC to the rostral cerebellum for eYFP+ terminal labeling. Areas with noticeable labeling were imaged at 10X on a Zeiss AxioImager.M2 microscope. Images were flat field corrected and terminal density was analyzed by measuring the percent-stained area relative to total area using ImageJ. Analysis was performed on four slices spanning the rostrocaudal extent of each region and averaged together to arrive at a single data point for each region. Analysis of secondary somatosensory cortex and dorsal hippocampus were included as negative controls.

Real-time place preference testing

After at least eight weeks to allow for sufficient viral transduction and transgene expression, rats were tested for real-time place preference using procedures adapted from previously published work (Stamatakis and Stuber, 2012). Rats were habituated to the tethering procedure for at least

three days prior to testing. Testing was performed in an unbiased, custom-made apparatus consisting of two contextually distinct compartments. On day one, rats were connected to a patch cable connected to a 473 nm laser and allowed to freely explore the apparatus for 20 min. Light was delivered immediately upon entry into one compartment of the apparatus at 10 mW intensity and 60 Hz for the duration of time spent in that compartment. Light delivery was terminated upon entry into the opposite compartment. To confirm that behavioral responding was light-mediated, rats were re-tested 24 hours later using identical procedures except that the compartment associated with light delivery was reversed. Time spent in each compartment was quantified from video recordings made with a camera mounted above the testing apparatus using Ethovision (Noldus, Leesburg, VA, USA).

cFos induction following aversive stimuli

cFos induction was measured in RMTg-projecting mPFC neurons following exposure to aversive stimuli using procedures adapted from Jhou et al. (2009a). Rats were allowed at least one week to recover following stereotaxic injection of CtB into the RMTg before beginning testing. All rats underwent three days of habituation during which they were tethered and could freely explore a standard operant testing apparatus (Med Associates, St Albans, VT, USA). The house light was illuminated for the duration of each habituation session and all subsequent sessions. On day four, rats in the Context (control) group were euthanized 90 min after an identical 20 min habituation session. Rats in the Shock group were euthanized 90 min after presentation of a series of 10 foot shocks (0.5 mA, 0.5 s duration, 60 s inter-stimulus interval) over the course of a 20 min testing session beginning 60 s after the start of the session. Rats in the Shock-paired group underwent standard fear conditioning over two consecutive days where tone (2.9 kHz, 65 dB, 20 s duration) presentation co-terminated with foot shock (0.5 mA, 0.5 s duration). Two tone-shock pairings were presented 20 min apart over the course of each 60 min conditioning session. Rats in the Shock-unpaired group were presented with the same stimuli as the Shock-paired group during two 60

min sessions except that stimuli were explicitly unpaired occurring 10 min apart. Following conditioning trials, rats from both the Shock-paired and Shock-unpaired groups were re-habituated to the testing apparatus during a 30 min session where the house light was illuminated but no stimuli were presented. On the test day, rats from both groups were euthanized 90 min after a 20 min test session consisting of presentation of eight tones for 30 s each (60 s inter-stimulus interval). Freezing during tone presentation was scored manually in the Shock-paired and Shock-unpaired rats using overhead video recorded during the test session. Following euthanasia, brains were processed for CtB (1:300,000) and cFos (Millipore #PC38, 1:10,000) expression using previously published procedures (Glover et al., 2016). The number of double-labeled neurons relative to all CtB+ neurons was quantified manually across 4-5 slices spanning the rostrocaudal extent of the mPFC. Rats with off-target injection sites were excluded from analysis.

In-situ hybridization

Rats were unilaterally injected with green retrobeads into the RMTg and allowed at least seven days to recover before testing. The animals were assigned to either Context or Shock groups and underwent testing identical to that described above for the cFos induction experiments. Rats were anesthetized with isoflurane and decapitated 90 min after the test session. The brains were then rapidly removed and placed in ice-cold PBS for ~5 minutes before being embedded in Tissue-Tek OCT media (Sukura Finetek Inc, Torrance, CA, USA) in Peel-A-Way cryo-embedding molds (Polysciences, Inc, Warrington, PA, USA) and covered with dry ice. The frozen tissue block was then extracted from the mold, wrapped in aluminum foil, and stored at -80 °C. Subsequently, 20 µm thick slices from the fresh-frozen brains were cut on a cryostat, mounted on SuperFrost Plus slides (Fisher Scientific, Hampton, NH), and stored at -80 °C until their use in in-situ hybridization experiments.

231 Fluorescence in-situ hybridization was performed using an Advanced Cell Diagnostics (ACD,
 232 Newark, CA) Multiplex RNAScope kit (catalog # 323100). RNA probes for dopamine D1 receptors
 233 (catalog # 317031), dopamine D2 receptors (catalog # 315641-C2), and cFos (catalog # 403591-
 234 C4) were also obtained from ACD. The RNAScope procedure was carried out according to the
 235 manufacturer's instructions (available for download at www.acdbio.com) with the exception that
 236 the protease digestion step was omitted. We observed that omission of this step not only improved
 237 the fluorescence intensity of the mRNA transcripts (visually observed as punctate dots), but was
 238 also required for preservation of the fluorescent intensity of the green (alexa-488) retrobeads. For
 239 multiplex hybridization of D1 and D2 mRNA transcripts, the probes were labeled with Cy3 and
 240 Cy5, respectively. For multiplex hybridization of cfos and D1 mRNA transcripts, the probes were
 241 labeled with Cy3 and Cy5, respectively. Images were acquired on a Zeiss LSM880 confocal
 242 microscope across three PFC slices (5 images/slice) using a 63X oil objective. Imaging was
 243 restricted to areas of the dmPFC that exhibited retrograde bead labeling, which was mainly
 244 observed in cortical layer V. Quantification and colocalization of mRNA transcript dot within cells
 245 was performed on the captured images using Imaris Software (Bitplane, Zurich, Switzerland)
 246 following a previously published method (Centanni et al., 2019). This analysis utilized the Cell
 247 Module of Imaris, and can be summarized as follows: 1) Define and interactively threshold the
 248 cell nucleus based on DAPI staining (we used a minimum size of 5 μm and a filter of 0.5); 2)
 249 Define and interactively threshold the cell body based upon the DAPI identified nucleus in step 1;
 250 3) Define and interactively threshold the mRNA transcript dots for each probe and for the
 251 retrobeads (we used a minimum size of 1 μm for both the transcript dots and retrobeads); 4)
 252 Calculation of the number and other parameters of the dots that lie within each defined cell. For
 253 a cell to be considered as positive for fluorescent beads or D1/D2 mRNA transcripts, it had to
 254 contain two or more dots/beads. We observed that a number of cells exhibited a variable level of
 255 background cfos mRNA irrespective of whether they were in the Control or Shock group.
 256 Therefore, for the purpose of assessment of the effect of shock on cfos mRNA expression, we

used a threshold of 15 or more cfos transcript dots in order to consider a cell as being cfos+. This threshold was determined based on a comparison of the distribution of the cfos mRNA transcript dots in the control versus shocked conditions.

Whole-cell patch-clamp slice electrophysiology

Rats were unilaterally injected with green retrobeads into the RMTg and allowed at least three days to recover before being assigned to either Context or Shock groups and undergoing testing identical to that described above for cFos induction experiments. Twenty-four hours following the final day of testing, the intrinsic excitability of dmPFC pyramidal neurons was determined using previously published procedures (Wayman and Woodward, 2018). In brief, rats were anesthetized with urethane (3.0mg/kg, i.p.) and perfused with an ice-cold sectioning solution consisting of (in mM): 200 sucrose, 1.9 KCl, 6 MgSO₄, 1.4 NaH₂PO₄, 25 NaHCO₃, 0.5 CaCl₂, 10 glucose, and 0.4 ascorbic acid; pH 7.35-7.45 with 310-320 mOsm. The brains were then immediately harvested and coronal brain sections (300 µm) containing the dmPFC were sliced on a Leica VT1000S vibratome (Leica Biosystems, Buffalo Grove, IL) in oxygenated (95% O₂; 5% CO₂) sectioning solution and then transferred to a holding chamber containing normal artificial cerebrospinal fluid (aCSF; in mM): 125 NaCl, 2.5 KCl, 25 NaHCO₃, 1.4 NaH₂PO₄, 1.3 MgCl₂, 2 CaCl₂, and 10 glucose; pH 7.35-7.45 with 310-320 mOsm. Brain slices were incubated at 34 °C for 30 minutes and allowed to recover at room temperature for an additional 45 minutes.

For current clamp recordings, brain slices were transferred to the recording chamber and perfused with oxygenated and heated (~34 °C) aCSF at a flow rate of 2 mL/min. The temperature was maintained during the course of the recordings with in-line and bath heaters (Warner Instruments, Hamden, CT). Retrobead-labeled layer V neurons within the dmPFC were visually identified using a Zeiss FS2 microscope (Zeiss, Thorndale, NY). Recording pipettes were constructed from thin-walled borosilicate capillary glass tubing (I.D.=1.17mm, O.D. 1.50mm; Warner Instruments,

Hamden, CT), pulled with a horizontal pipette puller (P-97 Sutter Instrument Co., Novata, CA). Pipettes were filled with an internal solution containing (in mM): 120 K-gluconate, 10 HEPES, 10 KCl, 2 MgCl₂, 2 Na₂ATP, 0.3 NaGTP, 1 EGTA and 0.2% biocytin; pH 7.35-7.45 with 285-295 mOsm and had resistances ranging from 3-5 MΩ. After a stable gigaohm seal was formed, light suction was applied to break through the cell membrane and achieve whole-cell access. Neurons with an access resistance of greater than 20 mΩ were not used for analysis. Recorded events were acquired with an Axon MultiClamp 700A amplifier (Molecular Devices, Union City, CA), digitized at a sampling rate of 10 kHz (filtered at 4 kHz) with an Instrutech ITC-18 analog-digital converter (HEKA Instruments, Bellmore, NY) controlled by AxographX software (Axograph Scientific, Sydney, Australia) running on a Macintosh G4 computer (Apple, Cupertino, CA). The resting membrane potential (RMP) and capacitance of all neurons was first recorded and then the RMP was adjusted to -70 mV for electrophysiological assessments of excitability. Action potential firing was induced by a series of 500 ms current steps (0-300 pA) incremented in +20 pA steps. Recordings were analyzed offline for the number of spikes in response to each current step, threshold (mV), rheobase (pA), action potential peak amplitude (mV), action potential half-width (ms) and after-hyperpolarization (AHP; mV) using AxographX software.

The caudal portion of the brain containing the RMTg was collected at the same time that dmPFC slices were generated, immersion fixed overnight in 4% PFA, and frozen on dry ice followed by storage at -80 °C until processing. Injection sites were confirmed by visual inspection of fluorescent retrobead labeling in slices containing the RMTg made using a cryostat.

Spine density analysis

A dual-virus, intersectional approach was used to label RMTg-projecting dmPFC neurons. After waiting at least eight weeks for optimal viral transduction and transgene expression, rats were assigned to either Context or Shock groups and underwent behavioral testing identical to that

described above for cFos induction experiments. Twenty-four hours later, rats were euthanized, and brains harvested as described for cell density analysis. Brains were sliced at 100 μm and eYFP signal was amplified (GFP, Abcam #ab290; 1:30,000) using immunofluorescence procedures optimized for thick slices (Kupferschmidt et al., 2015). Primary apical dendrites measuring 55 μm in length approximately 200-300 μm from the soma of eYFP+ neurons in the dmPFC were imaged using a 63.5X oil immersion objective on a Zeiss LSM880 confocal microscope. Images were analyzed in Imaris using previously published procedures (McGuier et al., 2015). Dendrite diameter, dendrite volume, and total spine density were analyzed in addition to analyses conducted by spine classification. Measures included density, length, diameter, and volume by spine class as well as diameter and volume of spine terminal point and spine neck volume, length, and diameter. Measures were collected in 3-5 dendrites per rat and averaged across dendrites to arrive at a single value for each rat.

Statistical analysis

Student's t-test and analysis of variance (ANOVA) were performed to analyze all functional data as indicated below in the Results section. Greenhouse-Geisser correction was performed on data that lacked sphericity. All analyses were performed using GraphPad Prism 8.0 and are presented as mean $\pm$ SEM. Effects were considered statistically significant at $p \leq 0.05$.

RESULTS

Cortical input to the RMTg is dense, glutamatergic, and exhibits extensive collaterals

While initial reports indicated the presence of cortical efferents to the RMTg, the magnitude of this input and subregional distribution was unclear. To investigate this, RMTg-projecting cell bodies were quantified in rat brains injected with the retrograde tracer CtB. Visual inspection of slices double stained for CtB and the neuronal marker, NeuN, revealed the presence of dense input spanning the medial wall of the PFC and the entire OFC (**Figure 1A**). In line with previous reports,

relatively low but consistent labeling was also observed in the anterior insular cortex (AIC) (Kaufling et al., 2009; Zhou et al., 2009b). In agreement with the well-understood layer specificity of cortico-subcortical projections in the rodent mPFC, the majority of CtB⁺ cell bodies originated in layer V (**Figure 1B**). Consistent cell body labeling was also apparent in the deepest portion of layer VI, albeit to a much smaller degree than was observed in layer V. Quantification of layer V CtB⁺ neurons relative to NeuN⁺ neurons in the mPFC revealed relatively uniform densities of RMTg-projecting neurons across the rostrocaudal extent of ACC ($3.96 \pm 0.28\%$), PL ($8.59 \pm 0.45\%$), and IL ($7.95 \pm 0.77\%$) subregions (**Figure 1C**). In contrast, the density of RMTg-projecting dorsopeduncular (DP) mPFC neurons increased substantially at more caudal levels relative to rostral DP mPFC ($13.02 \pm 2.91\%$).

