Prefrontal cortical and nucleus accumbens contributions to discriminative conditioned suppression of reward-seeking

Patrick T. Piantadosi, Dylan C.M. Yeates, and Stan B. Floresco

Department of Psychology and Djavad Mowafaghian Centre for Brain Health, University of British Columbia, Vancouver, British Columbia V6T 1Z4, Canada

Fear can potently inhibit ongoing behavior, including reward-seeking, yet the neural circuits that underlie such suppression remain to be clarified. Prior studies have demonstrated that distinct subregions of the rodent medial prefrontal cortex (mPFC) differentially affect fear behavior, whereby fear expression is promoted by the more dorsal prelimbic cortex (PL) and inhibited by the more ventral infralimbic cortex (IL). These mPFC regions project to subregions of the nucleus accumbens, the core (NAcC) and shell (NAcS), that differentially contribute to reward-seeking as well as affective processes that may be relevant to fear expression. Here, we investigated how these mPFC and NAc subregions contribute to discriminative fear conditioning, assessed by conditioned suppression of reward-seeking. Bilateral inactivation of the NAcS or PL reduced the expression of conditioned suppression to a shock-associated CS+, whereas NAcC inactivation reduced reward-seeking without affecting suppression. IL inactivation caused a general reduction in conditioned suppression following discriminative conditioning, but not when using a single-stimulus design. Pharmacological disconnection of the PL → NAcS pathway revealed that this projection mediates conditioned suppression. These data add to a growing literature implicating discrete cortico- striatal pathways in the suppression of reward-seeking in response to aversive stimuli. Dysfunction within related structures may contribute to aberrant patterns of behavior in psychiatric illnesses including substance use disorders.

Corresponding author: floresco@psych.ubc.ca

© 2020 Piantadosi et al. This article is distributed exclusively by Cold Spring Harbor Laboratory Press for the first 12 months after the full-issue publication date (see http://learnmem.cshlp.org/site/misc/terms.xhtml). After 12 months, it is available under a Creative Commons License (Attribution-NonCommercial 4.0 International), as described at http://creativecommons.org/licenses/by-nc/4.0/.
role for these regions in fear-induced suppression. We then conducted a pharmacological disconnection experiment aimed at isolating the PL → NAcS pathway responsible for the appropriate expression of conditioned suppression.

Results

NAcS inactivation

The experimental timeline of these studies is presented in Figure 1A, and the location of infusion for each region can be found in Figure 1B. Inactivation of NAcS (n = 13) markedly disrupted the expression of discriminative conditioned suppression, as compared to controls (n = 14) (Fig. 2A). Analysis of these data produced a significant CS Type × Treatment interaction (F(1,25) = 5.02, P < 0.035) that was driven by less suppression during presentation of the CS+ for animals in the NAcS inactivation group (F(1,25) = 4.24, P = 0.05). In contrast, lever-pressing during the CS− did not differ across treatments (F(1,25) = 0.20, P > 0.66). Inspection of Figure 2A (right) would suggest that NAcS inactivation had a greater effect on fear expression more prominently during the latter presentations of the CS+ compared to the first presentation. However, analyses failed to reveal a significant three-way interaction or any other two-way interactions (all F-values <1.3, all P-values >0.25). NAcS inactivation did not alter the total number of lever-presses made during the session (t(25) = −1.18, P > 0.24), nor the rate of pressing during the initial 5 min baseline period (t(25) = 0.15, P > 0.88; Table 1), indicating that NAcS inactivation did not induce a general disinhibition of reward-seeking. Similarly, there was no change in overall locomotion during the expression test session (t(25) = −1.21, P > 0.23; Table 1). Thus, the NAcS was required for response-suppression during the presentation of an aversive CS+, but not for discriminating between distinct stimuli or general reward-seeking motivation under these conditions.

NAcC inactivation

In contrast to the impact of NAcS inactivation, inactivation of the adjacent NAcC had no impact on fear expression (Fig. 2B). No main effect of Treatment was observed (F(1,19) = 0.05, P > 0.84), indicating that NAcC inactivated rats (n = 9) expressed levels of fear comparable to control rats (n = 12). Discrimination between the CS− and CS+ was intact, regardless of treatment condition (main effect of CS Type (F(1,19) = 102.36, P < 0.001; CS Type × Treatment interaction (F(1,19) = 0.54, P > 0.47)). Additionally, there was no significant three-way interaction (F(1,30) = 0.80, P > 0.50). However, NAcC inactivation reduced the total number of lever-presses (t(19) = 2.23, P = 0.04), and decreased the rate of lever-pressing during the first five min of the session relative to control animals, although this latter effect only approached statistical significance (t(19) = 1.74, P = 0.09; Table 1). NAcC inactivation also decreased locomotion (t(19) = 2.80, P < 0.02; Table 1). These data suggest that NAcC does not play an integral role in modulating fear responses based on cues predicting safety or an aversive consequence, but does promote behavioral activation, enhancing response vigor during reward-seeking.

PL cortex inactivation

Like the NAcS, neural activity in the PL cortex was found to be necessary for the appropriate expression of conditioned suppression (Fig. 3A). A significant main effect of Treatment (F(1,23) = 13.09, P < 0.001) suggested that PL inactivation (n = 13) diminished conditioned suppression, as compared to control animals (n = 12). This was confirmed by a significant CS Type × Treatment interaction (F(1,23) = 11.68, P < 0.005), with PL inactivation resulting in less fear to the CS+ as compared to control rats (F(1,23) = 19.77, P < 0.001). There was no change in the rate of lever-pressing prior to the first CS presentation (t(23) = 0.32, P > 0.75), suggesting that the disinhibition of pressing during the CS+ in PL-inactivated animals was not a result of general behavioral activation (Table 1). Although the total number of lever-presses made during the session was not affected by PL inactivation (t(23) = 1.28, P = 0.20), locomotor activity tended to be reduced, although this effect only approached significance (t(23) = −1.76, P > 0.09; Table 1). Thus, PL

---

**FIGURE 1.** Discriminative fear task diagram and histology. (A) Discriminative fear task procedure. (B) Histology figure diagraming location of infusions for animals used in the bilateral inactivation experiment. Red triangles represent NAcC placements, yellow pentagons indicate NAcC placements, blue circles represent PL placements and gray circles represent IL placements. Each dot represents the most ventral extent of the infusion, as observed in Nissl stained sections and the numbers beside each plate represent mm from bregma.
activity was necessary for the appropriate expression of fear toward a discriminative CS+, with inactivation markedly reducing the suppression of reward-seeking typically observed during its presentation.

