Competitive regulation of synaptic Ca²⁺ influx by D2 dopamine and A2A adenosine receptors

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Striatal D2-type dopamine receptors (D2Rs) have been implicated in the pathophysiology of neuropsychiatric disorders, including Parkinson's disease and schizophrenia. Although these receptors regulate striatal synaptic plasticity, the mechanisms underlying dopaminergic modulation of glutamatergic synapses are unclear. We combined optogenetics, two-photon microscopy and glutamate uncaging to examine D2R-dependent modulation of glutamatergic synaptic transmission in mouse striatopallidal neurons. We found that D2R activation reduces corticostriatal glutamate release and attenuates both synaptic- and action potential—evoked Ca²+ influx into dendritic spines by approximately 50%. Modulation of Ca²+ signaling was mediated by a protein kinase A (PKA)-dependent regulation of Ca²+ entry through NMDA-type glutamate receptors that was inhibited by D2Rs and enhanced by activation of 2A-type adenosine receptors (A2ARs). D2Rs also produced a PKA- and A2AR-independent reduction in Ca²+ influx through R-type voltage-gated Ca²+ channels. These findings reveal that dopamine regulates spine Ca²+ by multiple pathways and that competitive modulation of PKA controls NMDAR-mediated Ca²+ signaling in the striatum.

The striatum is the first processing stage in the basal ganglia, a collection of forebrain nuclei whose function is critical for the generation of coordinated, purposeful movements¹. Regulation of striatal activity by dopamine is important for a variety of psychomotor functions, including habit learning and serial movement, and perturbation of dopaminergic signaling is central to the pathogenesis of several neuropsychiatric diseases²⁻⁴. Dopamine release in the striatum activates two classes of G protein-coupled receptors that are expressed by distinct subpopulations of medium spiny neurons (MSNs), the principal striatal cells. D2Rs, expressed by striatopallidal MSNs, are linked to behavioral disorders, including Parkinson's disease, obsessivecompulsive disorder and schizophrenia^{2,4,5}. D2Rs are coupled to $G\alpha_i^{2,3,6}$, whose activation triggers a divergent cascade of signaling pathways, reducing cAMP production and PKA activity while simultaneously enhancing phospholipase C (PLC)-dependent processes and the release of Ca²⁺ from internal stores^{3,4,6}. Striatopallidal MSNs also express the Gα_c-coupled A2AR, whose activation elevates cAMP and PKA activity and may counteract the actions of D2Rs⁷. Nevertheless, the cellular targets of dopaminergic and adenosinergic signaling are not well characterized.

The expression of D2Rs on dendritic shafts and spines, as well as on afferent glutamatergic terminals, suggests that an important feature of D2R-mediated modulation may be the control of excitatory synaptic transmission⁸. Activation of D2Rs has been reported to inhibit postsynaptic glutamate receptors stimulated by exogenous agonists⁹ and to reduce glutamate release, although it remains unclear whether this effect is mediated by pre- or postsynaptic D2Rs^{10,11}. D2R activation also reduces whole-cell Ca²⁺ influx through voltage-gated Ca²⁺ channels (VGCCs)^{12–14}, although the consequences for synaptic Ca²⁺

influx are unknown. Indeed, recent work suggests that D2Rs might indirectly boost synaptic Ca^{2+} influx by decreasing acetylcholine release from local interneurons, relieving cholinergic inhibition of dendritic VGCCs¹⁵. Functionally, D2R activation promotes long-term depression (LTD) of striatal excitatory synapses while reducing NMDA-type glutamate receptor (NMDAR)-dependent long-term potentiation (LTP)^{16–19}. The former is thought to rely on a D2R-mediated, PLC-dependent enhancement of endocannabinoid release^{20,21}, whereas the mechanisms underlying the latter are unknown.