The density of CtB-labeled neurons was relatively similar across subregions of the OFC at rostral levels but began to diverge slightly more caudally (**Figure 1E**). On average, density was greatest and somewhat variable in the medial orbital (MO) cortex ($8.01 \pm 1.32\%$). By contrast, CtB labeling was lower and less variable in the dorsolateral orbital (DLO) cortex ($5.48 \pm 0.62\%$). The density of RMTg-projecting ventro-orbital (VO) and latero-orbital (LO) cortical projections varied substantially from rostral to more caudal levels within the brain. In the VO, density was greatest at the rostral-most point of the OFC ($9.18 \pm 1.43\%$) after which CtB labeling diminished ($4.35 \pm 0.98\%$) before increasing in density at its most caudal point ($7.03 \pm 1.87\%$). By contrast, RMTg-projecting neurons arising from the LO cortex are most dense at the rostral tip of the region ($7.45 \pm 1.43\%$) and become progressively less dense as one moves caudally with very little CtB⁺ cell bodies in the most caudal region ($1.05 \pm 0.73\%$).

Although substantially less dense than projections arising from the mPFC and OFC, CtB⁺ cell bodies were consistently observed in subregions of the AIC (**Figure 1F**). Density was greatest in the agranular AIC with approximately 4.5% of layer V neurons in dorsal (AID) and ventral (AIV)

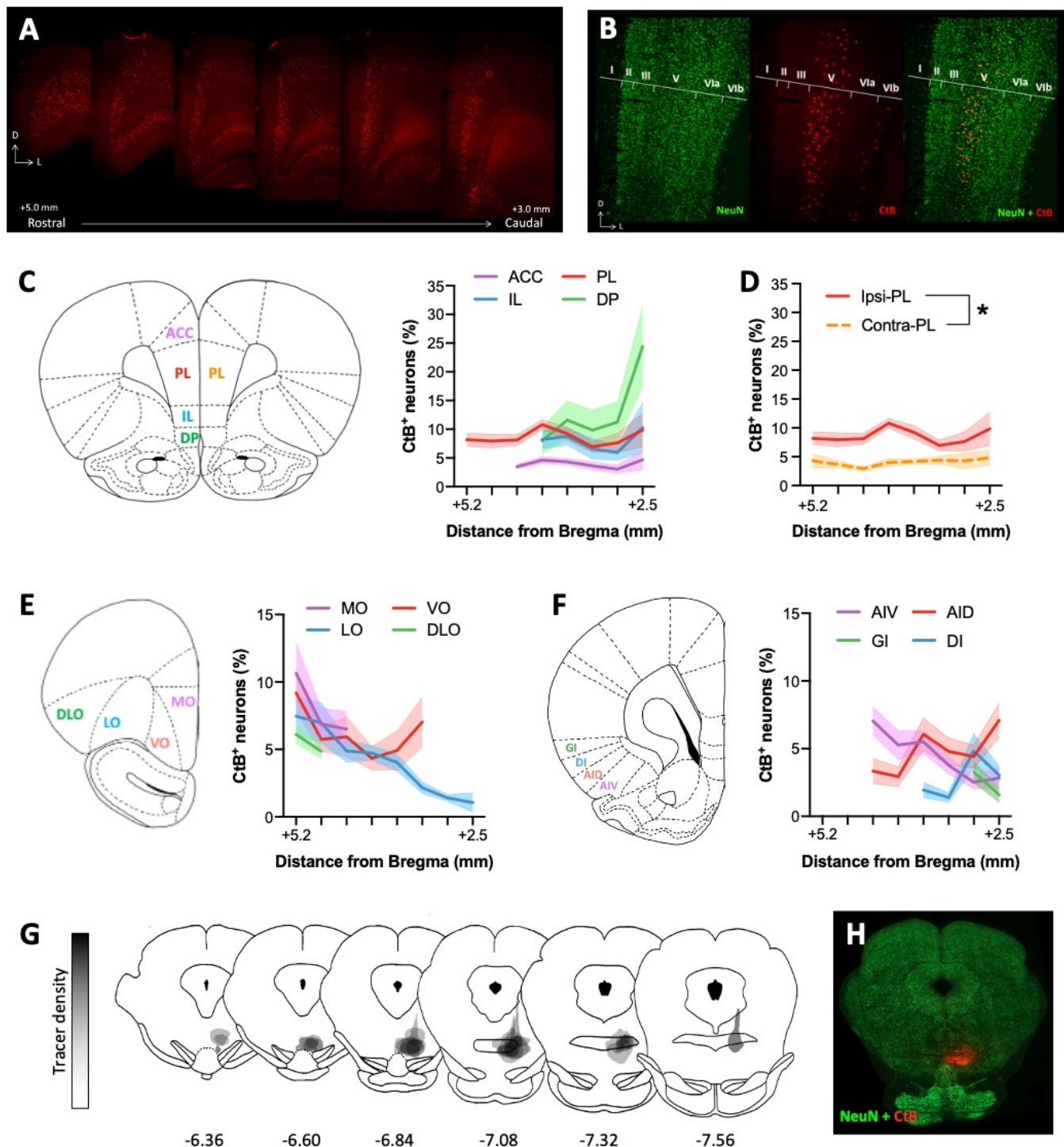


Figure 1. Anatomical distribution of cortical inputs to the RMTg. (A) Representative images demonstrating dense ipsilateral cortical labeling in brain areas injected with CtB into the RMTg. (B) Representative high magnification image showing that inputs to the RMTg arise primarily from layer V of the mPFC. (C) The percent of CtB+ neurons relative to all layer V NeuN+ neurons is relatively consistent across ACC, PL, and IL subregions of the mPFC whereas the density of RMTg-projecting DP mPFC neurons increases substantially at more caudal levels. (D) Contralateral cortical afferents are substantially less dense than ipsilateral inputs as exemplified by a comparison of RMTg-projecting PL mPFC neurons in both hemispheres. (E) The density of layer V OFC neurons projecting to the RMTg is similar to that observed in the mPFC with LO inputs diminishing at more caudal levels. (F) CtB labeling is consistently observed in the AIC, albeit to a lesser degree than that observed in mPFC and OFC. (G) Map of tracer injection sites for all animals included in quantification. (H) Representative injection site. Abbreviations: ACC = anterior cingulate cortex; AID = agranular insular cortex, dorsal; AIV = agranular insular cortex, ventral; DI = dysgranular insular cortex; DLO = dorsolateral orbitofrontal cortex; DP = dorsopeduncular cortex; GI = granular insular cortex; IL = infralimbic cortex; LO = lateral orbitofrontal cortex; MO = medial orbitofrontal cortex; PL = prelimbic cortex; VO = ventral orbitofrontal cortex.

subregions projecting to the RMTg (AID: $4.77 \pm 0.65\%$; AIV: $4.49 \pm 0.72\%$). By contrast, CtB labeling was approximately half that of agranular AIC subregions in the dysgranular (DI; $2.8 \pm 0.76\%$) and granular (GI; $2.43 \pm 0.84\%$) cortices.

As is often the case, CtB labeling was most dense in the hemisphere ipsilateral to the injection site with substantially less labeling apparent in the contralateral cortex. To measure this directly, we quantified the density of layer V RMTg-projecting neurons in the contralateral PL mPFC and found that contralateral cell density was approximately half that of the ipsilateral projection ($4.07 \pm 0.20\%$) (**Figure 1D**). A two-way RM ANOVA comparing cell density between ipsilateral and contralateral hemispheres across the rostrocaudal extent of the PL mPFC confirmed that significantly fewer RMTg-projecting cells arise in the contralateral compared to ipsilateral hemisphere regardless of rostrocaudal level [main effect of hemisphere: $F(1,16)=45.09$, $p<0.0001$].

Layer V cortical efferents are typically excitatory in nature, however, recent work has revealed the presence of long-range GABAergic projection neurons arising from various cortical regions including the mPFC (Lee et al., 2014; Basu et al., 2016; Rock et al., 2018). To determine the neurochemical composition of RMTg-projecting cortical neurons, CtB-expressing slices adjacent to those used in the cell density analysis were labeled with CaMKII α or GAD67. As shown in **Figure 2**, CtB-labeled neurons were predominantly CaMKII α +. In contrast, virtually no overlap in expression was observed between CtB and the GABAergic marker, GAD67. These data indicate that RMTg-projecting cortical neurons are excitatory projection neurons.

A number of cortico-subcortical projections purported to be involved in reward and aversion arise in layer V of the dmPFC. To examine whether RMTg-projecting dmPFC neurons may also overlap with populations of other subcortically projecting layer V neurons, neurons in this projection from

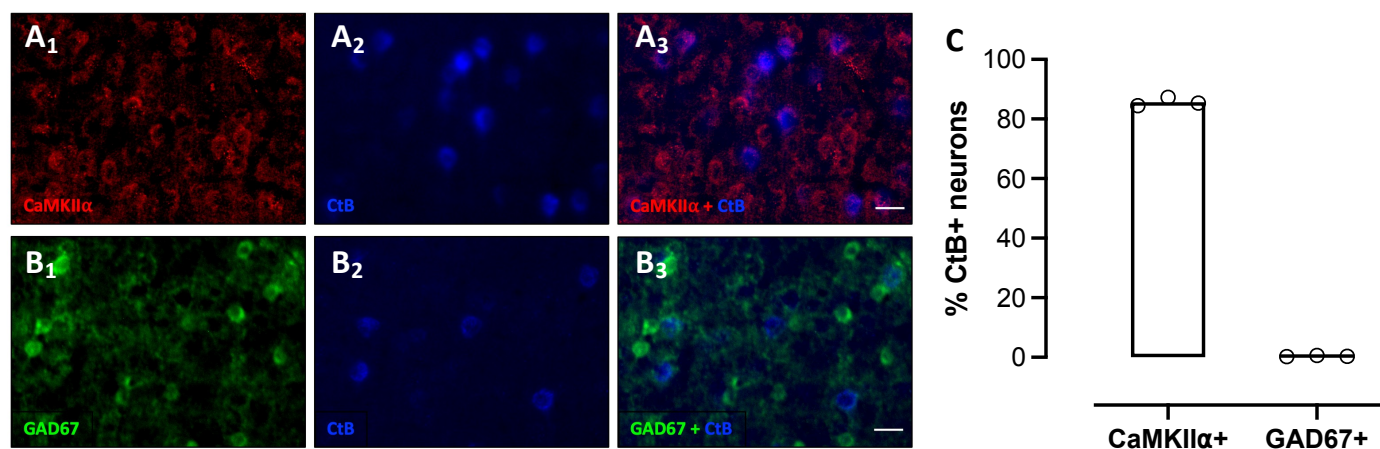


Figure 2. RMTg-projecting dmPFC neurons express CaMKIIα. Representative mPFC images co-labeled for (A₁₋₃) the glutamatergic marker CaMKIIα (red) and CtB (blue) and the (B₁₋₃) GABAergic marker GAD67 (green) and CtB (blue) from rat injected with CtB into the RMTg. (C) Quantification of co-labeling reveals that RMTg-projecting neurons are CaMKIIα⁺. Scale bar = 25 μm.

four rats were selectively filled with green fluorescent protein using an intersectional, dual-virus approach (**Figure 3A**). Labeling was absent in one rat that was, therefore, excluded from analysis. In the remaining three rats, labeling was targeted to the dmPFC, was restricted to the injected hemisphere, and was not apparent in cell bodies outside of the dmPFC indicating successful isolation of the dmPFC-RMTg circuit (**Figure 3B**). Dense punctate labeling, indicative of synaptic terminals, was evident in a number of subcortical regions. Quantification of staining density relative to background (**Figure 3C-D**) revealed the greatest density of collaterals in the dorsomedial striatum ($11.31 \pm 3.11\%$), whereas the dorsolateral striatum was virtually devoid of labeling ($0.07 \pm 0.01\%$). Ventrally, RMTg-projecting dmPFC neurons collateralized to a moderate degree in the nucleus accumbens core ($6.03 \pm 1.31\%$) as well as the shell ($0.34 \pm 0.19\%$), albeit very weakly. Substantial collateralization was also observed in the ventral pallidum ($11.05 \pm 3.98\%$), hypothalamus ($8.48 \pm 0.98\%$), periaqueductal gray ($6.54 \pm 1.50\%$), and lateral preoptic nucleus ($6.20 \pm 0.74\%$). Terminal labeling was much less dense in the lateral habenula ($5.15 \pm 1.55\%$) and ventral tegmental area ($2.58 \pm 1.07\%$) – two regions heavily interconnected with both the dmPFC and the RMTg. By comparison, terminal labeling in the RMTg itself was $3.58 \pm 0.17\%$. The amygdala also receives significant input from layer V dmPFC neurons and, like the RMTg, is well-known for its role in avoidance and aversive signaling. Despite this, only very weak terminal labeling was apparent in this region ($0.94 \pm 0.08\%$). The dorsal hippocampus (Hipp) and secondary somatosensory cortex (S2) were used as negative controls as it is well-known that these regions do not receive any input from the dmPFC (Hipp: $0.02 \pm 0.01\%$; S2: $0.01 \pm 0.00\%$).