we performed an additional experiment to clarify whether the suppression-reducing impact of IL inactivation was specific to discriminative conditioning. When a separate group of rats (n = 8; location of infusion displayed in Fig. 4A) were subjected to a fear

TABLE 1. Mean (±SEM) values for total locomotion, rate of lever-pressing, and total lever-presses during the discriminative fear expression test session

| Infusion timeline | Cannula placement | Treatment | Locomotion (photobeam breaks) | Lever-press rate (presses/min) | Total lever-presses |
|------------------|------------------|-----------|-----------------------------|-------------------------------|---------------------|
| Bilateral inactivation | NAcS | SAL | 1762 (±218) | 21.0 (±3.6) | 760.6 (±135.8) |
| | B/M | 2285 (±338) | 20.3 (±2.9) | 1028.7 (±189.2) |
| | NAcC | SAL | 1709 (±199) | 21.9 (±2.9) | 751.1 (±193.3) |
| | B/M | 989 (±131)* | 15.6 (±1.8)# | 483.0 (±60.9)* |
| | PL | SAL | 1557 (±188) | 22.2 (±3.8) | 1003.5 (±140.9) |
| | B/M | 1176 (±115)# | 23.8 (±3.0) | 1245.9 (±126.7) |
| | IL | SAL | 1560 (±165) | 21.5 (±2.8) | 1029.6 (±147.0) |
| | B/M | 1365 (±195) | 21.9 (±3.0) | 1173.2 (±127.9) |
| | Single-stimulus | IL | SAL | 2304 (±333) | 24.4 (±5.4) | 1450.9 (±242.2) |
| | B/M | 2412 (±518) | 25.4 (±4.6) | 2587.9 (±318.2)* |
| | PL-NAcS | Control | SAL | 1651 (±205) | 24.3 (±3.3) | 968.2 (±114.7) |
| | | Contra-Disc | B/M | 2155 (±272) | 23.6 (±4.0) | 1056.2 (±217.1) |
| | | Ipsi-Disc | B/M | 3384 (±357)* | 23.5 (±3.9) | 938.6 (±104.1) |
| | | Uni-Inact | B/M | 2455 (±371) | 19.8 (±2.9) | 947.5 (±144.5) |

* P < 0.05 versus SAL. # P = 0.09.

www.learnmem.org 431 Learning & Memory
conditioning protocol using a single CS, inactivation of the IL did not affect the expression of conditioned suppression compared to control animals \( (n=8; \text{Fig. 4B}) \). There was no main effect of Treatment \( (F_{1,14}=1.65, P>0.22) \), with both groups extinguishing at a comparable rate as indicated by a significant effect of CS Block \( (F_{5,70}=10.02, P<0.001) \), and no significant Treatment \times CS Block interaction \( (F_{15,70}=1.57, P>0.18) \). Although the rate of pressing during the first 5 min of the session was similar across Treatment \( (t_{14}=0.14, P>0.89) \), rats receiving IL inactivation made more lever presses throughout the session \( (t_{14}=2.84, P<0.013; \text{Table 1}) \). However, inactivation had no impact on overall locomotor activity \( (t_{14}=0.17, P>0.87; \text{Table 1}) \). Taken together, these results suggest that neural activity in the IL cortex may play a more selective role in the suppression of reward-seeking in situations requiring the disambiguation of conditioned aversive and neutral stimuli.

### PL-NAcS Disconnection

The qualitatively similar effects of PL mPFC and NAcS inactivation on discriminative fear expression led us to probe whether a functional PL→NAcS circuit promotes conditioned suppression. Separate groups of animals received contralateral disconnection \( (n=9) \), ipsilateral disconnection \( (n=9) \), saline \( (n=10) \), or unilateral inactivation \( (n=10; 5 \text{ PL and 5 NAcS} \text{)} \text{Fig. 5A}) \). Disconnection of the PL cortex from the NAcS prior to the expression test diminished conditioned suppression \( (\text{Fig. 5B)} \; \text{main effect of Treatment} \; (F_{1,34}=3.66, P<0.022, \text{CS Type} \times \text{Treatment interaction} \; (F_{13,34}=6.46, P<0.001)) \). There was no three-way interaction \( (F_{9,103}=1.31, P>0.24) \). Follow up simple-main effects analyses on the two-way interaction revealed that this effect was due to a difference between the treatment conditions on CS+ trials \( (F_{4,33}=8.42, P<0.001) \), but not CS− trials \( (F_{4,33}=0.66, P>0.58) \). Post-hoc comparisons with Tukey’s test indicated that suppression during the CS+ was similar between saline-treated animals and the unilateral inactivation group \( (P>0.80) \). In contrast, the contralateral \( (P<0.001) \) and ipsilateral disconnection \( (P<0.05) \) groups displayed less conditioned suppression during CS+ presentations compared to controls, although the ipsilateral disconnection and unilateral inactivation groups did not differ \( (P>0.15) \). Conversely, contralateral disconnection resulted in significantly less suppression during the CS+ than was observed in rats receiving unilateral inactivation \( (P<0.0013) \), suggesting that contralateral disconnections had a greater effect compared to ipsilateral ones. Notably, the impairment in conditioned suppression during the CS+ induced by contralateral PL→NAcS disconnection appeared to be smaller in magnitude to that induced by bilateral inactivation of the PL \( (\text{Fig. 2A)} \) This observation was confirmed by a direct, exploratory statistical comparison of the conditioned suppression to the CS+ between the two groups \( (F_{11,20}=17.41, P<0.001) \).
None of the disconnection treatments affected the total number of lever presses made during the expression test relative to control animals \((F_{(3,34)}=0.12, P>0.94)\), nor the rate of lever-pressing during the initial portion of the test session \((F_{(3,34)}=0.30, P>0.82; \text{Table 1})\). However, locomotor activity did differ as a function of treatment \((F_{(3,34)}=7.02, P<0.001; \text{Table 1})\), driven by an increase in locomotor activity in the ipsilateral disconnection group, as compared to all other groups \((\text{all } P\text{-values}<0.025)\). Collectively, these findings confirm that serial communication between the PL and the NAc aids in suppressing reward-seeking in response to a conditioned aversive CS.