To gain insight into D2R actions during synaptic transmission, we combined electrophysiological recordings and two-photon imaging of striatopallidal MSNs with optical activation of glutamatergic synapses using channelrhodopsin-2 (ChR2) and focal glutamate uncaging. We found that D2Rs reduced corticostriatal glutamatergic transmission without directly altering current through postsynaptic glutamate receptors. Instead, D2R activation selectively inhibited Ca²⁺ influx through both NMDARs and R-type VGCCs. Regulation of NMDARs was mediated by a PKA-dependent pathway and could be prevented by coactivation of A2ARs. In contrast, modulation of R-type Ca²⁺ influx was independent of PKA and A2AR activity. Our findings indicate that D2Rs engage a divergent signaling cascade controlling synaptic Ca²⁺ influx and suggest that A2ARs and D2Rs compete to regulate Ca²⁺ influx through NMDARs.

RESULTS

To investigate D2R actions on glutamatergic responses in striatopallidal MSNs, we made whole-cell current-clamp recordings from green fluorescent protein (GFP)-positive cells in acute slices prepared from mice expressing GFP under control of the Drd2 gene promoter (**Fig. 1a**) 15,22 .

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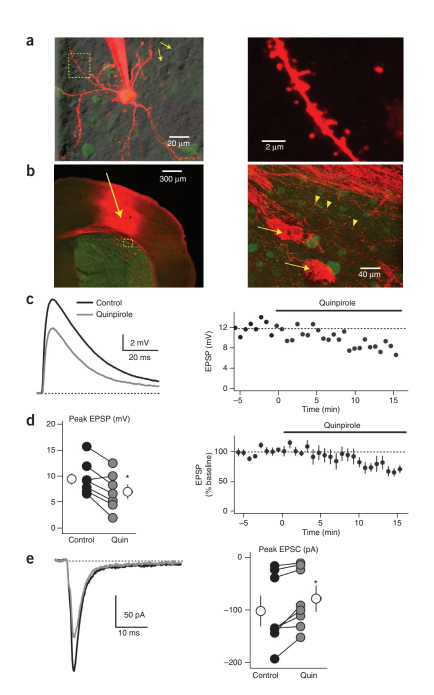


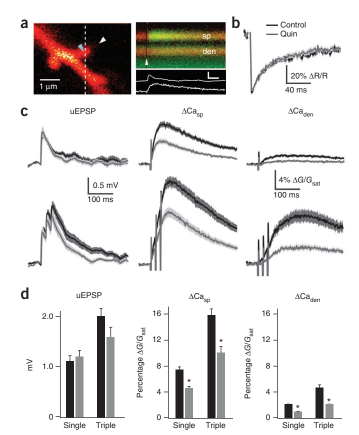
Figure 1 Modulation of corticostriatal synaptic responses in striatopallidal MSNs by D2Rs. (a) Left, two-photon laser-scanning microscopy image of a GFP-positive D2R-expressing striatopallidal MSN filled with the red fluorophore Alexa Fluor 594 through the recording pipette. Nearby nonrecorded GFP-positive cells are also visible. GFP-negative, putatively D2R-negative MSNs are shown by simultaneous laser-scanning differential interference contrast (yellow arrows). Right, higher magnification image of the region bounded by the yellow box on the left. Spines are visible along the dendritic branch. (b) Left, confocal image of a coronal section from the striatum of an Emx1-Cre/Drd2-GFP mouse. Green fluorescence indicates GFP expression in D2Rexpressing MSNs. Red fluorescence indicates the presence of the Channelrhodopsin-mCherry fusion protein in corticostriatal cells and axons. Arrow indicates site of adenoassociated virus injection, 10 d prior. Right, higher magnification image of the region bounded by the yellow box on the left. Arrows indicate large bundles of descending corticobulbar fibers passing through the striatum. Arrowheads indicate presumed presynaptic varicosities of axonal branches arborizing in the striatum. (c) Left, lightevoked EPSPs recorded from a striatopallidal MSN in control conditions and after wash-in of 8 μ M quinpirole. EPSPs are averages of five consecutive trials. Right, time course of the evoked EPSP amplitude from the experiment shown on the left. Quinpirole was applied during the time indicated by the horizontal bar. (d) Left, peak values for light-evoked EPSPs for each recorded cell (n = 7, closed circles) in control conditions (black) and after quinpirole wash-in (gray). The average population data are also shown (open circles). Right, average (n = 7) time course of the evoked EPSP amplitude for the cells shown on the left. Quinpirole was applied during the time indicated by the horizontal bar. (e) Left, light-evoked EPSCs recorded from a striatopallidal MSN in control conditions and after wash-in of 8 µM quinpirole. EPSCs are averages of five consecutive trials. Right, peak values for light-evoked EPSCs for each recorded cell (n = 8, closed circles) in control conditions (black) and after quinpirole wash-in (gray). The average population data are also shown

Membrane potential was held near -60 mV, and experiments were carried out in the presence of the muscarinic antagonist scopolamine.