Selective stimulation of RMTg-projecting dmPFC neurons drives avoidance behavior

To begin to investigate whether dmPFC inputs to the RMTg play a significant role in aversive signaling, in-vivo optogenetics was used to measure real-time place preference in response to activation of this neural circuit. Testing on day 1 revealed that stimulation of dmPFC terminals in the RMTg resulted in significant avoidance of the light-paired compartment relative to chance

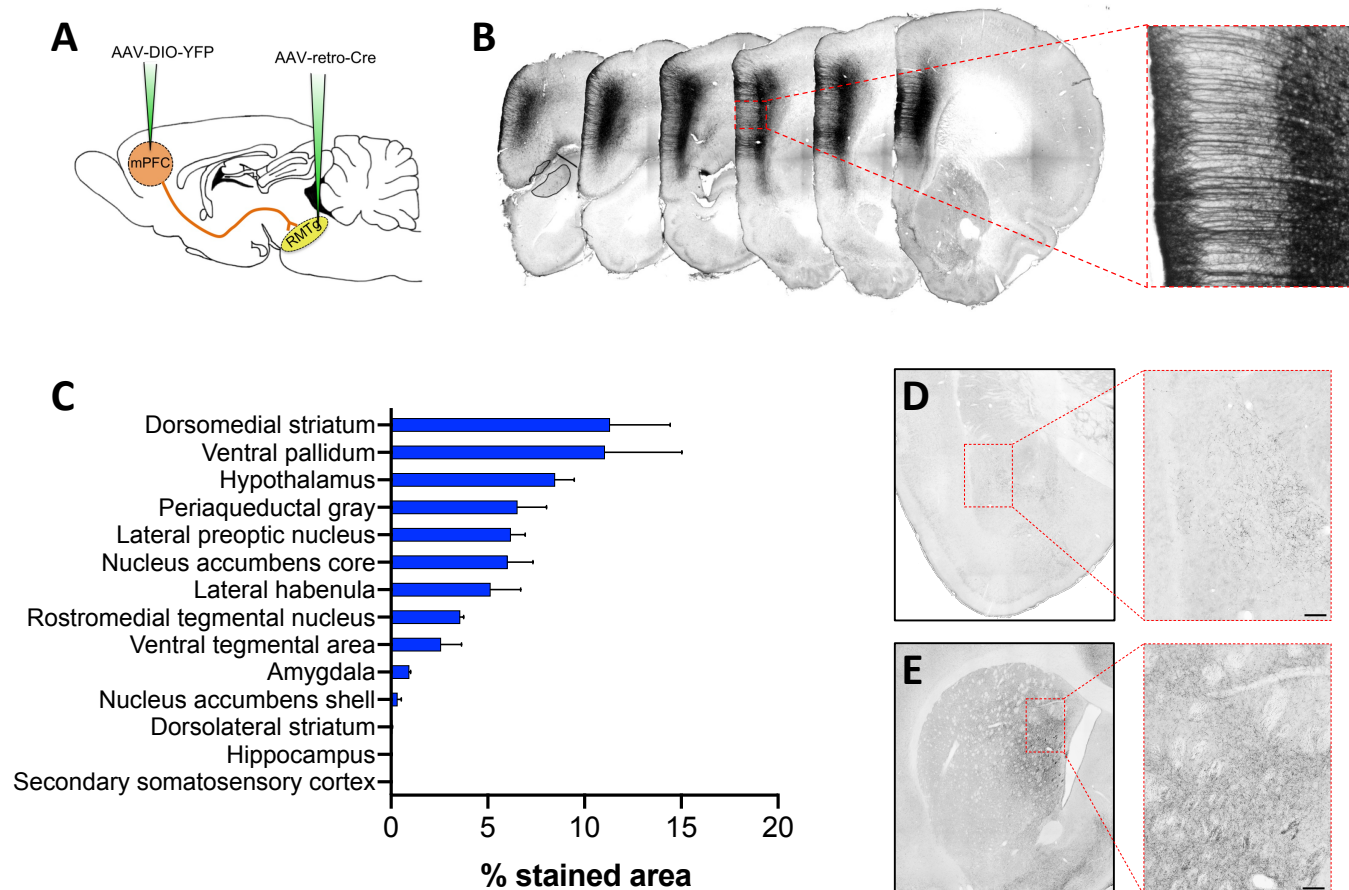


Figure 3. RMTg-projecting dmPFC neurons collateralize throughout the brain. (A) An intersectional dual-virus approach was used to fill RMTg-projecting dmPFC neurons with yellow fluorescent protein (YFP). (B) Representative images showing RMTg-projecting dmPFC neurons filled with YFP following amplification using standard immunohistochemistry. (C) Quantification of the average percent-stained area within ROIs placed within the respective brain regions. (D) Representative YFP staining in the amygdala shows relatively sparse collateralization of RMTg-projecting dmPFC neurons in the basolateral nucleus. (E) Representative YFP staining in the striatum shows dense collateralization in the dorsomedial but not dorsolateral striatum. Scale bar = 100 μ m.

(**Figure 4B**). This effect was replicated during testing on day 2 when the light-paired compartment was reversed [One-way ANOVA test 1 x test 2 x chance: $F(1.95, 7.75) = 22.74$; $p = 0.0006$]. The magnitude of this avoidance was similar to that observed during stimulation of LHb terminals in the RMTg (**Figure 4C**), which was also significantly lower than chance [One-way ANOVA test 1 x test 2 x chance: $F(1.59, 7.95) = 9.60$; $p = 0.0095$]. In contrast, stimulation of dmPFC terminals in the neighboring VTA resulted in neither preference nor avoidance of the light-paired compartment (**Figure 4D**) on either day 1 or day 2 [One-way ANOVA test 1 x test 2 x chance: $F(1.04, 3.11) = 0.095$; $p = 0.7866$]. Direct comparison of the effect of each circuit manipulation on real-time place preference revealed that stimulation of either dmPFC or LHb inputs to the RMTg drove avoidance behavior that was significantly different from stimulation of dmPFC inputs to the VTA (**Figure 4E**) [One-way ANOVA: $F(2, 12) = 7.30$, $p = 0.0084$]. Altogether, these data indicate that, similar to the LHb-RMTg projection, activation of dmPFC inputs to the RMTg provides an aversive signal to promote avoidance behavior.

RMTg-projecting mPFC neurons are activated following exposure to aversive stimuli

While optogenetic stimulation of dmPFC-RMTg neurons demonstrates that activation of this pathway can facilitate avoidance, we are unable to conclude from these data that neurons in this circuit are indeed active during avoidance and/or during similar behavioral responses to aversive stimuli. To explore this possibility, rats were euthanized 90 min following exposure to either neutral or aversive stimuli following injection of CtB into the RMTg (**Figure 5A**). Two groups of rats were exposed to a series of tones and foot shocks. In one group, tones were predictive of shock as in a standard fear conditioning paradigm. In contrast, in the other group, rats were exposed to the same number of tones and shocks but in an unpaired manner such that tones were not predictive of shock. As expected, rats in the Shock-paired tone group exhibited significantly greater freezing in response to tone presentation on test day than rats in the Shock-unpaired group (t-test; $p = 0.0005$; **Figure 5B**). Behavioral data was not collected on the two remaining groups of rats

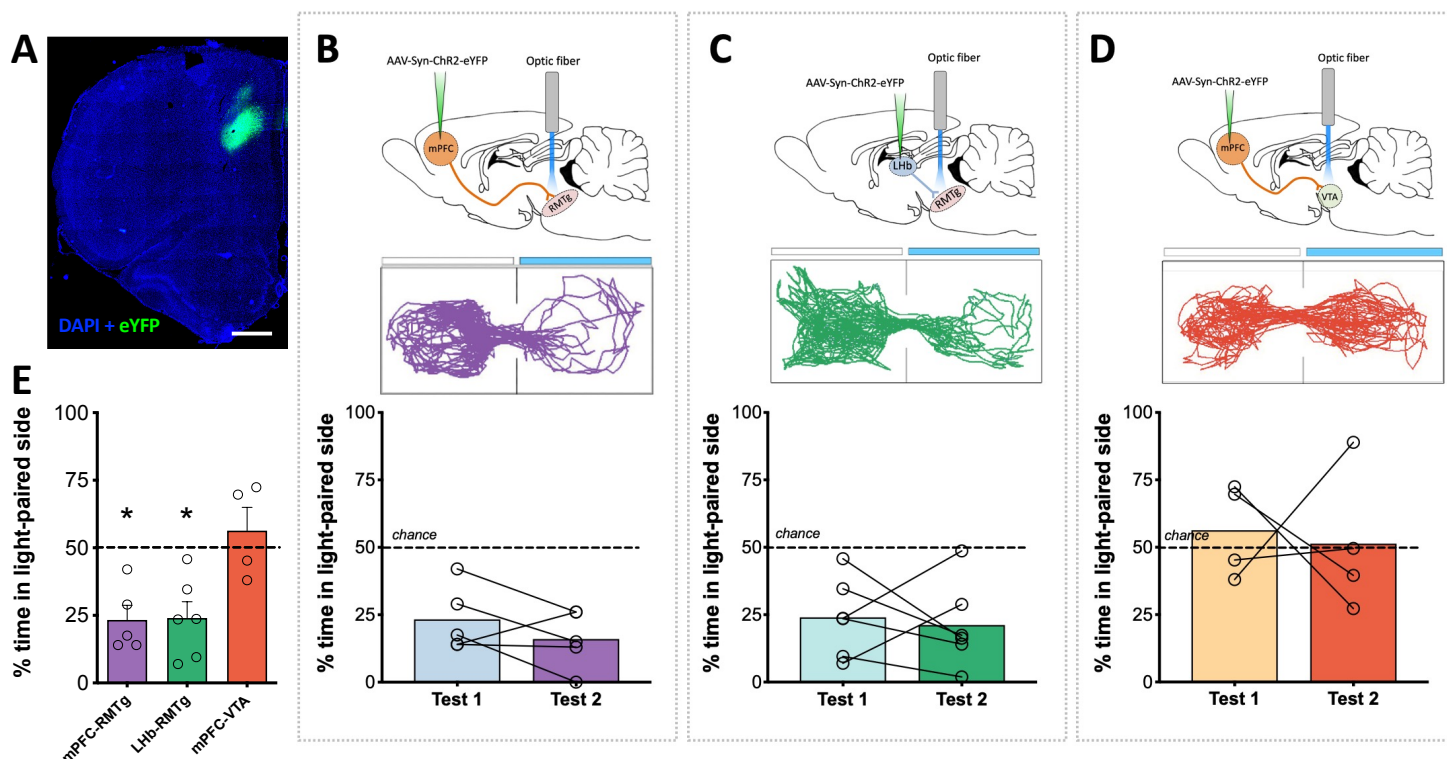


Figure 4. Optogenetic stimulation of RMTg-projecting dmPFC terminals drives avoidance. (A) Representative ChR2 expression in dmPFC. (B) Rats spend significantly less time relative to chance in the light-paired side of a two-compartment chamber during initial testing (test 1) and when the light-paired compartment is reversed (test 2) when light delivery results in stimulation of dmPFC terminals in the RMTg. (C) A similar degree of avoidance of the light-paired chamber is observed upon stimulation of lateral habenula inputs to the RMTg. (D) Unlike stimulation of dmPFC terminals in the RMTg, stimulation of dmPFC terminals in the VTA fails to produce either preference for or avoidance of the light-paired compartment. (E) Direct comparison of circuit manipulations reveals significant avoidance when stimulating inputs to the RMTg relative to the VTA. Light-paired side indicated by blue bar in representative maps above each dataset. * $p < 0.01$, scale bar = 1000 μ m.

exposed to either the neutral testing context or a series of foot shocks (without tone presentation). A one-way ANOVA comparing the magnitude of cFos expression in CtB⁺ neurons in the mPFC revealed a significant effect of treatment condition on cFos induction (**Figure 5C-D**) [$F(3,32)=11.00$, $p<0.0001$]. Post-hoc comparisons revealed that cFos expression was significantly greater in RMTg-projecting mPFC neurons of rats that were exposed to a series of either foot shocks or tones predictive of shocks relative to rats exposed to the neutral testing context (shock: $p<0.0001$; shock-paired tone: $p=0.006$). The magnitude of cFos expression in CtB⁺ mPFC neurons in shock-exposed rats was also significantly greater than was observed in rats exposed to tones that were explicitly unpaired with shocks ($p=0.004$). In combination with results from the real-time place preference testing, these data suggest that RMTg-projecting mPFC neurons are activated in response to learned and unlearned aversive stimuli and may play a role in regulating the behavioral response to such stimuli.