**Discussion**

These data demonstrate that separate subregions of the NAc and mPFC uniquely contribute to the expression of discriminative Pavlovian fear, as measured by conditioned suppression. Under control conditions, presentation of an aversive CS plus caused a marked suppression of ongoing reward-seeking, while presentation of a neutral CS minus did not alter behavior. Inactivation of either the NAc or PL diminished the expression of conditioned suppression while having no effect on responding during a neutral CS minus. Disconnection of these two structures reduced conditioned suppression during the expression test, consistent with the top-down regulation of reward-seeking by this discrete cortico-striatal pathway.

**NAc subregion-specific control of conditioned suppression**

Activity within the NAc was necessary for the reorganization of behavior during the presentation of an aversive stimulus, but did not affect overall levels of instrumental responding or other measures of activity. These data add to a growing literature implicating the NAc in the suppression of certain patterns of behavior. For example, the extinction of reinstated food seeking \((\text{Floresco et al. } 2008)\) and the extinction of alcohol or cocaine seeking \((\text{Peters et al. } 2008; \text{Millan et al. } 2010)\) are dependent upon NAcS integrity. Similarly, refining behavior through the learned cessation of instrumental responding during periods of reward unavailability or nonreinforcement is believed to be mediated by an inhibitory function of the NAc \((\text{Blaiss and Janak } 2009; \text{Ambroggi et al. } 2011; \text{Ghazizadeh et al. } 2012; \text{Floresco et al. } 2018)\). Populations of neurons that encode task-irrelevant stimuli and behaviors during reward-seeking are more numerous in the NAc, as compared to the NAcC \((\text{Ambroggi et al. } 2011)\), which may provide a neuronal mechanism for the NAcC-specific impact on response-inhibition. The ability of the NAc to suppress inappropriate actions may be mediated in part by projections from dopamine D1-receptor expressing medium spiny neurons to GABAergic neurons in the lateral hypothalamus, as this pathway has recently been shown to regulate the inhibition of alcohol-seeking acquired during extinction \((\text{Gibson et al. } 2018)\). Similarly, medial NAc D1-containing neurons projecting to the medial ventral tegmental area promote behavioral inhibition \((\text{Yang et al. } 2018)\). Finally, NAc activity is critical to withholding actions in the face of potential aversive consequences such as instrumental punishment \((\text{Piantadosi et al. } 2017; \text{Halladay et al. } 2020)\), predator odor exposure \((\text{Blomeley et al. } 2017)\), and inhibitory avoidance \((\text{Piantadosi et al. } 2018)\). These previous findings, in addition to the present data support the idea that the NAcS may refine action selection by suppressing inappropriate behaviors, including curtailing reward-seeking in response to potential threats \((\text{Floresco } 2015)\).

In contrast, inactivation of the NAcC did not alter conditioned suppression or affect discrimination between the CS plus and CS minus under the experimental conditions used here. Although the NAc has been suggested to control aspects of contextual Pavlovian fear learning \((\text{Haralambous and Westbrook, } 1999; \text{Wendler et al. } 2013)\), disrupting activity in this subregion does not generally affect the acquisition or expression of fear toward discrete cues \((\text{Jongen-Rêlo et al. } 2002; \text{Levita et al. } 2002; \text{McDannald and Galarce } 2011)\). Given that only cued fear was evaluated in the present experiments, the lack of a NAc inactivation effect on fear expression is perhaps not surprising. Intriguingly, recent evidence suggests that the NAcC may be recruited to aid in fear expression in situations where cue discrimination (and thus, the appropriate allocation of fear) is made more difficult by the inclusion of an ambiguous CS that is probabilistically associated with foot shock \((\text{Ray et al. } 2020)\). When viewed in light of the present data, this suggests that the NAcC may play a more prominent role in discriminative fear expression when threat is probabilistic, but not under more simple conditions when CSs are associated with shock in a deterministic manner.

Notably, manipulation of the NAcC was not without effect, as inactivation decreased both locomotion and reward-seeking \((\text{Table 1})\). Such effects are consistent with previous reports suggesting that this nucleus is involved in reward-related approach and other forms of behavioral activation \((\text{Ghods-Shariﬁ and Floresco } 2010; \text{Nicola } 2010; \text{Piantadosi et al. } 2017, 2018)\). Indeed, the NAcC is essential for the ability of an appetitive Pavlovian conditioned stimulus to invigorate behavior \((\text{Parkinson et al. } 2000; \text{Yun et al. } 2004; \text{Ambroggi et al. } 2011)\). Intact neural and dopaminergic activity within this nucleus is required for reward-predictive cues to promote efficient instrumental reward-seeking \((\text{Nicola } 2010; \text{Ambroggi et al. } 2011; \text{McGinty et al. } 2013)\). Yet, the present results show that NAcC activity is not essential for inhibition of reward-seeking by an aversive Pavlovian conditioned stimulus. Thus, the mechanisms through which the NAcC modulates the impact of Pavlovian cues on behavior appears to be biased toward response-promotion, rather than response-inhibition. More generally, these findings point to a double dissociation between these two NAc subregions in the organization of behavior.
subregions in negotiating motivational conflict during reward-seeking under threat, with the NAcS promoting the cessation of ongoing behaviors in response to aversive conditioned stimuli, and the NAcC promoting the vigor of reward-seeking.

**Prefrontal contribution to discriminative fear expression**

The finding that PL inactivation disrupted conditioned suppression is in keeping with numerous studies showing that this region of the mPFC is required for the expression of Pavlovian fear (Vidal-Gonzalez et al. 2006; Corcoran and Quirk 2007; Sierra-Mercado et al. 2011; Plantadosi and Florescu 2014; Sangha et al. 2014; Limpens et al. 2015). Models of PL function during the early stages of fear expression and extinction posit that activity within this subregion promotes defensive reactions such as freezing and conditioned suppression in part via interactions with the amygdala (Sierra-Mercado et al. 2011; Pendyam et al. 2013). In keeping with this, PL cortex has been shown to regulate response-inhibition induced by aversive conditioned stimuli during the seeking of cocaine (Chen et al. 2013; Limpens et al. 2015) or alcohol (Seif et al. 2013). Similarly, PL (and potentially IL) cortex appear to mediate the suppression of cocaine-seeking produced by periods of learned cocaine unavailability (Mihindou et al. 2013; Gutman et al. 2014). The present study supports these findings, demonstrating that the fear promoting aspect of the PL cortex is specific to a CS+ when animals are required to discriminate between aversive and neutral cues.

