Dopamine receptors mediate strategy abandoning via modulation of a specific prelimbic cortex–nucleus accumbens pathway in mice

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Edited by Anders Björklund, Lund University, Lund, Sweden, and approved April 19, 2018 (received for review September 29, 2017)

The ability to abandon old strategies and adopt new ones is essential for survival in a constantly changing environment. While previous studies suggest the importance of the prefrontal cortex and some subcortical areas in the generation of strategy-switching flexibility, the fine neural circuitry and receptor mechanisms involved are not fully understood. In this study, we showed that optogenetic excitation and inhibition of the prefrontal cortex–nucleus accumbens (NAc) pathway in the mouse respectively enhances and suppresses strategy-switching ability in a cross-modal spatial-egocentric task. This ability is dependent on an intact dopaminergic tone in the NAc, as local dopamine denervation impaired the performance of the animal in the switching of tasks. In addition, based on a brain-slice preparation obtained from Drd2-EGFP BAC transgenic mice, we demonstrated direct innervation of D2 receptor-expressing medium spiny neurons (D2-MSNs) in the NAc by prelimbic cortical neurons, which is under the regulation by presynaptic dopamine receptors. While presynaptic D1-type receptor activation enhances the glutamatergic transmission from the prefrontal cortex to D2-MSNs, D2-type receptor activation suppresses this synaptic connection. Furthermore, manipulation of this pathway by optogenetic activation or administration of a D1-type agonist or a D2-type antagonist could restore impaired task-switching flexibility in mice with local NAc dopamine depletions; this restoration is consistent with the effects of knocking down the expression of specific dopamine receptors in the pathway. Our results point to a critical role of a specific prelimbic cortex–NAc subpathway in mediating strategy abandoning, allowing the switching from one strategy to another in problem solving.

Significance

Strategy-switching flexibility is a critical executive function necessary for living in an ever-evolving environment, and this ability is often impaired in attentional deficit and hyperactivity disorder, schizophrenia, and early Parkinson’s disease. To date, the underlying brain circuitry and receptor mechanisms are not entirely clear. The results of the present study suggest the essential role of a specific projection from prefrontal cortex to nucleus accumbens (NAc) D2 medium spiny neurons as well as NAc dopamine and presynaptic dopamine receptors of this projection in controlling the strategy-switching flexibility. These findings promote a better understanding of circuitry and neurobiology of strategy-switching flexibility and could contribute to identifying novel therapeutic targets for patients suffering from strategy-switching inflexibility.


The authors declare no conflict of interest. This article is a PNAS Direct Submission.

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10. 1073/pnas.1717106115/-/DCSupplemental.
striatomesencephalic and striatopallidal pathways, some early studies suggested that D1-MSNs project to both the ventral tegmental area (VTA) in the midbrain and the ventral pallidum (VP) and that D2-MSNs preferentially project to the VP (19, 20). This finding has been refined by a recent study demonstrating that virtually 50% of VP neurons are under the innervation of D1-MSNs from the NAc core (21). Despite this, D2-MSNs, but not D1-MSNs, are suggested to be involved in the learning flexibility (22).

Dopaminergic innervation from the VTA represents a crucial control over behavioral flexibility executed by the prefrontal cortex and the medial striatum (23–25). The NAc also receives dopaminergic input from the VTA and, in addition to its known involvement in various learning and memory-associated behaviors (26–28), this mesolimbic dopamine pathway has been reported to regulate some forms of behavior flexibility (22, 29–31). However, there are contradictory conclusions regarding the types of dopamine receptors involved in the behavior flexibility (22, 31). Dopamine receptors are not expressed only by MSNs in the NAc. Physiological evidence indicates that they are also functionally present in the cortical terminals, including those from the prelimbic cortex (PrL), to regulate the glutamate release into the NAc (32–35). Thus, the presynaptic dopamine receptors, in addition to their postsynaptic counterparts, could act as important sites for mediating functions of dopamine. However, the available reports in the literature are still controversial regarding which dopamine receptor type, D1 or D2, mediates the presynaptic modulation of PrL–NAc glutamatergic transmission (32–35). Moreover, these studies did not take into account D1-MSNs and D2-MSNs being functionally segregated populations.

In this study, we elucidated the functional circuitry that links the roles of the prefrontal cortex and the NAc in behavioral flexibility by examining the involvement of PrL–NAc projection and NAc dopamine receptors in a spatial-egocentric strategy-switching task. A multidisciplinary approach was adopted. Our results unveiled a critical role of a PrL–NAc subpathway in mediating the strategy abandoning necessary for task-switching flexibility.