Functional & structural changes in RMTg-projecting dmPFC neurons following exposure to aversive stimuli

Given the above evidence that dmPFC inputs to the RMTg are activated in response to aversive stimuli, we next investigated the potential impact that exposure to such stimuli has on plasticity in this neural circuit (**Figure 6A**). A two-way repeated measures ANOVA of spiking measured during whole-cell patch-clamp recordings from RMTg-projecting dmPFC neurons revealed a significant interaction between current step and stimulus exposure [$F(15,315)=22.08$, $p<0.0001$] such that spike frequency was significantly reduced as current injection increased beyond 200 pA in rats exposed to the same foot shock procedure that induced significant cFos expression relative to Context controls (Sidak correction; all p values ≤ 0.03 ; **Figures 6B-C**). This effect was accompanied by a significantly higher rheobase (t-test; $p<0.0001$), significantly lower membrane resistance (t-test; $p<0.0001$), and higher membrane capacitance (t-test; $p<0.0434$) in shock-exposed rats compared to context controls. No significant difference in action potential threshold

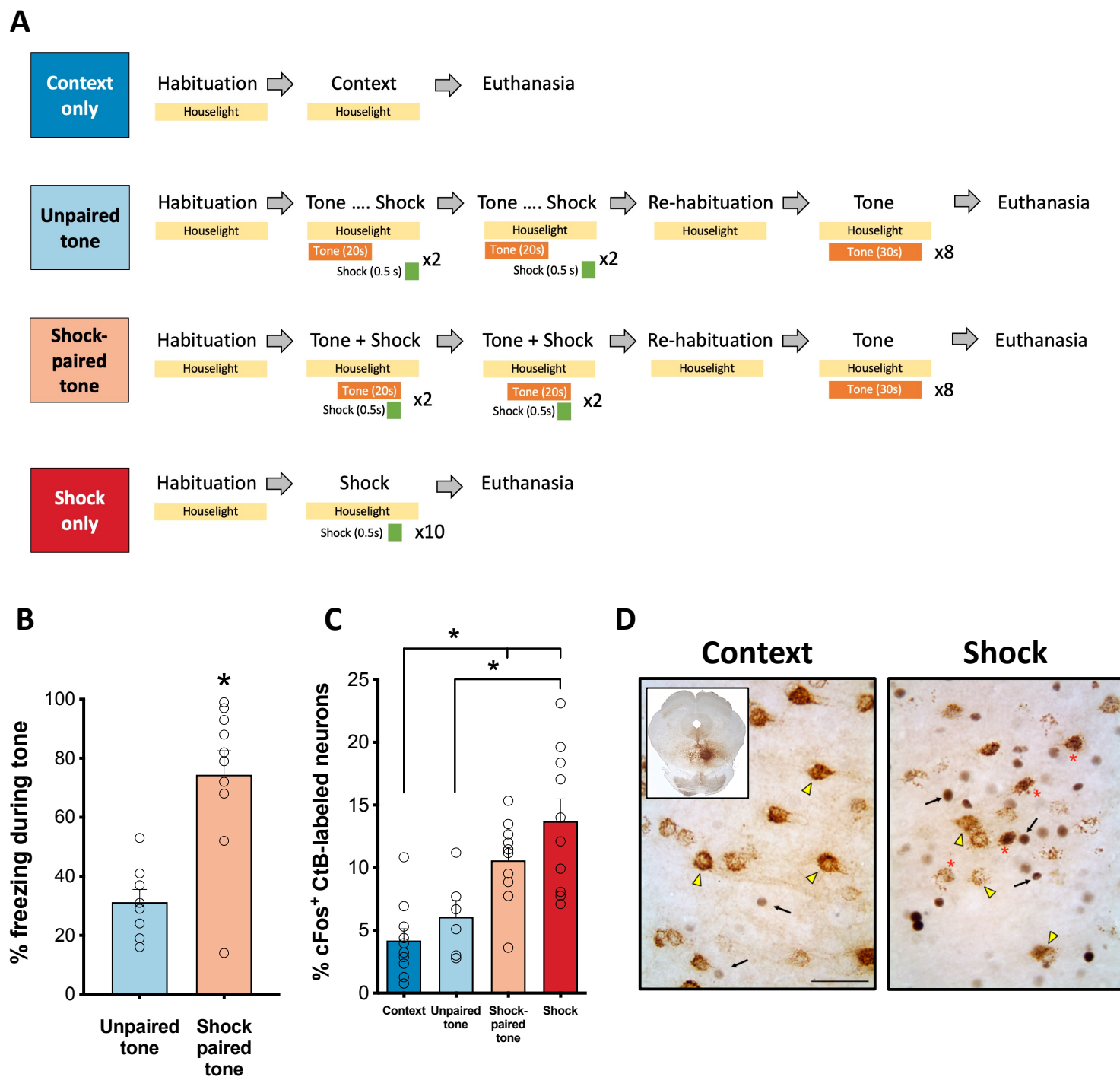


Figure 5. cFos induction in RMTg-projecting dmPFC neurons following exposure to aversive stimuli. (A) Experimental procedures. **(B)** Rats that had tone paired with foot shock delivery displayed significantly more freezing behavior in response to tone presentation than rats that were exposed to the same number of tone-shock presentations but in an unpaired manner. **(C)** Significantly greater cFos expression was observed in RMTg-projecting dmPFC neurons (CtB⁺) following exposure to either a series of foot shocks or a tone predictive of foot shock relative to a neutral tone or the testing context alone. **(D)** Representative images of CtB and cFos labeling in the dmPFC of a context-exposed rat and a rat exposed to foot shock. CtB⁺/cFos⁻ neurons are indicated with a yellow arrowhead; CtB⁻/cFos⁺ neurons are indicated with a black arrow; CtB⁺/cFos⁺ neurons are indicated by a red asterisk. Scale bar = 200 μ m.

was observed between groups (t-test; $p=0.1555$; **Figures 6D-G**). Action potential duration and amplitude were significantly different between groups with Shock-exposed rats exhibiting action potentials of greater amplitude (t-test; $p=0.0232$) and shorter duration (t-test; $p=0.0002$) than Context-exposed rats (**Figures 6H-I**). No significant difference in action potential after-hyperpolarization was observed between groups (t-test; $p=0.2625$; **Figure 6J**). Overall, these data are indicative of decreased intrinsic excitability in RMTg-projecting dmPFC neurons following exposure to an aversive stimulus.

To examine the impact of exposure to aversive stimuli on structural plasticity, we next quantified dendritic spine density and morphology in RMTg-projecting dmPFC neurons in rats exposed to either foot shock or a neutral context (**Figures 7A-B**). T-tests were used to analyze differences in dendrite diameter and volume as well as dendritic spine density collapsed across spine class. Two-way ANOVAs were used to analyze spine density and morphology by spine class between groups. No significant differences in dendrite diameter, volume or overall spine density were observed between Context- and Shock-exposed rats (p values > 0.50 ; **Table S1**). Analysis of spine density revealed a main effect of spine class [$F(3,40)=53.37$, $p<0.0001$] but no main effect of stimulus exposure [$F(1,40)=0.075$, $p=0.7856$] or interaction between the two factors [$F(3,40)=1.34$, $p=0.2756$]. Tukey corrected post-hoc comparisons of the main effect of spine class revealed that both Context- and Shock-exposed rats had a significantly greater density of mushroom-shaped spines relative to all other spine classes (all p values < 0.0001 ; **Figure 7C**). Of the eight other measures of spine morphology analyzed, no significant between-group differences were observed in dendritic spine length, diameter, or volume. Similarly, no significant between-group differences in spine terminal point diameter or volume were observed. Spine neck volume was not significantly different between groups either (**Table S1**). In contrast, a significant main effect of stimulus exposure [$F(1,33)=9.85$, $p=0.0036$] was observed for spine neck diameter in the absence of a significant interaction between stimulus exposure and spine class

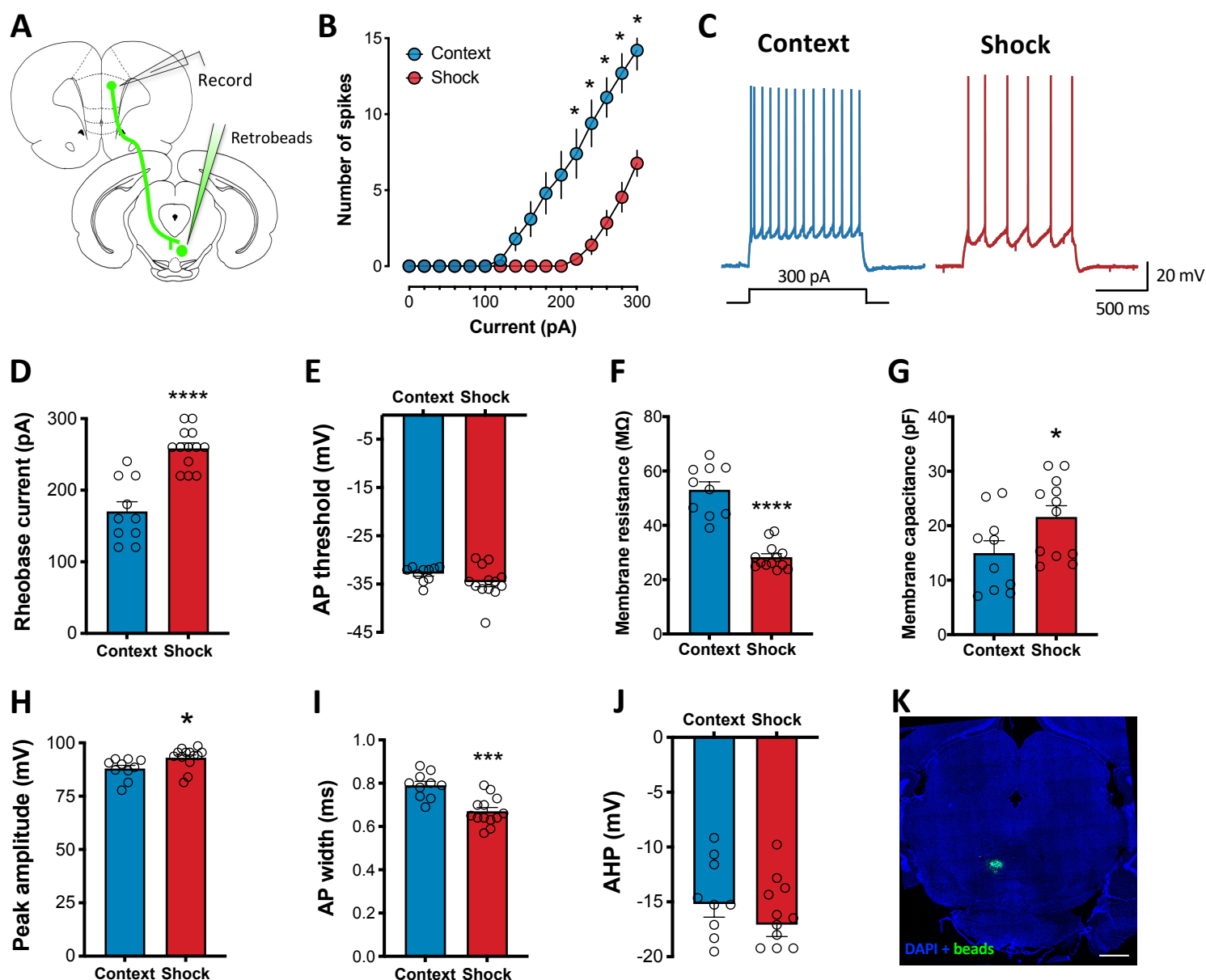


Figure 6. Decreased excitability in RMTg-projecting dmPFC neurons following exposure to aversive stimuli. (A) Experimental preparation. (B) Significantly fewer spikes were observed in shock-exposed rats relative to controls in current clamp recordings of retrobead-labeled dmPFC neurons. (C) Representative traces from a control and shock-exposed rat. Decreased spiking was associated with a significant increase in (D) rheobase, (G) membrane capacitance, and (H) peak action potential amplitude as well as a significant decrease in (F) membrane resistance and (I) action potential half-width. No significant difference was observed in (E) action potential threshold or (J) after-hyperpolarization. (K) Representative retrobead injection site in the RMTg. * $p \leq 0.05$, scale bar = 1000 μm .

[F(3,33)=1.69, p=0.1884] indicative of greater spine neck diameter in Shock-exposed rats across all spine classes compared to Context-exposed rats (**Figure 7D**). A significant main effect of spine class was also uncovered with post-hoc analyses revealing that for both context- and shock-exposed rats, long, thin spines had significantly greater spine neck diameter than other spine classes (all p values ≤ 0.002). Differences in spine neck length were also observed between Context- and Shock-exposed rats (**Figure 7E**) with significant main effects of both spine class [F(3,29)=180.60, p<0.0001] and stimulus exposure [F(1,290)=4.19, p=0.0498]. While a significant interaction between the two factors was also uncovered [F(3,29)=3.18, p=0.0388], Sidak's multiple comparisons failed to identify significant differences in spine neck length within any given spine class between Context- and Shock-exposed rats (all p values > 0.05).