Our observation that IL cortex inactivation decreased conditioned suppression when animals had to discriminate between aversive and neutral cues is somewhat surprising. Using a single-stimulus conditioning procedure, Sierra-Mercado et al. (2011) reported that inactivation of IL prolongs conditioned freezing, an effect opposite to that of PL cortex inactivation. Conversely, stimulation of this region has been shown to decrease fear, enhancing extinction either within-session or across sessions (Milad et al. 2004; Vidal-Gonzalez et al. 2006; Bukela et al. 2015). However, limited experimental evidence suggests that the fear-induced suppression of reward-seeking, unlike conditioned freezing, is either decreased (as seen here) or unaffected by IL inactivation (Rødested et al. 2008; Sierra-Mercado et al. 2011; Jean-Richard-Dit-Bressel and McNally 2016). Indeed, when we conducted a single-stimulus fear conditioning paradigm (as is common in the literature), conditioned suppression was intact despite IL inactivation (Fig. 4B). These data suggest that the comparable function of PL and IL observed here may relate to the discriminative nature of our task. In support of this, Sangha et al. (2014) have shown that these prefrontal subregions are not functionally dissociable during performance of a similar Pavlovian discrimination task. Inactivation of either prefrontal subregion altered discriminative fear expression in the same manner, decreasing conditioned freezing during an expression test session, while leaving intact the ability of a neutral, safe cue to ameliorate fear (Sangha et al. 2014). Thus, IL cortex may promote fear during situations that produce conflict between representations evoked by stimuli associated with safety versus fear.

**A PL → NAcS pathway contributes to conditioned suppression**

Given that the PL cortex projects to the NAcS (Sesack et al. 1989; Brog et al. 1993; Vertes 2004), and that bilateral inactivation of either region induced a qualitatively similar effect on discriminative conditioned suppression, we evaluated whether these regions form a functional circuit that promotes conditioned suppression. We observed that contralateral or ipsilateral disconnection of PL→NAcS circuitry disinhibited reward-seeking in the presence of an aversive CS+, although contralateral disconnection appeared to induce a greater effect. Notably, unilateral inactivation of either region by itself did not significantly disrupt fear expression, indicating that the effects of the disconnection treatment were not attributable to suppression of activity in either the NAcS or PL cortex alone. This combination indicates that both ipsilateral and contralateral communication within this cortico-striatal circuit promotes conditioned suppression of reward-seeking. Previous studies have also reported that ipsilateral disconnection of the NAc from its inputs can alter behavior. For example, ipsilateral manipulation of prefrontal or amygdalar projections to the NAc alters risk/reward decision making, while similar ipsilateral manipulation of prefrontal or ventral subicular projections to the NAc affects the reinstatement of drug-seeking behavior (Bossert et al. 2012, 2016; Jenni et al. 2017; St-Onge et al. 2012; van Holstein et al. 2020). One explanation for the similarity between our ipsilateral and contralateral disconnection effects is that a proportion of PFC neurons project to the contralateral hemisphere, meaning that exclusively ipsilateral manipulations will affect some contralaterally projecting neurons. These contralateral and ipsilateral projections may contribute similarly to behavior, as was demonstrated for a ventral PFC-NAcS projection during the context-induced reinstatement of heroin-seeking (Bossert et al. 2012). Notably, in the present study, the magnitude of the PL→NAcS disconnection effect on conditioned suppression (Fig. 5B) appeared to be smaller than that induced by bilateral PL inactivation (Fig. 2A). This suggests that prefrontal control of conditioned fear expression may manifest through multiple output pathways that include the NAcS, as well as the amygdala (Likhitik et al. 2005; Sotres-Bayon and Quirk 2010; Knapska et al. 2012) and periaqueductal gray (Rozeske et al. 2018).

The finding that a PL→NAc pathway regulates discriminative conditioned suppression adds a functional correlate to previous work demonstrating that some NAc-projecting mPFC neurons encode the behavioral relevance of an aversive CS+ and a neutral CS− (McGinty and Grace 2008). A similar microcircuit between the mPFC and lateral NAcS promotes the suppression of reward-seeking following instrumental punishment (Kim et al. 2017). In that study, a subset of mPFC neurons projecting to the lateral NAcS decreased their activity immediately prior to presses on a punished lever, suggesting that activity within this pathway promotes inhibitory control. Other excitatory inputs to the NAcS, including the amygdala, ventral hippocampus, and paraventricular thalamus have been shown to regulate behavioral inhibition in a variety of settings (Bagot et al. 2015; Millan et al. 2015, 2017; Schumacher et al. 2016, 2018; Yeates et al. 2019; Capuzzo and Florescu 2020; Lafferty et al. 2020), indicating that NAcS may be a hub controlling behavioral output in the face of conflicting motivations. Yet, how glutamatergic projections to the NAcS accomplish such regulation remains unclear. It is possible that PL input differentially affects distinct neuronal subpopulations within the NAcS that may be responsible for action promotion or inhibition, such as medium spiny neurons that express either dopamine D1 or D2 receptor subtypes, as has been shown in the NAc and other striatal regions (Ferguson et al. 2011; Lobo et al. 2011; Kravitz et al. 2012). For example, a projection from ventromedial PFC to the medial NAcS inhibits alcohol-seeking following instrumental punishment experience, potentially by affecting plasticity at D1-receptor expressing medium spiny neurons (Halladay et al. 2020). However, recent work has highlighted that unproductive behaviors governed by the NAcS are not distinctly regulated by particular medium spiny neuron subtypes (Lafferty et al. 2020). Clearly, further study is necessary to identify the circuit mechanisms by which excitatory afferents to the NAcS affect behavior, including the conditioned suppression of reward-seeking.
Relevance to fear circuitry and psychiatric illness

The studies reported here utilized a discriminative fear conditioning procedure that is similar to those used in translational settings, where CS− presentations serve as a baseline index of fear, and CS+ presentations induce fear. Using such designs, a relatively conserved fear circuit encompassing the amygdala, prefrontal cortex, and ventral striatum has been identified in the human brain (for reviews, see Delgado et al. 2008b; Peters et al. 2009; Milad and Quirk 2012; Adolphs 2013). Within the PFC, the dorsal anterior cingulate cortex (dACC; Brodmann’s area 32) and ventromedial PFC (vmPFC; Brodmann’s area 25) have been suggested to be functionally and anatomically homologous to the rodent PL and IL cortex, respectively (Milad and Quirk 2012; Heilbronner et al. 2016). Activity in the dACC occurs in response to CS+, presentations, and this activity (as well as the overall thickness of the region) correlates positively with physiological measures of fear in humans (Milad et al. 2007a). On the other hand, vmPFC activity appears to track extinction learning in humans, as this region displays patterns of activity consistent with deactivation during conditioning, but activation during extinction (Phelps et al. 2004; Milad et al. 2007b). Here, we provide tentative support for the dACC-PL homology suggested by these previous studies, as they apply to the expression of conditioned fear. However, our results seem to suggest that IL cortex performs a similar function, promoting conditioned suppression, in a manner inconsistent with human vmPFC activity. This may again stem from the nature of the defensive reaction measured, as freezing (in rats) and skin conductance or verbal scoring (in humans) do not produce a state of motivational conflict similar to that induced by the conditioned suppression of reward-seeking. Although conditioned suppression paradigms exist in humans (Greville et al. 2013; Allcoat et al. 2015), to date, the relevant functional imaging studies have not been performed to evaluate this hypothesis.