To examine modulation of corticostriatal synapses, we expressed ChR2 in motor cortex neurons projecting to the striatum (**Fig. 1b**, Online Methods). Pulses of blue light applied to the striatum surrounding the current-clamped MSN reliably evoked excitatory postsynaptic potentials (EPSPs; **Fig. 1c**). Bath application of the D2R agonist quinpirole reduced the average EPSP amplitude from 9.6 \pm 1.2 mV to 7.0 \pm 1.4 mV (n=7, Student's paired t test, P<0.01; **Fig. 1d**). In voltage-clamp recordings, application of quinpirole reduced light-evoked excitatory postsynaptic currents (EPSCs) from 101.4 ± 23.2 pA to 78.6 ± 19.4 pA (n=8, P<0.01; **Fig. 1e**). Notably, quinpirole failed to alter EPSPs evoked by local electrical stimulation (**Supplementary Fig. 1**), suggesting that electrical stimulation may trigger local release of dopamine, occluding the actions of exogenous agonists¹¹.

Previous studies found that D2R activation reduces glutamate release from corticostriatal terminals 10,11 , potentially explaining the inhibition of light-evoked EPSPs and EPSCs. Evidence for direct D2R modulation of postsynaptic glutamate receptors has been less conclusive. We combined two-photon laser-scanning microscopy and glutamate uncaging to bypass presynaptic terminals and examine modulation of glutamatergic responses at single postsynaptic sites. Cells were filled with the $\rm Ca^{2+}$ -insensitive red fluorophore Alexa Fluor 594 and the $\rm Ca^{2+}$ -sensitive green fluorophore Fluo-5F. Under current clamp, a 500- μs pulse from the uncaging laser directed 0.5 μm from a dendritic spine located within 60 μm of the cell body produced a small uncaging-evoked EPSP (uEPSP) and a $\rm Ca^{2+}$ -dependent increase in green fluorescence in the spine head and neighboring dendritic shaft (Fig. 2a). Fluorescence transients in the spine and dendrite were quantified as the percent increase in green signal relative to

(open circles). *P < 0.05.



the maximal green fluorescence in saturating Ca²⁺ ($\Delta G_{\rm sp}/G_{\rm sat}$ and $\Delta G_{\rm den}/G_{\rm sat}$, respectively; see Online Methods). For each synapse, laser power was adjusted such that the uncaging pulse applied directly to the spine head bleached ~50% of the red fluorescence (**Fig. 2b**)²³, and the periphery of the spine head was probed to find the uncaging position that evoked the largest uEPSP²⁴.