RMTg-projecting dmPFC neurons express both D1 and D2 dopamine receptors

D1 and D2 dopamine receptors play important roles in prefrontal regulation of behavioral flexibility and decision-making (Floresco, 2013). A number of studies suggest that, similar to their distribution in the striatum, D1- and D2-expressing neurons in the mPFC are anatomically and functionally distinct cell populations (e.g., Gaspar et al., 1995). RNAScope for D1 and D2 receptor mRNA was used in combination with fluorescent retrograde tracing to investigate whether RMTg-projecting dmPFC neurons exhibit a distinct dopamine receptor expression profile and whether dopamine receptor gene expression was altered in these neurons following exposure to an aversive stimulus. As shown in **Figures 8A-F**, fluorescent retrobead-labeled cells (bead+) comprised approximately 62% of the total population of cells analyzed. A two-way ANOVA indicated that there was no significant difference in the number of either total or bead+ cells between control and shock-exposed rats [Main effect of treatment: F(1,12)=0.002, p=0.0966; treatment x cell-type: F(1,12)=0.10, p=0.759]. In addition, no significant between-group differences were observed using a two-way ANOVA to compare the percent of bead+ cells that were also positive for either D1 or D2 mRNA in control and shock-exposed rats [F(3,24)=0.28,

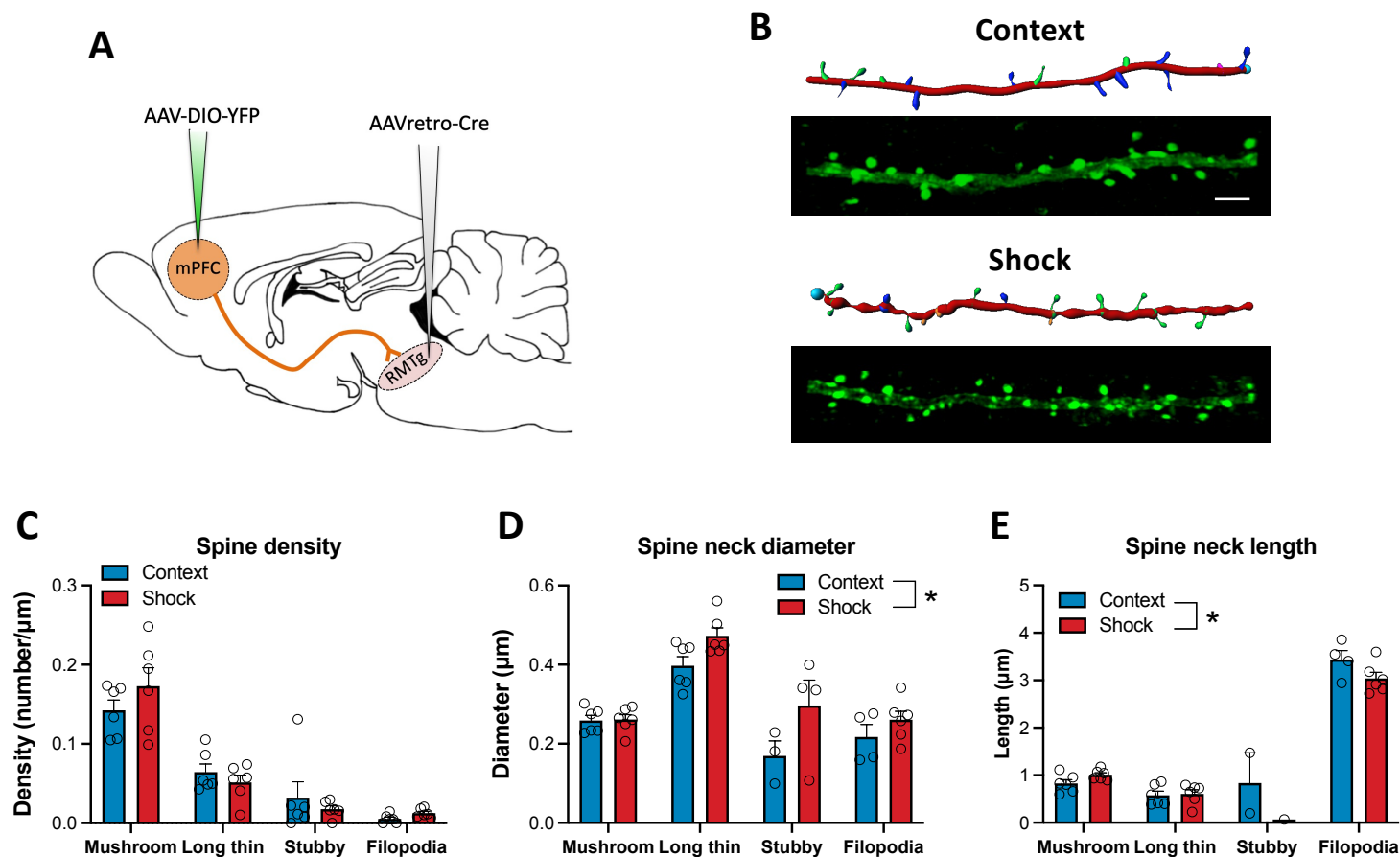


Figure 7. Exposure to aversive stimuli increases spine neck diameter in RMTg-projecting dmPFC neurons. (A) An intersectional dual-virus approach was used to fill RMTg-projecting dmPFC neurons with yellow fluorescent protein (YFP). (B) Representative YFP-filled primary apical dendrites in the dmPFC and accompanying IMARIS renderings for context- and shock-exposed rats. (C) Spine density did not differ between groups regardless of subclass. However, shock-exposed rats exhibited significantly greater spine neck diameter (D) and shorter spine length (E) across all subtypes (main effect of shock) relative to rats exposed to the neutral testing context. * $p \leq 0.05$; scale bar = 5 μm .

p=0.8411] (**Figure 8G**). Collapsing across groups, classification of bead⁺ cells by dopamine receptor mRNA expression revealed that RMTg-projecting dmPFC neurons are predominantly D1 receptor-expressing (87%) with a substantial proportion also expressing D2 receptors (59%) (**Figure 8H**). Only a small proportion of bead⁺ neurons expressed D2 mRNA in the absence of D1 mRNA (5%) and approximately 8% lacked both mRNA for either receptor. To examine whether exposure to an aversive stimulus altered the magnitude of dopamine receptor gene expression in RMTg-projecting dmPFC neurons, two-way ANOVAs were used to compare the average number of RNAScope dots present per cell across dopamine receptor-expressing cell types. As shown in **Figures 8I-J**, foot shock exposure had no significant effect on either D1 (Treatment: $F(1,18)=2.09$, $p=0.1651$; treatment x cell-type: $F(2,18)=0.21$, $p=0.8138$) or D2 (Treatment: $F(1,18)=1.38$, $p=0.2550$; treatment x cell-type: $F(2,18)=0.30$, $p=0.7445$) mRNA expression.

To further explore whether cFos induction observed following exposure to aversive stimuli is specific to a unique dopamine receptor-expressing population of dmPFC-RMTg neurons, we next measured induction of cfos mRNA colocalized with D1 receptor mRNA (the predominantly expressed dopamine receptor in these neurons). As expected, cfos expression was significantly enhanced in Shock-exposed rats relative to rats exposed to a neutral context (**Figure 9A**), and this was true regardless of retrobead labeling [two-way ANOVA main effect of treatment: $F(1,12)=4.72$, $p=0.0505$]. When comparing cfos expression in D1⁺ and D1⁻ RMTg-projecting dmPFC neurons, a two-way ANOVA revealed a main effect of treatment with Shock-exposed rats exhibiting a greater proportion of cfos in both cell types relative to Context controls (**Figure 9B**). However, this effect only trended toward statistical significance for a main effect of shock exposure [$F(1,12)=4.05$, $p=0.0673$]. Similarly, the magnitude of cFos mRNA expression, as measured by average number of cFos dots per cell, was significantly greater in RMTg-projecting dmPFC neurons of Shock-exposed rats compared to controls regardless of D1 receptor expression profile (**Figure 9C**; main effect: $F(1,12)=4.50$, $p=0.0555$). Altogether, these data

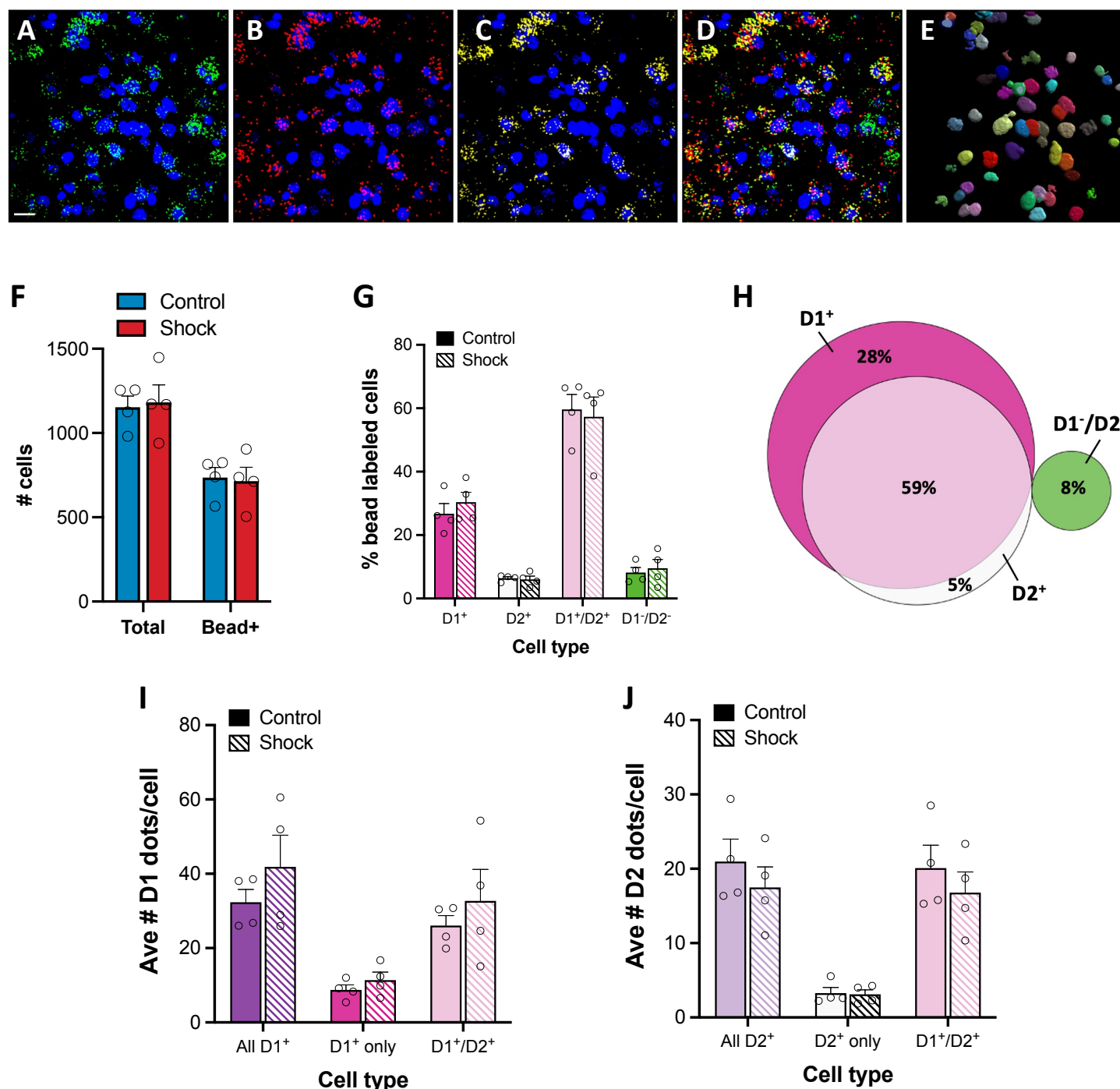


Figure 8. D1 & D2 receptor gene expression in RMTg-projecting dmPFC neurons. (**A-E**) Representative labeling in dmPFC neurons shows dense retrobead labeling in green (**A**) accompanied by D1 (**B**) and D2 (**C**) mRNA transcript dots indicated in red and yellow, respectively. The merged image (**D**) shows heterogeneous overlap of the retrograde beads and mRNA transcripts dots in cells identified using nuclear labeling with DAPI (blue). (**E**) IMARIS rendering of individual cells from A-D showing colocalization of the beads/dots in the 3D rendered soma. Foot shock had no effect on the total number of cells or number of bead positive cells analyzed (**F**). The prevalence of D1 and D2 mRNA transcript containing cells was similar between the treatment groups (**G**). When collapsed across groups (**H**), it is apparent that the majority of RMTg-projecting dmPFC neurons are D1+ with a large proportion of D1+ neurons also expressing D2 receptor mRNA. Foot shock exposure had no significant effect on the magnitude of either D1 (**I**) or D2 (**J**) mRNA expression. Scale bar = 20 μ m.