Results

PrL–NAc Pathway Contributes to Switching but Not Learning of a Strategy. To determine the participation of the prefrontal cortex–NAc projection in cognitive flexibility, we made use of optogenetic manipulation to functionally activate or inhibit this pathway and examined the effects on the learning and switching of strategy to get a food reward. Adeno-associated viruses (AAVs) encoding channelrhodopsin (ChR2)-EYFP or halorhodopsin (NpHR)-EYFP under the control of the CaMKII promoter were injected into the PrL, and optical cannula were implanted targeting the NAc (Fig. 1A, Upper). EYFP-positive terminals were found in the NAc, including the core region, suggesting the presence of a direct projection from the PrL to the NAc (Fig. 1A, Lower), which was consistent with results in patch-clamp recording experiments (see below). Indeed, the expression of functional ChR2 and NpHR in the PrL–NAc pathway responsive to light stimulation were verified in brain-slice preparations (Fig. S1). The experimental design is shown in the schematic in Fig. 1B, Upper. The behavioral paradigms involve a response-direction task (RDT) and a visual cue task (VCT) based on a four-arm cross-maze and the switching of these tasks (Fig. 1B, Lower). These spatial-egocentric tasks evaluate the ability of the animals to learn particular rules for rewards and their flexibility in switching to a new strategy by abandoning a previously learned one (22, 35), which was determined by the accumulated number of trials needed to reach the criterion as well as the number of different types of errors, including perseverative, regressive, and never-reinforced errors (for details see Materials and Methods).

In the mice with the expression of ChR2 in the PrL neurons, optogenetic activation of their terminals in the NAc did not affect the performance in task acquisition in both the RDT (total number of trials needed to reach the criterion: sham, 135.4 ± 23.7, n = 7; ChR2, 129.3 ± 24.1, n = 9; P > 0.05) and the VCT (total number of trials needed to reach the criterion: sham, 90.9 ± 7.8, n = 7; ChR2, 85.5 ± 11.0, n = 8; P > 0.05) tests compared with respective sham groups. The numbers of successful trials in consecutive sections (and the corresponding accuracy, in percentage) are shown in Fig. 2A and B, and the total

Fig. 1. Schematics of the optogenetic manipulation of the PrL–NAc pathway and behavioral tasks. (A, Upper) AAV-CaMKII-ChR2-EYFP and AAV-CaMKII-NpHR-EYFP were injected into the PrL of test animals, and an optical cannula was implanted directly above the NAc delivering 470 nm or 590 nm light. (Lower) Postmortem histological examination confirmed the expression of EYFP in soma of PrL neurons as well as in terminal-like structures in the NAc core region. aca, anterior commissure, anterior part. (B, Upper) The design and time-line of experiments. (Lower) In the RDT the animal is required to turn against its turn bias, and in the VCT the animal is required to turn to the arm associated with a visual cue. In task-switching test, the subject needs to abandon the previously learned rule and adopt another rule. The blue horizontal bar indicates a closed door in the corresponding paradigm.
number of trials needed to reach the criterion is shown in Fig. 2C. Optogenetic inhibition of the terminals through NpHR activation also had no influence on learning (total number of trials needed to reach the criterion: in RDT, NpHR, 141.0 ± 18.5, n = 8; P > 0.05; in VCT, NpHR, 87.4 ± 11.6, n = 7; P > 0.05, both compared with their respective sham groups) (Fig. 2 A–C). These data suggested that the PrL–NAc pathway is not required for learning these two strategies.

We next examined the ability of the animals in switching to learn the second strategy, after learning the first strategy in previous training sessions. A 1-h break was introduced between the two tasks. In the RDT-to-VCT switching paradigm, optogenetic activation of the PrL–NAc pathway significantly reduced the total number of trials needed to reach the criterion compared with the sham group (sham, 80.6 ± 4.3, n = 7; ChR2, 61.3 ± 5.8, n = 9; P < 0.05) (Fig. 2D). A similar result was found in VCT-to-RDT switching (sham, 145.7 ± 16.4, n = 7; ChR2, 102.0 ± 8.2, n = 8; P < 0.05) (Fig. 2D). On the other hand, when this pathway was inhibited optogenetically, the total number of trials needed to reach the criterion was significantly increased in both RDT-to-VCT switching (NpHR, 109.5 ± 7.3, n = 8; P < 0.01 compared with sham) and VCT-to-RDT switching (NpHR, 197.1 ± 15.5, n = 7; P < 0.05 compared with sham) (Fig. 2D). In addition to collecting data on the number of trials needed to reach the criterion, we classified error types to assess if impairments in shifting were due to deficits in the suppression of old modes of responding (perseverative errors), in exploring novel strategies (never-reinforced errors), or in the maintenance of novel strategies once perseveration has ceased (regressive errors). Analysis of these three subtypes of errors committed during strategy switching revealed that both perseverative errors and regressive errors (Fig. 2 E and F) were significantly altered and paralleled the changes in the total number of trials needed to reach the criterion. In contrast, there were no significant differences in the number of never-reinforced errors (Fig. 2G).