Under control conditions, a single uncaging pulse produced an average (n = 36) uEPSP and associated Ca²⁺ transients in the spine and neighboring dendritic shaft with amplitudes of 1.1 \pm 0.1 mV, 7.4 \pm 0.5% and 1.9 \pm 0.2%, respectively (**Fig. 2c,d**). Because MSN dendrites exhibit voltage-dependent conductances that can shape temporal integration of synaptic inputs^{25,26}, we also measured the average uEPSP (2.0 \pm 0.2 mV) and Ca²⁺ transients ($\Delta G_{\rm sp}/G_{\rm sat}$ = 15.8 \pm 1.1%, $\Delta G_{\rm den}/G_{\rm sat} = 4.7 \pm 0.2\%$) evoked by a burst of uncaging pulses (three stimuli, 20-ms interstimulus interval; Fig. 2c,d). In the presence of quinpirole (n = 23), there was no significant change in average peak uEPSP magnitude for either single (1.2 \pm 0.1 mV, P = 0.61) or triple $(1.6 \pm 0.2 \text{ mV}, P = 0.14) \text{ stimuli } (\text{Fig. 2c,d}) \text{ compared with control}$ conditions, although the decay of the triple response was slightly faster, possibly indicating changes in voltage-gated ion channel or NMDAR contributions. In contrast, quinpirole strongly reduced uncagingevoked Ca²⁺ influx following both single ($\Delta G_{\rm sp}/G_{\rm sat} = 4.2 \pm 0.4\%$, P < 0.0001; $\Delta G_{\text{den}}/G_{\text{sat}} = 0.9 \pm 0.1\%$, P < 0.0001) and triple ($\Delta G_{\text{sp}}/G_{\text{sat}} = 0.9 \pm 0.1\%$). 9.4 ± 0.8%, P < 0.0001; $\Delta G_{\rm den}/G_{\rm sat} = 1.9 \pm 0.2$ %, P < 0.001) stimuli (Fig. 2c,d). Quinpirole did not alter the kinetics of recovery following photobleaching of the red fluorophore (50.7 \pm 5.2 ms versus 58.6 \pm 10.4 ms for control and quinpirole, respectively, P = 0.45; Fig. 2b), indicating a lack of change in the diffusional properties of the spine. Furthermore, temporal summations of uEPSPs and Ca²⁺ transients were not significantly altered by quinpirole (P > 0.05 for all comparisons; Supplementary Fig. 2). In summary, when bypassing the presynaptic

Figure 2 D2R modulation of uncaging-evoked Ca²⁺ transients in active spines. (a) Left, two-photon laser-scanning microscopy image of a spiny region from a striatopallidal MSN dendrite filled with the red fluorophore Alexa Fluor 594 and the green Ca²⁺-sensitive indicator Fluo 5F. Right, red and green fluorescence in the spine head (sp) and neighboring dendrite (den) measured in line scan over the region indicated by the dashed line on the left. The white arrowheads on the left and right indicate the location and timing, respectively, of a 500-µs pulse of 725-nm laser light used to focally uncage glutamate near the spine head. The increase in green fluorescence indicates increased intracellular Ca2+. Background green fluorescence visible at the bottom of the image is a result of GFP in an adjacent unrecorded MSN. The corresponding uEPSP and spine Ca²⁺ transient (white traces) are shown below. Inset scale bars represent 1 mV, 5% $\Delta G/G_{sat}$, 50 ms. (b) Recovery of red fluorescence following photobleaching was used to set laser power. The uncaging laser was directed to the spot indicated by the blue arrowhead in a and red fluorescence was measured in line scan from the region indicated in a. Power was adjusted such that a 500-µs pulse produced ~50% reduction in red fluorescence. Average photobleaching transients are shown for control conditions (black, n = 36) and in the presence of quinpirole (gray, n = 23). Solid lines indicate the mean and the shaded regions indicate the mean \pm s.e.m. (c) Uncagingevoked uEPSPs (left), $\Delta {\rm Ca_{sp}}$ (middle) and $\Delta {\rm Ca_{den}}$ (right) produced by single (upper traces) or 50-Hz triple (lower traces) uncaging stimuli are shown for control conditions (black) and in the presence of quinpirole (gray). Mean (solid lines) and mean \pm s.e.m. (shaded regions) are shown. (d) Population data for peak uEPSP (left), ΔCa_{sp} (middle) and ΔCa_{den} (right) evoked by single or triple uncaging stimuli in control conditions (black) and in the presence of quinpirole (gray) are shown. *P < 0.05.

terminal, activation of D2Rs in striatopallidal MSNs reduced synaptic Ca^{2+} influx by nearly 50% with minimal effects on somatic potentials.