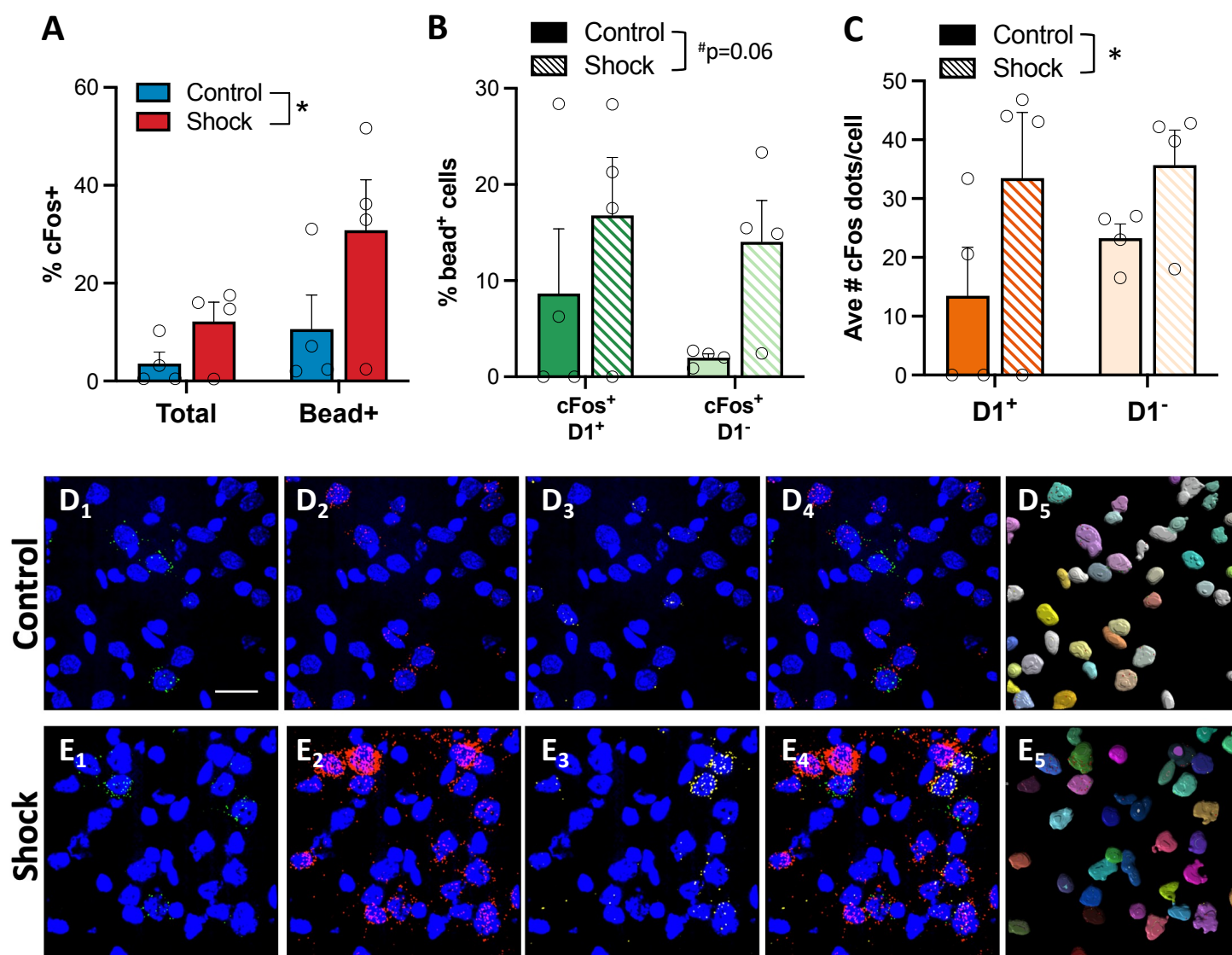


Figure 9. Shock-induced enhancement of cFos expression occurs in D1⁺ and D1⁻ RMTg-projecting dmPFC neurons. (A) Shock exposure significantly increased cFos mRNA expression in both bead⁺ and bead⁻ dmPFC neurons. (B) cFos expression was similarly increased in both D1⁺ and D1⁻ neurons in shock-exposed rats relative to controls. (C) cFos mRNA expression was significantly greater in shock-exposed rats compared to controls, but there was no significant difference between D1⁺ and D1⁻ cell populations. Representative RNAScope labeling from a context-exposed control (D) and shock-exposed (E) rat. Retrobead labeling is depicted in green (D-E₁), cFos mRNA in red (D-E₂), D1 receptor mRNA in yellow (D-E₃), merged image (D-E₄) and IMARIS rendering of colocalized beads and dots the cell soma (D-E₅). Nuclear labeling by DAPI depicted in blue. Scale bar = 20 μ m; *p $\leq$ 0.05.

indicate that RMTg afferents arising in the dmPFC are highly enriched in D1 dopamine receptors (with D2 receptors colocalized to many of these neurons), and that the effects of exposure to aversive stimuli are similar across dmPFC-RMTg neurons with differing dopamine receptor expression profiles.

DISCUSSION

Findings from the present study demonstrate the presence of dense cortical input to the RMTg spanning the entire rostrocaudal extent of the mPFC, OFC and AIC. The greatest density of afferents arises from the neurons in the dmPFC. The neurons are primarily D1⁺ with a large proportion also expressing D2 receptor mRNA. RMTg-projecting dmPFC neurons collateralize extensively in regions critically involved in regulating motivated behavior and flexible decision-making. Stimulation of dmPFC terminals in the RMTg drives avoidance and exposure to aversive stimuli induces cFos expression in this neural circuit. Finally, repeated exposure to an aversive stimulus results in significant alterations in RMTg-projecting dmPFC neurons in the form of increased excitability and changes in spine neck morphology. Together these data suggest that dmPFC neurons play an important role in governing the behavioral response to aversive stimuli.

If anatomical density is any indication of the influence a particular circuit may have over behavior, our data suggest that mPFC inputs to the RMTg are likely to play, at minimum, an equally important role in guiding adaptive responding to environmental stimuli as other heavily researched cortico-subcortical circuits. In a quantitative analysis of cortico-subcortical projection density in the mPFC, Gabbot et al. (2005) reported that ~8% of layer V PL and IL mPFC neurons project to the amygdala, whereas raphe- and PAG-projecting mPFC neurons each account for ~1-5% of neurons in layer V. Despite being sparse relative to PL and IL input to the ventral striatum (~18%), for example, subsequent work has implicated each of these discrete circuits in crucial aspects of motivated behavior (e.g., Rozeske et al., 2011; Warden et al., 2012; Bukalo et al., 2015). By

comparison, the density of RMTg-projecting PL and IL neurons observed in the current study (~10%) is one of the denser subcortical projections arising from layer V of the mPFC. A large body of work has shown that the PL and IL mPFC exert opposing effects on many types of behavior. For example, PL mPFC neurons facilitate behavioral responding in Pavlovian and Instrumental assays associated with either appetitive or aversive outcomes, whereas IL mPFC activity has been shown to facilitate extinction of such behavioral responses (Peters et al., 2009; Gourley and Taylor, 2016). While the current study showed that stimulation of dmPFC terminals in the RMTg (which includes the PL subregion) facilitate avoidance, it remains unknown whether the IL-RMTg projection regulates RMTg signaling in an inverse manner that is similar to what is often observed when PL and IL mPFC are manipulated at a regional level. Unlike the PL and IL mPFC, the DP mPFC has been largely neglected with very few studies investigating this region at either anatomical or functional levels. In one of the few existing DP studies, Kataoka et al (2020) revealed a role for DP mPFC inputs to the dorsomedial hypothalamus in sympathetic stress response and stress-induced avoidance of social interactions. Combined with the current analysis, which reveals a remarkably dense projection from the DP mPFC to the RMTg that increases to an astonishing degree in the caudal mPFC, these data highlight the need for further investigation into the role of both the DP mPFC and its connections with the RMTg in aversion.

Consideration of collateral input is particularly important when investigating circuit function, as the possibility that manipulations of circuit activity affect signaling in sites that receive collaterals has the potential to influence interpretation of results. Using an intersectional, virally-mediated approach, our data reveal dense collateralization of RMTg-projecting dmPFC neurons to a number of regions critically involved in motivated behavior. While the extent of collateralization may be somewhat surprising, this frequently underappreciated aspect of neuronal structure is not uncommon. Indeed, recent methodological advancements have enabled researchers to map the extent of a single neuronal projection throughout the brain and demonstrate that collateralization

is often widespread (Economo et al., 2016; Kebschull et al., 2016). The current findings indicate that RMTg-projecting dmPFC neurons collateralize most densely in the dorsal striatum. Interestingly, dmPFC afferents to the dorsal striatum are well-characterized for their role in guiding goal-directed behavior (Simmler and Ozawa, 2019) including avoidance (Loewke et al., 2021). Previous work found that, relative to a number of other cortico-subcortical projections, dmPFC input to the dorsal striatum was among the densest of projections, comprising ~19% of layer V neurons (Gabbott et al., 2005). Thus, it is not necessarily surprising that there is overlap in the population of dorsal striatum-projecting dmPFC neurons and those of other subcortical afferents. Unexpectedly, terminal density in the RMTg itself was relatively low by comparison to other brain regions. However, it should be noted that collaterals are often comprised of very thin branches (Rockland, 2013), and as a result the density measurement obtained in the present study (percent area stained) may not provide a full picture of the extent of collateralization of this projection. It is also unclear from the present data whether the observed collateralization was indicative of dense arborization of a select few dmPFC neurons, or of a high number of dmPFC cells each providing relatively weak collateral input to a given region. Recent work reporting very little overlap in cell body labeling between NAc- and RMTg-projecting mPFC neurons using dual retrograde tracer approach (Cruz et al., 2021) suggests that the former may be the more likely scenario. Additional experiments using multiple retrograde tracers to examine overlap in dmPFC cell body labeling will be essential to understand the potential functional implications of synergistic neurotransmission in regions receiving collateral input. Certainly, the current data present intriguing possibilities for coordinated signaling across brain regions involved in guiding behavioral responding to environmental stimuli.

The results of the present study also revealed that RMTg-projecting dmPFC neurons are glutamatergic and predominantly express D1 dopamine receptor mRNA. Of note, D2 receptor mRNA is also colocalized in the majority of these neurons. These findings agree in large part with

existing data showing that D1 receptor expression is greater than that of D2 in the mPFC (Santana et al., 2009). While D1 and D2 receptor-expressing neurons are often thought of as discrete cell populations, a number of studies have observed colocalization of both receptors, particularly in layer V of the mPFC where D2 receptors are most abundant (Vincent et al., 1995; Gaspar et al., 1995; Santana et al., 2009). Recent work has highlighted the importance of dopaminergic regulation of cortical control in aversive signaling (Vander Weele et al., 2018; Huang et al., 2020). Of particular interest is data suggesting that dopamine signaling alters mPFC responses to aversive stimuli by altering the signal-to-noise ratio of incoming sensory inputs (Vander Weele et al., 2018). Whether this dopaminergic modulation is circuit- or cell-type specific is not well-understood. Nevertheless, a rich literature demonstrates that D1 and D2 receptors regulate behavioral flexibility in complex ways in the mPFC (Floresco and Magyar, 2006). The expression and function of both receptor subtypes is frequently mechanistically linked to neuropsychiatric illnesses characterized by deficits in decision-making including schizophrenia, addiction, and anxiety disorders (Volkow et al., 2004; Perez de la Mora et al., 2012; McCutcheon et al., 2019). While foot shock exposure did not significantly affect D1 or D2 mRNA expression in dmPFC-RMTg neurons in the current study, it is possible that repeated or prolonged insults uniquely affect dopamine receptor modulation of this neural circuit or that other measures of D1/D2 receptor function not explored in the current study are altered by exposure to aversive stimuli. Thus, the role that dopaminergic regulation of dmPFC-RMTg circuitry plays in aversive signaling and how it is altered in models of neuropsychiatric illness present intriguing areas for future investigation.

The present study focused on the potential role of dmPFC-RMTg circuit in aversion based on previous work characterizing the RMTg for its involvement in aversive signaling and the functional overlap exhibited by the dmPFC. However, some data suggests that PL-RMTg neurons respond to both aversive and rewarding stimuli (Li et al., 2019a). In addition, recent work suggests that activity in this neural circuit is particularly crucial when responding to conditioned rather than

unconditioned stimuli (Li et al., 2019b; Cruz et al., 2020). It is therefore clear that additional work is needed to determine the potential diversity of signals encoded by these neurons and how this information differs across afferents arising from various mPFC subregions (i.e., PL vs IL).

Somewhat unexpectedly, exposure to repeated foot shock resulted in a significant decrease in excitability in RMTg-projecting dmPFC neurons. This contrasts with what has been observed in LHb neurons, the densest source of input to the RMTg (Jhou et al., 2009b), which exhibit a significant increase in excitability following a similar foot shock exposure paradigm (Lecca et al., 2016). Interestingly, the decrease in excitability observed in the present study was accompanied by significant changes in neck length and diameter of spines localized to the primary apical dendrites of RMTg-projecting dmPFC neurons. Dendritic spines are the primary recipients of incoming excitatory signals in pyramidal neurons. Spine neck morphology, in particular, plays a fundamental role in compartmentalizing electrical and biochemical signals in the head of the spine. In particular, spine neck diameter has been identified as the single greatest contributor to such compartmentalization (Tønnesen et al., 2014). In combination with data demonstrating an inverse relationship between spine neck diameter and excitatory potential (Araya et al., 2014), our data suggest a potential reduction in the synaptic strength of inputs to RMTg-projecting dmPFC neurons in shock-exposed rats relative to controls. Although speculative, given the role that layer V cortical apical dendrites are thought to play in modulating cortical oscillations (LaBerge and Kasevich, 2013), it is possible that the observed change in spine neck morphology alters oscillatory frequency in this neuronal population. Altogether, these data suggest a loss of top-down modulation of RMTg activity following exposure to aversive stimuli. Whether these physiological and structural adaptations serve to promote adaptive responding to future aversive stimuli is an important avenue for future exploration.