The significant difference in perseverative and regressive errors from optogenetically manipulated groups suggest the PrL–NAc pathway is necessary for successful disengagement from a previously relevant strategy.

**NAc Dopamine Depletion Impairs Strategy-Switching Flexibility.** To examine if accumbal dopamine also plays a role in the same strategy-switching task, stereotaxic injection of 6-hydroxydopamine hydrochloride (6-OHDA) into the NAc together with i.p. injection of desipramine were performed to selectively damage dopaminergic terminals. Tyrosine hydroxylase (TH) immunostaining in the slices confirmed that NAc TH-containing fibers were reduced 2 wk following 6-OHDA lesion (Fig. 3A). The normalized optical density of the NAc from the lesioned side was significantly lower than that of the unlesioned side (unlesioned, 2.29 ± 0.43; lesioned, 1.31 ± 0.13; n = 10 mice, P < 0.05) (Fig. 3B), which suggested that dopamine was partially but significantly depleted. In contrast, there was no change in the optical density of TH signals in the dorsolateral striatum (unlesioned, 2.04 ± 0.29; lesioned, 1.77 ± 0.24; n = 10 mice, P > 0.05) (Fig. 3B) and ventromedial striatum (unlesioned, 2.02 ± 0.26; lesioned, 1.82 ± 0.23; n = 10 mice, P > 0.05) (Fig. 3B). The partial depletion of dopamine was also confirmed by measurement of in vivo dopamine levels in the NAc by standard HPLC (unlesioned, 12.09 ± 1.21 ng/mg; lesioned, 6.09 ± 0.76 ng/mg; n = 3 mice, 

![Fig. 2. The PrL–NAc pathway contributes to task-switching flexibility. (A–D) Learning of the RDT (A) and VCT (B) tests by the mice, which could reach the criterion within 1 d. For the RDT, data were pooled from seven, nine, and eight animals for the sham, ChR2, and NpHR groups, respectively. For the VCT, seven, eight, and seven animals were used in the sham, ChR2, and NpHR groups, respectively. Compared with the sham group, ChR2 or NpHR activation of the PrL–NAc projection did not affect the total number of trials needed to reach the criterion in either the RDT (A and C) or the VCT (B and C). On the other hand, in both the RDT–VCT and VCT–RDT switching paradigms, activation of the PrL–NAc pathway in the ChR2 group significantly reduced the total number of trials needed to reach the criterion, and the number of trials needed to reach the criterion was significantly increased when this pathway was inhibited in the NpHR group (D). (E and F) The perseverative errors (E) and regressive errors (F) committed exhibited a pattern of change similar to the total number of trials needed to reach the criterion in both switching paradigms. (G) Never-reinforced errors (G) committed exhibited a pattern of change similar to the total number of trials needed to reach the criterion in both switching paradigms. (D) Never-reinforced errors (G) committed exhibited a pattern of change similar to the total number of trials needed to reach the criterion in both switching paradigms. (D) Never-reinforced errors (G) committed exhibited a pattern of change similar to the total number of trials needed to reach the criterion in both switching paradigms. (D) Never-reinforced errors (G) committed exhibited a pattern of change similar to the total number of trials needed to reach the criterion in both switching paradigms. (D) Never-reinforced errors (G) committed exhibited a pattern of change similar to the total number of trials needed to reach the criterion in both switching paradigms. (D) Never-reinforced errors (G) committed exhibited a pattern of change similar to the total number of trials needed to reach the criterion in both switching paradigms.
To verify that unilateral depletion of dopamine would not impose any asymmetry in motor ability, e.g., in executing turning movements, we quantified the locomotor behaviors of the animals in the open-field arena and also the turn bias in the T-maze. We found that no rotational behaviors were induced by the unilateral NAc dopamine depletion. There were no differences between sham and lesioned mice in general mobility, rotation ratio (Fig. S2 A and B), and turn bias in the T-maze (Fig. S2C).