In addition to prefrontal homology, discriminative aversive conditioning produces activity in the human ventral striatum (Jensen et al. 2003; Delgado et al. 2008a,b; 2009; Klucken et al. 2009; Pohlack et al. 2012). This activity is generally differential, with activity increasing in response to a CS+ to a greater degree than a CS−, a pattern which develops over the course of the conditioning session (Klucken et al. 2009). Activity within this nucleus has been shown to track fear into motivated action, as learning to avoid an aversive CS+ also recruits the NAc (Delgado et al. 2009). In the present study, NAcS activity was necessary for the appropriate expression of discriminative conditioned suppression. Thus, it is possible that the NAc activity observed in human imaging studies of fear learning may reflect preferential activation of the NAcS. Interestingly, diffusion tractography was used to differentiate the NAcS and NAcC in the human brain, with results indicating that the putative NAcS responds in anticipation of thermal pain, while NAcC responds particularly to the offset of a painful stimulus (Baliki et al. 2013). Whether this anticipatory activity relates to behavior is currently unknown, but may partially explain the anticipatory activity observed in NAc prior to presentation of a conditioned aversive stimulus (Jensen et al. 2003).

A number of neuropsychiatric disorders are characterized by dysfunction within cortical and striatal nodes that contribute to abnormal decision-making processes. For example, prefrontal hypofunction appears to be related to inhibitory control deficits in substance abuse (for review, see Goldstein and Volkow 2011). In cocaine users, deficits in inhibitory control are known to correlate with reduced dACC activity, the same region suggested to promote fear expression previously (Kaufman et al. 2003; Hester and Garavan 2004; Li et al. 2008; Goldstein et al. 2009). In rats, hypofunction of the functionally homologous PL cortex recapitulates key aspects of addictive behavior, including drug-seeking under threat of punishment (Chen et al. 2013; Limpens et al. 2015). Such a deficit may be related to the loss of a response-inhibitory function within the PL or dACC, as a function of addiction progression. Additionally, meta-analytic studies have consistently shown that patients with anxiety disorders express more fear to a CS− than do control individuals (Lissek et al. 2005; Duits et al. 2015). This deficit may be related to aberrant function of prefrontal circuitry, as trait anxiety is associated with diminished coupling between the amygdala and the vmPFC and a heightened coupling between the amygdala and the dorsomedial PFC, patterns that were opposite that observed in healthy comparison subjects (Kim et al. 2011). Specifically, vmPFC activity is negatively modulated by similarity to a CS+, while dorsomedial PFC activity is positively modulated by the CS+ similarity. This effect has recently been reported to be disturbed in individuals with PTSD, suggesting that imbalanced prefrontal discrimination mechanisms may contribute to anxiety (Kaczkuri et al. 2017). In the present study, the fear expressed toward a CS− was normal regardless of treatment. Thus, other regions, such as the BLA, which has been shown to encode the valence of discriminative stimuli in rats, nonhuman primates, and humans (Schiller et al. 2008; Genu-Gabai et al. 2013; McHugh et al. 2013; Sangha et al. 2013), may be causally related to fear generalization.

Experimental limitations

The experiments described here are subject to several important limitations. First, only male rats were tested. Many fear and anxiety-related disorders disproportionately affect women (Kessler et al. 1994, 1995; Breslau et al. 1999; McLean et al. 2011), and rodent studies have demonstrated sex differences in fear expression and its underlying neural circuitry (Rey et al. 2014; Gruene et al. 2015a,b; Fenton et al. 2016). Given that female rats (as compared to males) display distinct active and passive defenserreactions (Gruene et al. 2015a), future studies should compare the relevance of these defensive reactions to situations where reward-seeking and fear overlap, as in conditioned suppression paradigms.

Second, there was a degree of drift in conditioned fear expression across experimental cohorts that complicates direct cross-region comparisons. This was particularly notable in the IL control groups (Figs. 3B, 4B), where conditioned suppression was lower than other SAL control groups. This difference may relate to the damage caused to somewhat distinct regions by cannula implantation, based on the differences in stereotaxic coordinates required to target individual regions. For example, IL implants were conducted without an angle, which likely resulted in damage to the overlying PL cortex. As shown here and elsewhere, PL cortex is necessary for appropriate fear expression, and thus animals with cannula implanted into the IL may display submaximal fear as a result. A similar rationale may explain the differences in fear expressed by control rats in the NAcS (medial cannula placement) versus NAcC (more lateral cannula placement) experiments. Conditioned suppression was somewhat higher in the NAcS SAL group (Fig. 2A, open circles), as compared to the NAcC SAL (Fig. 2B, open circles), or PL SAL (Fig. 3A, open circles) conditions. However, it is less clear how the damage induced by medial NAc cannula placement (mostly affecting posterior PFC and lateral septum) may predispose rats to be more fearful. Critically, our statistical analyses focused primarily on single brain regions, to ensure that differences in fear were assessed in relation to control groups that had comparable cannulation damage (that is, comparing Treatment within IL, rather than across brain regions).
Conclusion
Here we demonstrate that distinct subnuclei of the rat NAc and mPFC contribute to aspects of reward-seeking and its conditioned suppression. Both the NACs and PL were required for the appropriate expression of conditioned suppression, while NACC activity enhanced response vigor during reward-seeking. IL activity tended to bias rats away from reward-seeking during threat in a manner that was specific to contexts requiring cue discrimination. Finally, we show that a circuit between the PL and NACC is necessary to instantiate the appropriate inhibition of reward-seeking during aversive stimulus presentations. These findings extend our knowledge of cortico-striatal circuits mediating flexible reward-seeking, identifying prefrontal input to the medial NAC as a critical component of such a network.