In conclusion, the current work presents a fresh perspective on the degree of subregion- and circuit-specific cortical regulation of RMTg-mediated aversive signaling. These data provide a strong foundation from which future studies can begin to dissect out the distinct (or complementary) roles of parallel cortico-subcortical circuits involved in motivated behavior. How these circuits are altered in models of neuropsychiatric illness will be crucial for understanding the neural mechanisms underlying disruptions in the balance of neural signals mediating reward and aversion that is altered in a number of disease states.

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REFERENCES

- Alexander WH, Brown JW (2019) The Role of the Anterior Cingulate Cortex in Prediction Error and Signaling Surprise. *Top. Cogn. Sci.* 11:119–135
- Araya R, Vogels TP, Yuste R (2014) Activity-dependent dendritic spine neck changes are correlated with synaptic strength. *Proc. Natl. Acad. Sci. U. S. A.* 111:E2895-2904
- Basu J, Zaremba JD, Cheung SK, Hitti FL, Zemelman BV, Losonczy A, Siegelbaum SA (2016) Gating of hippocampal activity, plasticity, and memory by entorhinal cortex long-range inhibition. *Science* 351:aaa5694
- Bukalo O, Pinard CR, Silverstein S, Brehm C, Hartley ND, Whittle N, Colacicco G, Busch E, Patel S, Singewald N, Holmes A (2015) Prefrontal inputs to the amygdala instruct fear extinction memory formation. *Sci. Adv.* 1:e1500251
- Burgos-Robles A, Vidal-Gonzalez I, Quirk GJ (2009) Sustained conditioned responses in prelimbic prefrontal neurons are correlated with fear expression and extinction failure. *J. Neurosci. Off. J. Soc. Neurosci.* 29:8474–8482
- Centanni SW, Morris BD, Luchsinger JR, Bedse G, Fetterly TL, Patel S, Winder DG (2019) Endocannabinoid control of the insular-bed nucleus of the stria terminalis circuit regulates negative affective behavior associated with alcohol abstinence. *Neuropsychopharmacol. Off. Publ. Am. Coll. Neuropsychopharmacol.* 44:526–537
- Corbett D, Wise RA (1980) Intracranial self-stimulation in relation to the ascending dopaminergic systems of the midbrain: a moveable electrode mapping study. *Brain Res.* 185:1–15
- Corcoran KA, Quirk GJ (2007) Activity in prelimbic cortex is necessary for the expression of learned, but not innate, fears. *J. Neurosci. Off. J. Soc. Neurosci.* 27:840–844
- Cruz AM, Kim TH, Smith RJ (2021) Monosynaptic Retrograde Tracing From Prelimbic Neuron Subpopulations Projecting to Either Nucleus Accumbens Core or Rostromedial Tegmental Nucleus. *Front. Neural Circuits* 15:639733
- Cruz AM, Spencer HF, Kim TH, Jhou TC, Smith RJ (2020) Prelimbic cortical projections to rostromedial tegmental nucleus play a suppressive role in cue-induced reinstatement of cocaine seeking. *Neuropsychopharmacol. Off. Publ. Am. Coll. Neuropsychopharmacol.*
- Do-Monte FH, Manzano-Nieves G, Quiñones-Laracuenta K, Ramos-Medina L, Quirk GJ (2015) Revisiting the role of infralimbic cortex in fear extinction with optogenetics. *J. Neurosci. Off. J. Soc. Neurosci.* 35:3607–3615
- Economo MN, Clack NG, Lavis LD, Gerfen CR, Svoboda K, Myers EW, Chandrashekar J (2016) A platform for brain-wide imaging and reconstruction of individual neurons. *eLife* 5:e10566
- Elmer GI, Palacorolla H, Mayo CL, Brown PL, Jhou TC, Brady D, Shepard PD (2019) The rostromedial tegmental nucleus modulates the development of stress-induced helpless behavior. *Behav. Brain Res.* 359:950–957

721 Floresco SB (2013) Prefrontal dopamine and behavioral flexibility: shifting from an “inverted-U”
722 toward a family of functions. *Front. Neurosci.* 7:62

723 Floresco SB, Magyar O (2006) Mesocortical dopamine modulation of executive functions: beyond
724 working memory. *Psychopharmacology (Berl.)* 188:567–585

725 Gabbott PLA, Warner TA, Jays PRL, Salway P, Busby SJ (2005) Prefrontal cortex in the rat:
726 projections to subcortical autonomic, motor, and limbic centers. *J. Comp. Neurol.* 492:145–177

727 Gaspar P, Bloch B, Le Moine C (1995) D1 and D2 receptor gene expression in the rat frontal
728 cortex: cellular localization in different classes of efferent neurons. *Eur. J. Neurosci.* 7:1050–1063

729 Glover EJ, McDougale MJ, Siegel GS, Jhou TC, Chandler LJ (2016) Role for the Rostromedial
730 Tegmental Nucleus in Signaling the Aversive Properties of Alcohol. *Alcohol. Clin. Exp. Res.*
731 40:1651–1661

732 Gourley SL, Taylor JR (2016) Going and stopping: Dichotomies in behavioral control by the
733 prefrontal cortex. *Nat. Neurosci.* 19:656–664

734 Hong S, Jhou TC, Smith M, Saleem KS, Hikosaka O (2011) Negative reward signals from the
735 lateral habenula to dopamine neurons are mediated by rostromedial tegmental nucleus in
736 primates. *J. Neurosci.* 31:11457–11471

737 Huang S, Zhang Z, Gambeta E, Xu SC, Thomas C, Godfrey N, Chen L, M'Dahoma S, Borgland
738 SL, Zamponi GW (2020) Dopamine Inputs from the Ventral Tegmental Area into the Medial
739 Prefrontal Cortex Modulate Neuropathic Pain-Associated Behaviors in Mice. *Cell Rep.* 33:108393

740 Jhou TC, Fields HL, Baxter MG, Saper CB, Holland PC (2009)(a) The rostromedial tegmental
741 nucleus (RMTg), a GABAergic afferent to midbrain dopamine neurons, encodes aversive stimuli
742 and inhibits motor responses. *Neuron* 61:786–800

743 Jhou TC, Geisler S, Marinelli M, Degarmo BA, Zahm DS (2009)(b) The mesopontine rostromedial
744 tegmental nucleus: A structure targeted by the lateral habenula that projects to the ventral
745 tegmental area of Tsai and substantia nigra compacta. *J. Comp. Neurol.* 513:566–596

746 Kataoka N, Shima Y, Nakajima K, Nakamura K (2020) A central master driver of psychosocial
747 stress responses in the rat. *Science* 367:1105–1112

748 Kaufling J, Veinante P, Pawlowski SA, Freund-Mercier M-J, Barrot M (2009) Afferents to the
749 GABAergic tail of the ventral tegmental area in the rat. *J. Comp. Neurol.* 513:597–621

750 Kebschull JM, Garcia da Silva P, Reid AP, Peikon ID, Albeanu DF, Zador AM (2016) High-
751 Throughput Mapping of Single-Neuron Projections by Sequencing of Barcoded RNA. *Neuron*
752 91:975–987

753 Kupferschmidt DA, Cody PA, Lovinger DM, Davis MI (2015) Brain BLAQ: Post-hoc thick-section
754 histochemistry for localizing optogenetic constructs in neurons and their distal terminals. *Front.*
755 *Neuroanat.* 9:6

756 LaBerge D, Kasevich R (2013) The cognitive significance of resonating neurons in the cerebral
757 cortex. *Conscious. Cogn.* 22:1523–1550

758 Lecca S, Pelosi A, Tchenio A, Moutkine I, Lujan R, Hervé D, Mameli M (2016) Rescue of GABAB
759 and GIRK function in the lateral habenula by protein phosphatase 2A inhibition ameliorates
760 depression-like phenotypes in mice. *Nat. Med.* 22:254–261

761 Lee AT, Vogt D, Rubenstein JL, Sohal VS (2014) A Class of GABAergic Neurons in the Prefrontal
762 Cortex Sends Long-Range Projections to the Nucleus Accumbens and Elicits Acute Avoidance
763 Behavior. *J. Neurosci.* 34:11519–11525

764 Li H, Pullmann D, Cho JY, Eid M, Jhou TC (2019)(a) Generality and opponency of rostromedial
765 tegmental (RMTg) roles in valence processing. *eLife* 8

766 Li H, Vento PJ, Parrilla-Carrero J, Pullmann D, Chao YS, Eid M, Jhou TC (2019)(b) Three
767 Rostromedial Tegmental Afferents Drive Triply Dissociable Aspects of Punishment Learning and
768 Aversive Valence Encoding. *Neuron*

769 Loewke AC, Minerva AR, Nelson AB, Kreitzer AC, Gunaydin LA (2021) Frontostriatal Projections
770 Regulate Innate Avoidance Behavior. *J. Neurosci. Off. J. Soc. Neurosci.* 41:5487–5501

771 Matsumoto M, Hikosaka O (2007) Lateral habenula as a source of negative reward signals in
772 dopamine neurons. *Nature* 447:1111–1115

773 McCutcheon RA, Abi-Dargham A, Howes OD (2019) Schizophrenia, Dopamine and the Striatum:
774 From Biology to Symptoms. *Trends Neurosci.* 42:205–220

775 McGuier NS, Padula AE, Lopez MF, Woodward JJ, Mulholland PJ (2015) Withdrawal from chronic
776 intermittent alcohol exposure increases dendritic spine density in the lateral orbitofrontal cortex of
777 mice. *Alcohol Fayettev. N* 49:21–27

778 Mena-Segovia J, Bolam JP (2017) Rethinking the Pedunculopontine Nucleus: From Cellular
779 Organization to Function. *Neuron* 94:7–18

780 National Research Council (2011) Guide for the Care and Use of Laboratory Animals: Eighth
781 Edition 8th ed. Washington DC: National Academies Press.

782 Nieh EH, Matthews GA, Allsop SA, Presbrey KN, Leppla CA, Wichmann R, Neve R, Wildes CP,
783 Tye KM (2015) Decoding neural circuits that control compulsive sucrose seeking. *Cell* 160:528–
784 541

785 Omelchenko N, Bell R, Sesack SR (2009) Lateral habenula projections to dopamine and GABA
786 neurons in the rat ventral tegmental area. *Eur. J. Neurosci.* 30:1239–1250

787 Paxinos G, Watson C (2007) The Rat Brain in Stereotaxic Coordinates, Sixth Edition: Hard Cover
788 Edition 6 edition. Amsterdam ; Boston: Academic Press.

789 Perez de la Mora M, Gallegos-Cari A, Crespo-Ramirez M, Marcellino D, Hansson AC, Fuxe K
790 (2012) Distribution of dopamine D(2)-like receptors in the rat amygdala and their role in the
791 modulation of unconditioned fear and anxiety. *Neuroscience* 201:252–266

792 Peters J, Kalivas PW, Quirk GJ (2009) Extinction circuits for fear and addiction overlap in
793 prefrontal cortex. *Learn. Mem. Cold Spring Harb. N* 16:279–288

794 Rock C, Zurita H, Lebby S, Wilson CJ, Apicella AJ (2018) Cortical Circuits of Callosal GABAergic
795 Neurons. *Cereb. Cortex N. Y. N* 1991 28:1154–1167

796 Rockland KS (2013) Collateral branching of long-distance cortical projections in monkey. *J.*
797 *Comp. Neurol.* 521:4112–4123

798 Rozeske RR, Evans AK, Frank MG, Watkins LR, Lowry CA, Maier SF (2011) Uncontrollable, But
799 Not Controllable, Stress Desensitizes 5-HT_{1A} Receptors in the Dorsal Raphe Nucleus. *J.*
800 *Neurosci.* 31:14107–14115

801 Santana N, Mengod G, Artigas F (2009) Quantitative analysis of the expression of dopamine D1
802 and D2 receptors in pyramidal and GABAergic neurons of the rat prefrontal cortex. *Cereb. Cortex*
803 *N. Y. N* 1991 19:849–860

804 Schultz W (1986) Responses of midbrain dopamine neurons to behavioral trigger stimuli in the
805 monkey. *J. Neurophysiol.* 56:1439–1461

806 Schultz W, Dayan P, Montague PR (1997) A neural substrate of prediction and reward. *Science*
807 275:1593–1599

808 Sharpe MJ, Marchant NJ, Whitaker LR, Richie CT, Zhang YJ, Campbell EJ, Koivula PP,
809 Necarsulmer JC, Mejias-Aponte C, Morales M, Pickel J, Smith JC, Niv Y, Shaham Y, Harvey BK,
810 Schoenbaum G (2017) Lateral Hypothalamic GABAergic Neurons Encode Reward Predictions
811 that Are Relayed to the Ventral Tegmental Area to Regulate Learning. *Curr. Biol. CB* 27:2089-
812 2100.e5

813 Simmler LD, Ozawa T (2019) Neural circuits in goal-directed and habitual behavior: Implications
814 for circuit dysfunction in obsessive-compulsive disorder. *Neurochem. Int.* 129:104464

815 St Laurent R, Martinez Damonte V, Tsuda AC, Kauer JA (2020) Periaqueductal Gray and
816 Rostromedial Tegmental Inhibitory Afferents to VTA Have Distinct Synaptic Plasticity and Opiate
817 Sensitivity. *Neuron* 106:624-636.e4

818 Stamatakis AM, Stuber GD (2012) Activation of lateral habenula inputs to the ventral midbrain
819 promotes behavioral avoidance. *Nat. Neurosci.* Available at:
820 <http://www.ncbi.nlm.nih.gov/pubmed/22729176> [Accessed June 30, 2012].