Under the condition of local dopamine depletion in the NAc, the performances of the sham groups and lesioned groups were comparable in both the RDT and VCT. Fig. 3C shows their performance in successive sessions. In the RDT, the total number of trials needed to reach the criterion was 135.4 ± 23.7, n = 7, in the sham group and was 160.0 ± 17.7, n = 6, in the lesioned group (P > 0.05) (Fig. 3D). For the VCT, the total number of trials needed to reach the criterion was 90.9 ± 7.8, n = 7, in the sham group and was 92.6 ± 9.7, n = 7, in the lesioned group (P > 0.05) (Fig. 3D). These data suggested that acquisition of the tasks was not impaired by NAc dopamine depletion. However, in the mice switching from the RDT to the VCT, the performance of the lesioned group was degraded (Fig. 3C), and a significantly larger number of the trials was required to reach the criterion (sham, 80.6 ± 4.3, n = 7; lesion, 116.0 ± 7.4, n = 6; P < 0.01) (Fig. 3E). Likewise, lesioned mice undergoing VCT-to-RDT switching needed more trials to reach the criterion (sham, 145.7 ± 16.4, n = 7; lesion, 222.9 ± 16.5, n = 7; P < 0.01) (Fig. 3 C and E). These results indicate that NAc dopamine depletion could impair strategy-switching flexibility. Again, these impairments were contributed by perseverative errors and regressive errors. In both RDT–VCT and VCT–RDT switching, the numbers of these errors were significantly higher in the lesioned groups than in the sham groups (Fig. 3 F and G), while the numbers of never-reinforced errors remained unaffected (Fig. 3H).

**Impairment in Task-Switching Flexibility Can Be Rescued by Optogenetic Activation of the PrL–NAc Pathway.** The previous data imply that the PrL–NAc projection plays a significant role in mediating task-switching flexibility by facilitating the abandoning of an old strategy. This ability is also dependent on intact dopamine innervation within the NAc. We next asked whether impaired task-switching flexibility under NAc dopamine depletion could be rescued by enhancing the PrL–NAc pathway. Indeed, while optogenetic activation of the PrL–NAc pathway in the dopamine-depleted mice did not affect the learning of the RDT and VCT per se (Fig. S3A), this paradigm reduced the total number of trials needed to reach the criterion in both RDT-to-VCT switching experiments and VCT-to-RDT switching experiments to levels comparable to those of the respective sham groups (Fig. 3H).

(A) A typical section showing that TH immunoreactivity was reduced in the NAc core and shell regions 2 wk after focal injection of 6-OHDA into the NAc. CPu, caudate putamen. (B) Normalized data of optical density in the NAc (Left), dorsolateral CPu (Middle), and ventromedial CPu (Right) from 10 animals. *P < 0.05; paired t test. (C) Tracking of success trials in RDT and VCT learning as well as their switching in the sham, 6-OHDA lesioned, and lesioned+ChR2 groups. (D–H) Pooled data suggest that NAc dopamine depletion did not affect the ability of the animals to acquire the RDT and the VCT (D). However, in the RDT–VCT and VCT–RDT switching tests, NAc dopamine depletion resulted in a significantly higher number of trials before reaching the criterion (E), accompanied by increases in perseverative errors (F) and regressive errors (G), but not in never-reinforced errors (H). Optogenetic activation of ChR2 of the PrL–NAc pathway in the lesioned mice restored switching performance to that in the sham group. *P < 0.05; **P < 0.01; ***P < 0.001, unpaired t test. Data are presented as mean ± SEM.
PrL Innervates D2-MSNs in the Core Subregion of the NAc. Our data strongly suggest an interaction between the PrL–NAc pathway and the dopaminergic input from midbrain in the NAc in task-switching flexibility. An obvious question to address is whether dopamine released in the NAc directly modulates the PrL–NAc glutamatergic transmission. Among the MSNs in NAc, there is strong evidence showing that D2-MSNs in NAc, rather than D1-MSNs, are crucial in learning flexibility (22, 36), but the underlying mechanism has yet to be explored. Thus, as a first step in deciphering the neural circuitry, we focused on the role of the projection from the PrL to the core region of the NAc. To achieve this, we established an in vitro preparation that would allow us to study the modulation of the synaptic connections mediated by AMPA receptors (Fig. 4D). The current–voltage relationship of the eEPSC as shown in Fig. 4E indicated that Na⁺ is the main ion mediating the synaptic current. Among our samples, 81.0% of D2-MSNs and 88.0% of putative D1-MSNs in core subregion responded to PrL stimulation. D2-MSNs and putative D1-MSNs had similar eEPSC amplitude, latency, and paired-pulse ratio (Fig. S4B). The fact that the evoked synaptic event was monosynaptic rather than multisynaptic in nature was demonstrated by experiments based on optogenetic stimulation of PrL terminals in NAc expressing ChR2 (Fig. S1B). Under this condition, the application of blue light could directly evoke optical EPSCs (oEPSCs) in D2-MSNs. While perfusion of TTX abolished oEPSCs, the addition of the K-channel blocker 4-aminopyridine (4-AP) could largely restore them (Fig. 4F). Together, these results confirmed that an excitatory glutamatergic projection is sent from the PrL to innervate NAc D2-MSNs directly.