Materials and Methods

Animals
All procedures were approved by the Animal Care Committee at the University of British Columbia, in accordance with the Canadian Council on Animal Care guidelines. Separate groups of male Long Evans rats (Charles River) arrived weighing 250–300 g. Rats were initially housed in groups (4–5 rats/cage) with ad libitum access to food and water. After 5–10 d of colony acclimatization, rats were stereotaxically implanted with stainless-steel guide cannula, as described below. Upon recovery, rats were singly housed and food-restricted to ~90% of their free-feeding weight. Rats were allowed to gain weight following this initial period of restriction, such that they were maintained on a delayed growth curve.

Apparatus
Behavior was assessed using eight standard Med Associates operant chambers, enclosed in sound attenuating chambers (30.5 x 24 x 21 cm; Med Associates), as previously described (Piantadosi et al. 2017). Each chamber was capable of delivering sucrose reinforcement (45 mg pellet; BioServ), and contained a house light and two 100 mA cue lights. An auditory speaker allowed for the delivery of discriminative auditory stimuli via a programmable generator (ANL-926, Med Associates). Locomotor activity was measured by four infrared photobeams located just above the grid floor, which was wired to a shock source and solid-state grid scanner for foot shock delivery.

Stereotaxic surgery and disconnection rationale
Due to changes in institutional policies regarding anesthesia, rats were anesthetized either with a combination of ketamine/xylazine (100/10 mg/mL at 100/10 mg/kg, i.p.) or a half dose of ketamine/xylazine (same mg/mL, i.p) followed by maintenance using Isoflurane anesthetic (2.5–3% Isoflurane concentration) throughout surgery.

For the bilateral inactivation experiments, twenty-three gauge bilateral stainless-steel guide cannula were implanted into the NACs, NACC, PL, or IL, according to the following stereotaxic coordinates (in mm):

- **NACd**—from bregma, **AP:** +1.3, **ML:** ±1.0, from dura, **DV:** −6.3
- **NACC**—from bregma, **AP:** +1.6, **ML:** ±1.8, from dura, **DV:** −6.3
- **PL**—from bregma: **AP:** +3.2, **ML:** ±0.7; from dura: **DV:** −2.8
- **IL**—from bregma: **AP:** +2.8; **ML:** ±0.7; from dura: **DV:** −4.1

Guide cannula were beveled at the tip to minimize damage when implanted, which in turn would be expected to curtail backflow of infusate to more dorsal regions. A subsequent series of experiments used a pharmacological disconnection approach to probe whether a functional pathway from the PL to the NACs may control the expression of conditioned suppression. Briefly, this approach entailed perturbing neural activity in one region (for example, the PL) within one hemisphere, which prevents transmission of task-relevant information to another region of interest (for example, the NACs). This is combined with an inactivation of an efferent target in the contralateral hemisphere (the NACs). As a result, direct communication between two brain regions within a neural circuit is disrupted in both hemispheres of the brain following a contralateral disconnection. In a separate group, neural activity can be disrupted within each region in the ipsilateral hemisphere (ipsilateral disconnection), leaving an intact circuit in the opposite hemisphere. Finally, unilateral inactivation of each region individually can be performed to assess whether the disconnection effect was due to the partial loss of a functional pathway, or whether the effect is mediated by a single node within this putative circuit.

For the disconnection experiments, single 23 gauge stainless steel guide cannula were implanted aimed at the PL and NACs in the contralateral or ipsilateral hemispheres, or unilaterally in the PL or NACs, according to the stereotaxic coordinates listed above. The particular hemisphere selected for each placement was counterbalanced across experimental conditions, such that equivalent numbers of rats received cannula in each combination of hemispheres.

For all surgical procedures, four stainless-steel skull screws were inaudited with dental acrylic to secure cannula in place. Stainless-steel obturators flush with the end of the guide cannula were inserted after surgery. Rats were given 5–10 d to recover from surgery before beginning behavioral training.

Lever training
The day before their initial operant training session, all rats were provided with ~30 sucrose pellets in their home cage, to reduce neophobia to the reinforcer. Training was conducted at a consistent time each day. Rats were initially trained to press a lever on the left side of the chamber on a fixed ratio 1 (FR1) schedule of reinforcement to a criterion of 40 total presses during the 30 min session (Fig. 1A). After reaching criterion, rats were trained over three consecutive days on increasing variable interval (VI) schedules, whereby reward was provided after approximately 15 (VI15), 30 (VI30), or 60 (VI60) sec of pressing (one session at each schedule, per day). Rats were then trained on the VI60 schedule for 10–13 d, after which rats received a fear conditioning session. A VI60 schedule engenders a high rate of lever-pressing in rats while maintaining a consistent reward rate, allowing for the accurate assessment of conditioned suppression as a proxy for fear (Kamin et al. 1963; McAllister 1997; Quirk et al. 2000; Piantadosi andFloresco 2014). The house light was illuminated during all sessions, including the test of fear expression.

Discriminative fear conditioning

**Conditioning session**
Rats underwent discriminative fear conditioning in an identical fashion as we have reported previously (Piantadosi andFloresco 2014). During this conditioning session, the reward lever was not inserted into the chamber and no food could be obtained. Thus, even though rats were placed in the same individual chambers throughout training, conditioning, and testing, CS-shock pairings likely were perceived to have occurred in a different context compared to the one where they lever pressed for food and would later be tested for in fear expression. This contextual shift is notable, as previous studies examining the effects of PL/IL inactivation on conditioned suppression have allowed animals to lever-press for food during the fear conditioning session, potentially complicating the dissociation between contextual and cued fear as assessed during the test session (Sierra-Mercado et al. 2011).

Rats were given eight presentations each of a 30 sec neutral conditioned stimulus (CS−; 1 kHz, 80 dB tone + cue light illumination, no house light) and a 30 sec aversive conditioned stimulus (CS+: 9 kHz, 80 dB tone + flashing house light coterminating with a 0.5 mA foot shock delivered over 0.5 sec), separated by an average
interstimulus interval of 180 sec (min: 100 sec, max: 240 sec). CS− and CS+ presentations occurred in a pseudorandom sequence over the course of the conditioning session, with the exception that the last two stimulus presentations were always the CS+ paired with shock. During CS presentations, the house light was turned off to maximize the salience of the visual component of each CS. The day after this conditioning session, animals were given a baseline VI60 session (no foot shocks or conditioned stimuli).