821 Sun Y, Cao J, Xu C, Liu X, Wang Z, Zhao H (2020) Rostromedial tegmental nucleus-substantia
822 nigra pars compacta circuit mediates aversive and despair behavior in mice. *Exp. Neurol.*
823 333:113433

824 Tønnesen J, Katona G, Rózsa B, Nägerl UV (2014) Spine neck plasticity regulates
825 compartmentalization of synapses. *Nat. Neurosci.* 17:678–685

826 Vander Weele CM, Siciliano CA, Matthews GA, Namburi P, Izadmehr EM, Espinel IC, Nieh EH,
827 Schut EHS, Padilla-Coreano N, Burgos-Robles A, Chang C-J, Kimchi EY, Beyeler A, Wichmann
828 R, Wildes CP, Tye KM (2018) Dopamine enhances signal-to-noise ratio in cortical-brainstem
829 encoding of aversive stimuli. *Nature* 563:397–401

830 Vincent SL, Khan Y, Benes FM (1995) Cellular colocalization of dopamine D1 and D2 receptors
831 in rat medial prefrontal cortex. *Synap. N. Y. N* 1991 19:112–120

- 832 Volkow ND, Fowler JS, Wang G-J (2004) The addicted human brain viewed in the light of imaging
833 studies: brain circuits and treatment strategies. *Neuropharmacology* 47:3–13
- 834 Warden MR, Selimbeyoglu A, Mirzabekov JJ, Lo M, Thompson KR, Kim S-Y, Adhikari A, Tye KM,
835 Frank LM, Deisseroth K (2012) A prefrontal cortex-brainstem neuronal projection that controls
836 response to behavioural challenge. *Nature* 492:428–432
- 837 Wayman WN, Woodward JJ (2018) Exposure to the Abused Inhalant Toluene Alters Medial
838 Prefrontal Cortex Physiology. *Neuropsychopharmacol. Off. Publ. Am. Coll.*
839 *Neuropsychopharmacol.* 43:912–924

840

FIGURE LEGENDS

Figure 1. Anatomical distribution of cortical inputs to the RMTg. (A) Representative images demonstrating dense ipsilateral cortical labeling in brain areas injected with CtB into the RMTg. (B) Representative high magnification image showing that inputs to the RMTg arise primarily from layer V of the mPFC. (C) The percent of CtB+ neurons relative to all layer V NeuN+ neurons is relatively consistent across ACC, PL, and IL subregions of the mPFC whereas the density of RMTg-projecting DP mPFC neurons increases substantially at more caudal levels. (D) Contralateral cortical afferents are substantially less dense than ipsilateral inputs as exemplified by a comparison of RMTg-projecting PL mPFC neurons in both hemispheres. (E) The density of layer V OFC neurons projecting to the RMTg is similar to that observed in the mPFC with LO inputs diminishing at more caudal levels. (F) CtB labeling is consistently observed in the AIC, albeit to a lesser degree than that observed in mPFC and OFC. (G) Map of tracer injection sites for all animals included in quantification. (H) Representative injection site. Abbreviations: ACC = anterior cingulate cortex; AID = agranular insular cortex, dorsal; AIV = agranular insular cortex, ventral; DI = dysgranular insular cortex; DLO = dorsolateral orbitofrontal cortex; DP = dorsopeduncular cortex; GI = granular insular cortex; IL = infralimbic cortex; LO = lateral orbitofrontal cortex; MO = medial orbitofrontal cortex; PL = prelimbic cortex; VO = ventral orbitofrontal cortex.

Figure 2. RMTg-projecting dmPFC neurons express CaMKII α . Representative mPFC images co-labeled for (A₁₋₃) the glutamatergic marker CaMKII α (red) and CtB (blue) and the (B₁₋₃) GABAergic marker GAD67 (green) and CtB (blue) from rat injected with CtB into the RMTg. (C) Quantification of co-labeling reveals that RMTg-projecting neurons are CaMKII α +. Scale bar = 25 μ m.

Figure 3. RMTg-projecting dmPFC neurons collateralize throughout the brain. (A) An intersectional dual-virus approach was used to fill RMTg-projecting dmPFC neurons with yellow fluorescent protein (YFP). (B) Representative images showing RMTg-projecting dmPFC neurons filled with YFP following amplification using standard immunohistochemistry. (C) Quantification of the average percent-stained area within regions of interest placed within the respective brain regions. (D) Representative YFP staining in the amygdala shows relatively sparse collateralization of RMTg-projecting dmPFC neurons in the basolateral nucleus. (E) Representative YFP staining in the striatum shows dense collateralization in the dorsomedial but not dorsolateral striatum. Scale bar = 100 μ m

Figure 4. Optogenetic stimulation of RMTg-projecting dmPFC terminals drives avoidance. (A) Representative ChR2 expression in dmPFC. (B) Rats spend significantly less time relative to chance in the light-paired side of a two-compartment chamber during initial testing (test 1) and when the light-paired compartment is reversed (test 2) when light delivery results in stimulation of dmPFC terminals in the RMTg. (C) A similar degree of avoidance of the light-paired chamber is observed upon stimulation of lateral habenula inputs to the RMTg. (D) Unlike stimulation of dmPFC terminals in the RMTg, stimulation of dmPFC terminals in the VTA fails to produce either preference for or avoidance of the light-paired compartment. (E) Direct comparison of circuit manipulations reveals significant avoidance when stimulating inputs to the RMTg relative to the VTA. Light-paired side indicated by blue bar in representative maps above each dataset. * $p \leq 0.01$, scale bar = 1000 μ m.

Figure 5. cFos induction in RMTg-projecting dmPFC neurons following exposure to aversive stimuli. (A) Experimental procedures. (B) Rats that had tone paired with foot shock delivery displayed significantly more freezing behavior in response to tone presentation than rats that were exposed to the same number of tone-shock presentations but in an unpaired manner.

(C) Significantly greater cFos expression was observed in RMTg-projecting dmPFC neurons (CtB+) following exposure to either a series of foot shocks or a tone predictive of foot shock relative to a neutral tone or the testing context alone. **(D)** Representative images of CtB and cFos labeling in the dmPFC of a context-exposed rat and a rat exposed to foot shock. CtB⁺/cFos⁻ neurons are indicated with a yellow arrowhead; CtB⁻/cFos⁺ neurons are indicated with a black arrow; CtB⁺/cFos⁺ neurons are indicated by a red asterisk. Scale bar = 200 μ m.

Figure 6. Decreased excitability in RMTg-projecting dmPFC neurons following exposure to aversive stimuli. **(A)** Experimental preparation. **(B)** Significantly fewer spikes were observed in shock-exposed rats relative to controls in current clamp recordings of retrobead-labeled dmPFC neurons. **(C)** Representative traces from a control and shock-exposed rat. Decreased spiking was associated with a significant increase in **(D)** rheobase, **(G)** membrane capacitance, and **(H)** peak action potential amplitude as well as a significant decrease in **(F)** membrane resistance and **(I)** action potential half-width. No significant difference was observed in **(E)** action potential threshold or **(J)** after-hyperpolarization. **(K)** Representative retrobead injection site in the RMTg. * $p \leq 0.05$, scale bar = 1000 μ m.

Figure 7. Exposure to aversive stimuli increases spine neck diameter in RMTg-projecting dmPFC neurons. **(A)** An intersectional dual-virus approach was used to fill RMTg-projecting dmPFC neurons with yellow fluorescent protein (YFP). **(B)** Representative YFP-filled primary apical dendrites in the dmPFC and accompanying Imaris renderings for context- and shock-exposed rats. **(C)** Spine density did not differ between groups regardless of subclass. However, shock-exposed rats exhibited significantly greater spine neck diameter **(D)** and shorter spine length **(E)** across all subtypes (main effect of shock) relative to rats exposed to the neutral testing context. * $p \leq 0.05$; scale bar = 5 μ m

Figure 8. D1 & D2 receptor gene expression in RMTg-projecting dmPFC neurons. (A-E)

Representative labeling in dmPFC neurons shows dense retrobead labeling in green **(A)** accompanied by D1 **(B)** and D2 **(C)** mRNA transcript dots indicated in red and yellow, respectively. The merged image **(D)** shows heterogeneous overlap of the retrograde beads and mRNA transcripts dots in cells identified using nuclear labeling with DAPI (blue). **(E)** Imaris rendering of individual cells from A-D showing colocalization of the beads/dots in the 3D rendered soma. Foot shock had no effect on the total number of cells or number of bead positive cells analyzed **(F)**. The prevalence of D1 and D2 mRNA transcript containing cells was similar between the treatment groups **(G)**. When collapsed across groups **(H)**, it is apparent that the majority of RMTg-projecting dmPFC neurons are D1+ with a large proportion of D1+ neurons also expressing D2 receptor mRNA. Foot shock exposure had no significant effect on the magnitude of either D1 **(I)** or D2 **(J)** mRNA expression. Scale bar = 20 μ m

Figure 9. Shock-induced enhancement of cFos expression occurs in D1⁺ and D1⁻ RMTg-

projecting dmPFC neurons. (A) Shock exposure significantly increased cFos mRNA expression in both bead⁺ and bead⁻ dmPFC neurons. **(B)** cFos expression was similarly increased in both D1⁺ and D1⁻ neurons in shock-exposed rats relative to controls. **(C)** cFos mRNA expression was significantly greater in shock-exposed rats compared to controls, but but there was no significant difference between D1⁺ and D1⁻ cell populations. Representative RNAScope labeling from a context-exposed control **(D)** and shock-exposed **(E)** rat. Retrobead labeling is depicted in green **(D-E₁)**, cFos mRNA in red **(D-E₂)**, D1 receptor mRNA in yellow **(D-E₃)**, merged image **(D-E₄)** and Imaris rendering of colocalized beads and dots the cell soma **(D-E₅)**. Nuclear labeling by DAPI depicted in blue. Scale bar = 20 μ m; *p $\leq$ 0.05

Measure	Statistical test	Effect	Result
Dendrite diameter	Unpaired t-test		t(10)=0.6430, p=0.3960
Dendrite volume	Unpaired t-test		t(10)=0.3731, p=0.1613
Overall spine density	Unpaired t-test		t(10)=0.2997, p=0.4101
Spine density by class	Two-way ANOVA	Class	F(3,40)=53.37, p<0.0001
		Treatment	F(1,40)=0.0750, p=0.7856
		Class x Treatment	F(3,40)=1.338, p=0.2756
Spine length by class	Two-way ANOVA	Class	F(3,36)=224.7, p<0.0001
		Treatment	F(1,36)=1.427, p=0.2401
		Class x Treatment	F(3,36)=1.573, p=0.2129
Spine diameter by class	Two-way ANOVA	Class	F(3,36)=23.77, p<0.0001
		Treatment	F(1,36)=2.492, p=0.1232
		Class x Treatment	F(3,36)=0.8049, p=0.4994
Terminal point diameter by class	Two-way ANOVA	Class	F(3,36)=13.77, p<0.0001
		Treatment	F(1,36)=0.0579, p=0.8112
		Class x Treatment	F(3,36)=0.0967, p=0.9614
Spine volume by class	Two-way ANOVA	Class	F(3,36)=15.89, p<0.0001
		Treatment	F(1,36)=1.832, p=0.1843
		Class x Treatment	F(3,36)=0.7693, p=0.5188
Spine neck volume by class	Two-way ANOVA	Class	F(3,36)=18.86, p<0.0001
		Treatment	F(1,36)=2.404, p=0.1298
		Class x Treatment	F(3,36)=0.9704, p=0.4173
Terminal point volume by class	Two-way ANOVA	Class	F(3,36)=5.712, p=0.0026
		Treatment	F(1,36)=0.9216, p=0.3435
		Class x Treatment	F(3,36)=0.6379, p=0.5955
Neck length by class	Two-way ANOVA	Class	F(3,29)=180.6, p<0.0001
		Treatment	F(1,29)=4.190, p=0.0498
		Class x Treatment	F(3,29)=3.179, p=0.0388
Neck diameter by class	Two-way ANOVA	Class	F(3,33)=27.33, p<0.0001
		Treatment	F(1,33)=9.847, p=0.0036
		Class x Treatment	F(3,33)=1.689, p=0.1884

Table S1. Complete analysis of dendritic spine density & morphology in RMTg-projecting dmPFC neurons following exposure to aversive stimuli. Bolded results indicate statistical significance of $p \leq 0.05$.