Presynaptic D1-Type Receptors and D2-Type Receptors Respectively Facilitate and Suppress PrL–NAc D2-MSN Transmission. To investigate the potential modulatory role of dopamine and its receptors on D2 MSN-associated PrL–NAc projections, in our brain-slice preparation, we examined the effects of specific D1-type and D2-type receptor agonists on membrane excitability as well as synaptic transmission. In these neurons, despite the expression of D2 receptors, the D2-type receptor agonist quinpirole (100 nM) did not induce a consistent change in membrane current. Thus, the membrane excitability was not acutely decreased.
modulated by the postsynaptic D2-type receptors, at least not with 100 nM quinpirole. To study the effects on the synaptic transmission, paired electrical stimulation with a 50-ms interval was delivered to the PrL. The amplitudes of the eEPSCs recorded from D2-MSNs were measured, and the paired-pulse ratios (PPRs) were also determined. The changes in PPR are inversely correlated with the changes in synaptic release probability of presynaptic terminals and thus can indicate if there is presynaptic modulation and in what direction (38, 39). We found that 20 nM of the D1-type agonist SKF8393 induced an increase in normalized eEPSC amplitude (without SKF8393: 1.00 ± 0.18; with SKF8393: 1.18 ± 0.19; n = 12, P < 0.05) (Fig. 5A and B) with a concomitant decrease in normalized PPR (without SKF8393: 1.00 ± 0.12; with SKF8393: 0.83 ± 0.08; n = 12, P < 0.05) (Fig. 5A and B). These changes suggest that presynaptic D1-type receptors are involved in enhancing D2 MSN-associated PrL–NAc transmission. The full D1-type receptor agonist SKF81297 (50 nM) also exerted a similar effect on normalized eEPSC amplitude (without SKF81297: 1.00 ± 0.26; with SKF81297: 1.12 ± 0.25; n = 8, P < 0.05) (Fig. 5C and D) and normalized PPR (without SKF81297: 1.00 ± 0.05; with SKF81297: 0.89 ± 0.05; n = 8, P < 0.01) (Fig. 5C and D). Intriguingly, the D2-type receptor agonist quinpirole exerted opposite effects on this pathway. As shown in Fig. 5E and F, quinpirole at 100 nM significantly decreased the normalized eEPSC amplitude (without quinpirole: 1.00 ± 0.25; with quinpirole: 0.63 ± 0.18; n = 8, P < 0.05), which was accompanied by an increase in normalized PPR (without quinpirole: 1.00 ± 0.08; with quinpirole: 1.49 ± 0.35; n = 8, P < 0.05). Consistently, in the eEPSCs, which have been shown to be monosynaptically activated (Fig. 4F), the D1-type and D2-type receptor agonists exerted effects similar to those they exerted in the eEPSCs (Fig. S5). These changes suggest that presynaptic D2-type receptors suppress D2 MSN-associated PrL–NAc transmission. Through reducing excitatory glutamatergic drive, this presynaptic D2-type receptor activation should lead to reduced excitability of D2-MSNs. Together, these results demonstrated that both D1-type receptors and D2-type receptors contribute to the acute modulation of excitatory neurotransmission from the PrL to NAc D2-MSNs. Furthermore, both D1-type and D2-type receptors can be expressed presynaptically, but they modulate the transmission in an opposite manner. Since activation of presynaptic D1-type receptors, but not D2-type receptors, can enhance neurotransmission from the PrL to NAc D2-MSNs, dopamine release from a VTA projection targeting these receptors may play a crucial role in facilitating task-switching ability.