**Expression test session**

During the fear expression test, a 5 min baseline VI60 period preceded the presentation of four CS− presentations, followed by four presentations of the CS+ (30 sec each, no foot shock; 5 min interstimulus interval). The suppression of lever-pressing during each CS presentation served as an index of fear, as rats suppress seeking behavior in the presence of an aversive CS+ (Kamin et al. 1963; Quirk et al. 2000; Sierra-Mercado et al. 2011; Piantadosi and Floresco 2014). Suppression was calculated using the formula \( \frac{(A - B)}{(A + B)} \), where \( A \) was the number of lever-presses made in the 30 sec epoch prior to CS presentation, and \( B \) was the number of lever-presses made during the 30 sec CS presentation. A value of 1 indicates complete suppression, while values at 0 or below indicate no suppression or facilitation, respectively. Rarely, rats did not press during the pretone and tone period; a suppression value of 0 was applied to all such instances, as has been done previously (Quirk et al. 2000). An a priori inclusion criteria of greater than 200 presses made during the test session was established, as levels of pressing under this threshold can produce unreliable suppression ratios. Across all experimental cohorts, data from \( n = 3 \) rats were eliminated as a result of this criterion.

**Single-stimulus fear conditioning: pretest IL inactivation**

We conducted an additional experiment to ascertain whether inactivation of the IL affected the expression of conditioned suppression under conditions where a single shock-associated stimulus was used during conditioning, as opposed to the discriminative nature of the design used in our other experiments. Animals were implanted with cannula into the IL cortex and given an identical lever training protocol as described above. During the conditioning session, animals received eight presentations of a single, 30 sec CS+ (identical to the CS+ used in the discriminative protocol), similar to conditioning procedures typically used to study IL function (e.g., Akirav et al. 2006; Sierra-Mercado et al. 2011). Forty-eight hours later, rats were given a test session. Again, this session started with rats lever pressing for food delivered on a VI60 schedule. Five minutes into the session, rats received the first of 12 presentations of the 30 sec CS+ (no foot shock), each separated by a 3 min interstimulus interval.

**Microinfusion**

All animals underwent mock infusions 10 min prior to the final VI60 session before discriminative conditioning, as described previously (Piantadosi and Floresco 2014). On the infusion day, stainless-steel injectors extending 0.8 mm beyond the guide cannula were lowered into the region of interest. For the bilateral inactivation experiments, rats received infusions of 0.9% saline (SAL, 0.3 µL/side), or a solution containing the GABA<sub>A</sub>-receptor agonist baclofen and the GABA<sub>A</sub>-receptor agonist muscimol (B/M; 75 ng/µL of each drug at a volume of 0.3 µL/side). Infusions were conducted over 45 sec, with injectors left in place for an additional 60 sec to allow for diffusion. The surgical and microinfusion procedures used here are identical or similar to published reports used to dissociate between these and other closely apposed brain regions on a wide variety of behavioral measures (Floresco et al. 2008, 2018; Stopper and Floresco 2011; Dalton et al. 2014, 2016; Piantadosi et al. 2017, 2018; van Holstein and Floresco 2020). Electrophysiological and immunohistochemical studies estimate the functional spread of GABA agonist-induced neural inactivations to be ~1 mm (Martin and Ghez 1999; Allen et al. 2008; Hamel et al. 2017). Implantation coordinates in the present study were chosen in part to minimize potential spread into neighboring regions.

For the disconnection experiment, control rats received unilateral infusions of 0.9% SAL into the PL and contralateral NAcS (0.3 µL/side). Disconnection animals received infusion of B/M (75 ng/µL of each drug at a volume of 0.3 µL/side) into the contralateral PL and NAcS (contralateral disconnection), or ipsilateral PL and NAcS (ipsilateral disconnection). A separate group underwent unilateral inactivations of the PL or NAcS (same infusion parameters). Infusion timing was identical to the procedures described previously.

**Histology**

All rats were euthanized with CO<sub>2</sub>, brains were removed and fixed in a 4% phosphate buffered formalin solution. Brains were sectioned at 50 µm, following which tissue was mounted and Nissl stained using Cresyl Violet. Placements were examined under a light microscope.

**Data analysis**

For the bilateral inactivation experiments, the suppression ratio during each CS presentation during the expression test was analyzed using between-within-subjects three-way ANOVAs with Treatment group (SAL vs. B/M) as the between-subjects variable, and CS Type (CS− vs. CS+) and CS Number (1–4) as the within-subjects variables. Separate ANOVAs were conducted on data from animals infused within each brain region (NAcS, NAcc, PL, and IL). For the single-stimulus IL inactivation experiment, suppression ratios for each of the 12 CS+ presentations were calculated and binned into six separate bins of two CS+ presentations. These data were analyzed via a two-way ANOVA, with Treatment group (SAL vs. B/M) as the between-subjects variable, and CS Block (1–6) as the within-subjects variable. Follow-up simple main effects analyses were conducted using one-way ANOVAs or t-tests, where appropriate. Locomotion (photobeam breaks/session) during the conditioning session or expression test were analyzed using separate independent samples t-tests. The rate of lever-pressing (presses/min) in the first 5 min of the expression test session and the total number of lever-presses made during the session were analyzed in an identical fashion.

Analysis of the disconnection experiment was identical, with the exception that the between-subjects Treatment factor was made up of four levels: saline, contralateral disconnection, ipsilateral disconnection, unilateral inactivation. There was no significant difference between the mean suppression ratio during the CS− and CS+ for animals in the unilateral PL inactivation group (CS−: 0.07 ± 0.06 SEM, CS+: 0.82 ± 0.10 SEM) versus the unilateral NAcS inactivation group (CS−: 0.14 ± 0.06 SEM, CS+: 0.85 ± 0.08 SEM) (\( F_{1,16} = 0.39, P > 0.55 \)); therefore these groups were combined into a single unilateral inactivation group for all analyses. All other analyses were conducted as described previously for the bilateral inactivation experiments.

**Acknowledgments**

This work was supported by a Discovery Grant (RGPIN-2018-04295) from the Natural Sciences and Engineering Research Council of Canada to S.B.F. and a University of British Columbia Doctoral Fellowship to P.T.P. We thank Marie T. Tse, Katie Pezarro, Matthew Wilkins, and Magdalene Schluter for their assistance with behavioral testing.

**References**

Adolphs R. 2013. The biology of fear. *Curr Biol* 23: R79–R93. doi:10.1016/j.cub.2012.11.055

Akirav I, Raizel H, Maroun M. 2006. Enhancement of conditioned fear extinction by infusion of the GABA(A) agonist muscimol into the rat prefrontal cortex and amygdala. *Eur J Neurosci* 25: 758–764. doi:10.1111/j.1460-9568.2006.04603.x

www.learnmem.org 437 Learning & Memory
Piantadosi PT, Floresco SB. 2014. Prefrontal cortical GABA transmission.