**Opposite Roles of NAc D1-Type Receptors and D2-Type Receptors in Modulating Task-Switching Flexibility.** The role of the PrL projection onto NAc D2-MSNs in mediating task-switching ability is supported by pharmacological evidence. In view of the facilitating action of D1-type receptors and the suppressing action of D2-type receptors on this pathway, we speculated that activation of the D1-type receptors or blockade of the D2-type receptors is beneficial in alleviating the cognitive impairment induced by dopamine depletion. To test this hypothesis, we implanted an infusion cannula into the NAc core in dopamine-depleted mice (Materials and Methods) and infused the D1-type receptor agonist SKF81297 or the D2-type receptor antagonist sulpiride into the NAc 10 min before behavioral training. Compared with the lesion group, the pharmacological infusion did not affect the acquisition of the RDT and VCT tasks per se (Fig. S3A) but restored the task-switching performance in both RDT-to-VCT switching tasks (total number of trials needed to reach the criterion: sham, 80.6 ± 4.3, n = 7; lesion, 116.0 ± 7.4, n = 6; lesion+SKF81297, 81.6 ± 4.5, n = 5; lesion+sulpiride, 92.0 ± 8.0, n = 6;
**Discussion**

The involvement of the prefrontal cortex, including its subregions, and also the subcortical NAc in strategy-switching flexibility has been pursued and confirmed in separate studies (4, 5, 22, 36, 40, 41). It is known that the prefrontal cortex is connected with the NAc. Trials using pharmacological inactivation and disconnection lesions further suggested that the prefrontal cortex connections with the mediodorsal nuclei of the thalamus and NAc core mediate strategy switching by inhibiting perseveration (42). Here we demonstrated optogenetically that activating or inhibiting the terminals of neurons of the PrL in the NAc core could respectively enhance or suppress strategy-switching flexibility. Thus, our results extend the previous understanding of the role of prefrontal cortex–subcortical area connections, and in particular the contribution of the PrL–NAc pathway, in the expression of cognitive flexibility.

At the same time, accumulating evidence has long supported a critical control by dopamine of strategy-switching flexibility. In the prefrontal cortex or its subregion, the infusion of dopamine receptor agonists or antagonists differentially alters strategy-switching flexibility (23, 24). The dopaminergic innervation of the prefrontal cortex arises from the VTA that also sends projections to the NAc. In the present study, we observed that local depletion of dopamine in the NAc induced an impairment in switching different strategies (egocentric response-based vs. visual cue-based) to get a food reward, demonstrating the causal role of NAc dopamine on this type of cognitive flexibility. Thus, the interconnections among the VTA, prefrontal cortex, and the NAc together constitute an important circuitry that mediates expression of cognitive flexibility.

By taking advantage of the Drd2-EGFP BAC transgenic mice, we further dissected the circuitry between the prefrontal cortex and ventral striatum by confirming that both D2-MSNs and putative D1-MSNs in the NAc receive input from the PrL, as this notion has remained inconclusive from previous studies (53). While the role of innervation from the PrL to D1-MSNs is

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**Fig. 6.** D1-type receptor activation and D2-type receptor suppression ameliorate dopamine depletion-induced task-switching inflexibility. (A) In both the RDT→VCT and VCT→RDT switching paradigms, pretreatment with the D1-type agonist SKF81297 or the D2-type antagonist sulpiride restored the increased total number of trials needed to reach the criterion caused by dopamine lesion in the NAc. (B–D) The effects of the drugs are also reflected in the number of perseverative errors (B) and regressive errors (C) committed but not in the number of never-reinforced errors (D). In A–D, sham and lesion groups as shown in Fig. 3 E–H are included here for comparisons. *P < 0.05; **P < 0.01; ***P < 0.001, unpaired t test. Data are presented as mean ± SEM.
currently unknown, we provide evidence from electrophysiological and optogenetic approaches in brain slices that the glutamatergic connection to D2-MSNs is under tight regulation by presynaptic D1-type and D2-type receptors, which enhance and suppress the pathway, respectively. Although contributions by other dopamine receptors, including postsynaptic dopamine receptors on NAc MSNs (22), could not be excluded, observations from our in vivo and in vitro experiments together implicate key roles of these presynaptic dopamine receptors in the NAc in mediating task-switching flexibility. Thus, local dopamine depletion demonstrated not only the importance of an intact dopamine tone in this region but also that this deficit could be restored by the local administration of a D1-type agonist and a D2-type antagonist, as is consistent with their facilitating actions determined in in vitro experiments. On the other hand, although knockdown of D1 and D2 receptors via the AAV-shRNAs is not confined to the terminal regions of the PrL–NAc pathway, the fact that these manipulations resulted in suppression and facilitation of task-switching ability similar to that observed after administration of a D1-type agonist and a D2-type antagonist into the NAc is also consistent with the roles of the presynaptic dopamine receptors located in NAc.

It has been argued that cross-modal shift facilitated by the prelimbic–infralimbic cortex represents higher-order processing compared with intramodel shift because a new strategy is required to solve a task (9). We have shown in our study that the PrL–NAc projection, probably to the D2-MSNs, is critical in facilitating the abandoning of the old strategy but has minimal role in the learning of a new strategy. This notion is supported by the observation that the deficit in behavioral flexibility could be contributed by perseverative errors and regressive errors (23, 39), both implying a choice associated with the previously acquired but now incorrect or inefficient strategy. The key difference in the two types of errors is when the errors occur in the choice sequence. Perseverative errors are responses in which a previously reinforced strategy continues to be used despite a switch in the category rule and termination of positive feedback. Regressive errors are trials in which mice identify the newly reinforced response choice but then are unable to maintain this new response set and instead revert to the previously reinforced strategy. On the other hand, the never-reinforced errors, which can be interpreted as an index of how quickly animals are able to parse out an ineffective strategy and explore new response set, were unaffected in all manipulations in the present study. Our
results are in line with a previous finding by Florescu et al. (17) that functional inactivation of NAc core neurons by the infusion of GABA agonists did not impair initial learning of strategies but disrupted the ability to shift to a different strategy. However, in their study, regressive errors, but not perseverative errors, were impaired after NAc inactivation, which may reflect the consequence of a different interventional strategy. Combined together, these findings strongly indicate that the neuroplasticity process underlying the learning of a new task is a function distinct from that of task switching and probably is mediated by a different mechanism or circuit. Similar conclusions about the dissociation between the learning of a strategy and strategy switching have also been indicated in studies of the rodent medial prefrontal cortex (10, 11) and the primate lateral prefrontal cortex (54). However, we point out the lack of impact of optogenetic manipulation of prelimbic input on learning could also be due to the methodology in our study, i.e., unilateral rather than bilateral manipulations. On the other hand, it is possible that the activation of opsins during the learning phase could have a carryover effect on the strategy-switching phase, although we gave a 1-h break between the two phases. Other than methodology concerns, it is of particular interest to ascertain the specific functional neural circuitries that facilitate different components of strategy switching.

In conclusion, we demonstrated the critical involvement of PrL–NAc projections, in particular to D2-MSNs, and presynaptic dopamine receptors of this synapse in strategy-switching flexibility. Interestingly, perseverative types of errors are committed more by human patients with ADHD or schizophrenia who show deficits in cognitive flexibility (3, 55). Findings in this study may contribute to the development of novel therapeutic strategies for the impaired behavioral flexibility observed in ADHD, schizophrenia, and early Parkinson’s disease patients.

**Materials and Methods**

Animals, chemicals, stereotaxic surgery for in vivo studies, optogenetic stimulation, in vivo drug infusion, local dopamine denervation, immunohistochemical studies, brain-slice preparation, whole-cell recording, and the construction of AAV-shiRNAVs are described in SI Materials and Methods. All animals were handled in strict accordance with the guidelines by The Chinese University of Hong Kong on animal ethics. All animal procedures were approved by the Chinese University of Hong Kong Animal Experimentation Ethics Committee.

**Strategy Acquisition and Switching.** Strategy acquisition and switching in goal-directed behavior were tested with an RDT and a VCT and using a custom-built four-arm cross-maze, as described in other studies (22, 56). The four-arm cross-maze was made of a clear plastic wall with a gray floor and placed 90 cm above the floor of the room. Each arm was 25 cm long and 5 cm wide, and the center platform was 5 × 5 cm. The position of a mouse was detected by a video camera (C515; Logitech) suspended over the maze and was analyzed by the Any-Maze software (version 4.7; Stoelting Co.).