Piantadosi PT, Yeates DCM, Wilkins M, Floresco SB. 2017. Contributions of the anterior cingulate cortex and nucleus accumbens core to mediating extinction learning in the rat: an anterograde tract-tracing study with Phascolus vulgaris leucoagglutinin.

Phelps EA, Delgado MR, Nearing KI, Ledoux JE. 2004. Extinction learning in the ventral striatum during aversive contextual conditioning in humans.

Peters J, LaLumiere RT, Kalivas PW. 2008. Infralimbic prefrontal cortex is responsible for inhibiting cocaine seeking in extinguished rats.

Rodriguez-Romaguera J, Monte FHMD, Quirk GJ. 2012. Deep brain stimulation of the infralimbic and prelimbic cortices.

Sangha S, Robinson PD, Greba Q, Davies DA, Howland JG. 2014. Alterations in reward, fear and safety cue discrimination after inactivation of the rat prelimbic and infralimbic cortices. Neuropsychopharmacology.

Schiller D, Levy I, Niv Y, LeDoux JE, Phelps EA. 2008. From fear to safety and back: reversal of fear in the human brain.

Schumacher A, Vlassov E, Ito R. 2016. The ventral hippocampus, but not the dorsal hippocampus is critical for learned approach avoidance decision making. Hippocampus.

Schumacher A, Villarreal FR, Ussling A, Riaz S, Lee ACH, Ito R. 2018. Ventral hippocampal CA1 and CA3 differently mediate learned approach–avoidance conflict processing. Curr Biol.

Schwienbacher I, Fendt M, Richardson R, Schnitzler HU. 2004. Temporary inactivation of the nucleus accumbens disrupts acquisition and expression of fear-terminated startle in rats. Brain Res.

Seif T, Chang S-J, Simms JA, Gibb SL, Dadgar J, Chen BT, Harvey BK, Ron D, Messing RO, Bonci A, et al. 2013. Cortical activation of accumbens hyperpolarization-activated NMDARs mediates aversion-resistant alcohol intake. Nat Neurosci.

Sesack SR, Deutch AY, Bunney BS. 1989. The organization of the efferent projections of the medial prefrontal cortex in the rat: An anterograde tract-tracing study with Phascolus vulgaris leucoagglutinin. J Comp Neurol.

Sierra-Mercado D, Padilla-Coreano N, Quirk GJ. 2011. Dissociable roles of prefrontal and infralimbic cortices, ventral hippocampus, and basolateral amygdala in the expression and extinction of conditioned fear.

Sotres-Bayon F, Quirk GJ. 2010. Prefrontal control of fear: more than just extinction. Curr Opin Neurobiol.

St Onge JR, Stopper CM, Zahm DS, Floresco SB. 2012. Separate prefrontal-subcortical circuits mediate different components of risk-based decision making. J Neurosci.

Stwahl MF, Deutch AY, Bunney BS. 1989. Flexible approach hypothesis: unidimensional hypothesis for the role of nucleus accumbens in decision making. Prog Neuropsychopharmacol Biol Psychiatry.

Steffes M, Steketee M, Dyer C, LeDoux JE, Phelps EA. 2008. Pavlovian approach behavior: further evidence for limbic cortical-ventral striatalal systems. Behav Neurol.

Stumpf MA, Aiba T, Siuciak JA, Everitt BJ. 2000. Disconnection of the anterior cingulate cortex and nucleus accumbens striatum in rats with lesions of the infralimbic cortex.

S therapies of the prelimbic and infralimbic cortices, ventral hippocampus, and basolateral amygdala in the expression and extinction of conditioned fear.

Svendsen CN, Deutch AY. 1983. Galanin-like immunoreactivity in the rat brain. Brain Res.

Tanaka S, Okada Y, Takeuchi H, Taya K, Tanaka T, Kuroda T, Ushio K, Kita H, Kikumoto T, Ogawa K. 2018. Classic fear conditioning induces expression of the cannabinoid receptor CB1 in dorsal striatum.

Teruel DCM, Floresco SB. 2018. Preclinical development and phase 1b clinical trial of TAD-130527: A novel CNS selective cannabinoid 1 receptor inverse agonist.

Vertes RP. 2004. Differential projections of the infralimbic and prelimbic cortex in the rat. Neurobiol Learn Mem.

Vidal-Gonzalez I, Vidal-Gonzalez B, Rauch SL, Quirk GJ. 2006. CA1 and CA3 of the hippocampus are critical for the recovery of extinguished fear.

Vidal-Gonzalez I, Vidal-Gonzalez B, Quirk GJ. 2008. The role of the prelimbic cortex in cue-guided risk/reward decision making. Proc Natl Acad Sci USA.

Wagner RA, McNaughton NB. 2005. The role of the hippocampus in memory consolidation: an update. Hippocampus.

Wang Z, Xu Z, Jiang J, Zhang S, Li P, Zhang H, Wang H, Zou X, Niu Q, Gao L. 2016. The role of the prelimbic cortex in the recovery of fear extinction.

Wright CI, Beijer AV, Groenewegen HJ. 1996. Basal amygdaloid complex involvement in Pavlovian fear conditioning and instrumental action. J Neurosci.

Wright CI, Beijer AV, Groenewegen HJ. 1998. Basal amygdaloid complex involvement in Pavlovian fear conditioning and instrumental action.

Xue T, Xu Z, Jiang J, Zhang S, Li P, Zhang H, Zou X, Niu Q, Gao L. 2018. The role of the prelimbic cortex in the recovery of fear extinction.

Yee Y, Wang Z, Xu Z, Jiang J, Zhang S, Li P, Zhang H, Zou X, Niu Q, Gao L. 2017. The role of the prelimbic cortex in the recovery of fear extinction.

Yang H, de Jong JW, Tak Y, Peck J, Bateup HS, Lammel S. 2018. Nucleus accumbens subnuclei regulate motivated behavior via direct inhibition and disinhibition of VTA dopamine neurons. Nat Neurosci.

Yusen DCM, Ussling A, Lee ACH, Ito R. 2018. The roles of the nucleus accumbens core and dorsomedial striatum in learning: performance and extinction of Pavlovian fear-conditioned responses and instrumental avoidance responses. Neurobiol Learn Mem.

Zahn DS, Brog JS. 1992. On the functional significance of subterritories in the “accumbens” part of the rat ventral striatum. Neuroscience.

Received May 4, 2020; accepted in revised form July 2, 2020.