**Habitation and turn bias.** Animals were food-restricted to maintain about 85% of the original ad libitum weight from the beginning of behavioral task, which was started with habituation. The complete test consisted of several components including habituation, turn bias, RDT, VCT, and their switching. For animal treated with 6-OHDA injection, habituation training started 1 wk after surgery. Before each day’s habituation, mice were handled for 10 min. On the first habituation phase, three reward pellets were placed in each of the arms of the cross-maze (two in the food well at the arm end and one down the length of the arm). The mice were allowed to navigate freely and consume the food pellets for 15 min. If a mouse consumed all 12 pellets within 15 min, it was removed from the maze and placed in the holding cage. After the maze was re baited with eight additional pellets in the food well at the arm end, the mouse was placed back in the center of the maze and was allowed to consume all the pellets. On the second habituation phase, the procedure was almost the same, except that whenever the mouse traversed the entire length of an arm and consumed the two food pellets in the food well, it was picked up and placed at the entrance of a different arm, habituating the animal to repeat handling after consuming the food reward. On the third habituation phase, each arm was baited with only two pellets in the food well. A piece of black-and-white striped cardboard (10 cm wide × 50 cm long × 0.5 cm thick, as a visual cue) was placed outside and adjacent to one arm. After consuming all eight pellets, the mouse was placed in the holding cage, the visual cue was moved to a different arm, and the food was rebaited, until the mouse had consumed eight food pellets four times within 15 min. All mice finished the habituation training within 10 d (averaged 5 d, range 3–10 d).

Immediately after maze habituation, the turn bias was assessed in a T-maze (blocking one arm’s entry). No food was provided in this procedure. Mice were put in the stem arm and could turn 90° left or right when entering the center platform. After choosing an arm and reaching its end, the mouse was picked up, placed in the stem arm, and allowed to make the next choice. The direction being turned four or more times over seven trials was considered the mouse’s turn bias.

**RDT and VCT.** In the RDT, mice were required always to turn in the opposite direction of their turn bias to receive a food pellet, regardless of the location of the visual cue. In the VCT, mice were trained to enter the arm indicated by the visual cue. Twelve consecutive trials were set as one session. The starting arm for each trial and the position of the visual cue were determined pseudorandomly such that they were equally occurred in each experimental session. Each mouse was given an equal frequency for every consecutive set of 12 trials. Each task continued until the mouse reached the acquisition criterion of more than 10 correct choices in two consecutive sessions. Accuracy was calculated as the percentage of correct choices per session, and the total number of trials needed to reach the criterion was also recorded.

**RDT–VCT shifting.** After successful acquisition of RDT task, the mouse was placed back in the holding cage for 1 h and then was shifted to the VCT. Errors committed during the set shift determined the animal’s ability to abandon a previously learned strategy and acquire a new one. Perseverative errors were defined as those committed with the visual cue as a preceding cue. If a mouse entered the incorrect arm on three or more trials per block of four trials that required it to enter the arm indicated by the visual cue, which was always opposite to the turning direction required in the previous RDT. After the first time a mouse made fewer than three perseverative errors in a block, subsequent errors were no longer counted as perseverative errors because at this point the mouse started to choose an alternative strategy at least half of the time. Instead, the stem arm was randomly selected in the following trials, and the subsequent errors following the previous RDT rule were counted as regressive errors. The third type of error, never-reinforced errors, was scored when a mouse entered the incorrect arm during trials in which the visual cue was placed in the same arm as previous RDT.

**VCT–RDT shifting.** Animals for this set of experiment were initially trained on the VCT, followed by testing on the RDT after a 1-h rest in the holding cage. All other aspects were as described above, and the three subtypes of errors were evaluated in the same way.

**Data Analysis and Statistics.** Imaging data were analyzed using ImageJ (NIH). Patch-clamp recording data were analyzed using Clampfit 10.2 (Molecular Devices). GraphPad Prism 7 was used for performing statistics and graphically depicting the population data. Normal distributions were assumed for all datasets in the present study. Population data in the main text are presented as mean ± SEM. Error bars in the figures represented the SEM. An unpaired t test was performed on comparisons of two groups of independent samples, and a paired t test was used for comparing paired data. One-way ANOVA was used to compare multiple independent groups. Statistical significance was preset at P < 0.05.

**ACKNOWLEDGMENTS.** We thank Leo Yan, Curtis Wong, Howard Chan, and Ada Fu for their assistance in some of the experiments and Prof. Savio Chan of the Feinberg School of Medicine, Northwestern University for generous support of this study. This work was supported by Hong Kong Research Grants Council (HKRGC-General Research Fund Grants 14111715 (to W.-H.Y. and Y.K.) and 14107616 (to Y.K. and W.-H.Y.), HKRGC-Collaborative Research Fund Grants C6003-14G (to N.Y.I. and W.-H.Y.) and C6042-15G (to W.-H.Y.), and Area of Excellence Grant AoEM-604/16 (to N.Y.I. and W.-H.Y.).


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