Multiple sources contribute to synaptic Ca2+ transients

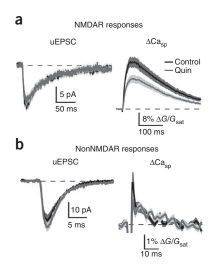
To identify potential targets of dopaminergic regulation underlying the inhibition of Ca²⁺ influx, we examined the Ca²⁺ sources that contribute to synaptic signaling in striatopallidal MSNs. Blockade of NMDARs with the selective antagonist 3-(2-carboxypiperazin-4-yl)propyl-1phosphonic acid (CPP) significantly increased the average (n = 21)amplitude of the single stimulus uEPSP to $146.3 \pm 20.7\%$ of the control value (P < 0.05; Fig. 3). This counterintuitive result may reflect the role of NMDARs in promoting the opening of SK-type Ca²⁺-activated potassium channels that inhibit synaptic potentials in a variety of cell types^{23,27}. Indeed, application of the SK channel blocker apamin (0.1 μ M) also significantly increased uEPSP amplitude (P < 0.001; Supplementary Fig. 3). A 50-Hz burst stimulus revealed that temporal summation was decreased by NMDAR blockade as a result of reduced uEPSP duration (Fig. 3a and Supplementary Fig. 2). CPP reduced the uncaging-evoked Ca²⁺ influx in the spine head to 14.0 \pm 1.3% (P < 0.0001) and $9.1 \pm 1.0\%$ (P < 0.0001) of control values for single and triple stimuli, respectively (Fig. 3), consistent with a dominant contribution of NMDARs to synaptically evoked Ca²⁺ transients.

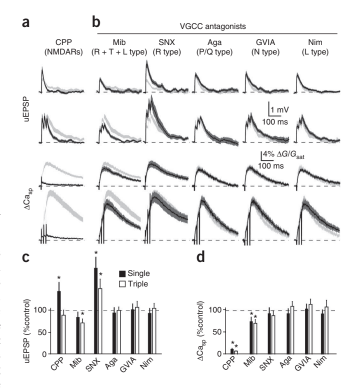
We also analyzed the contribution of VGCCs to synaptic signaling. Selective blockade of P/Q- (n=24), N- (n=26) or L-type (n=26) channels (with 0.2 μ M ω -agatoxin-IVA, 1 μ M ω -conotoxin GVIA or 3 μ M nimodipine, respectively) had no significant effect on synaptic potentials and Ca²⁺ transients evoked by single or triple stimuli (P>0.05 for all comparisons; **Fig. 3b-d**). These blockers also failed to alter the temporal summation of uncaging-evoked potentials and Ca²⁺ transients (**Supplementary Fig. 2**). As shown previously²⁸, 20 μ M nimodipine reduced Ca²⁺ influx (**Supplementary Fig. 4**), indicating that either channel selectivity decreases at this high concentration²⁹ or that relatively dihydropyridine-resistant Ca_V1.3 L-type channels are present³⁰.

Combined blockade of R-, L- and T-type channels with mibefradil had no effect on the average (n = 18) uEPSP evoked by a

single stimulus, although the uEPSP following the triple stimulus was reduced relative to control (Fig. 3b,c). However, mibefradil reduced the average relative amplitude of the Ca²⁺ transients evoked by both single (77.2 \pm 9.5%, *P* < 0.05) and triple (72.9 \pm 9.1%, P < 0.01) stimuli (**Fig. 3b,d**). In contrast, selective blockade of R-type channels with SNX-482 produced a significant increase in average (n = 20) uEPSP magnitude (P < 0.001; **Fig. 3b,c**). These results suggest that, similar to the hippocampus, blocking R-type channels disengages a negative feedback loop involving SK channels, thereby boosting synaptic potentials and NMDAR-mediated Ca2+ influx23. The lack of uEPSP boosting and the reduction in burst-evoked uEPSPs by mibefradil suggests that the broad actions of this nonselective agent may influence the generation of synaptic potentials. In summary, our results indicate that the largest fraction of spine Ca²⁺ influx evoked by activation of a single postsynaptic terminal in striatopallidal MSNs arises from NMDAR opening, with additional contributions from R- and either T- or dihydropyridine-resistant L-type VGCCs, and suggest that these sources are potential targets for dopaminergic control of synaptic Ca²⁺ influx.

D2Rs inhibit NMDAR- and R-type VGCC-mediated Ca²⁺ influx To examine the potential modulation of NMDAR-mediated synaptic responses, we voltage-clamped MSNs at -70 mV using a cesium-based internal solution and bathed them in artificial cerebrospinal fluid (ACSF) containing nominally zero extracellular Mg²⁺, tetrodotoxin, a full cocktail of VGCC antagonists (Online Methods) and the AMPA-type glutamate receptor (AMPAR) antagonist





6-nitro-2,3-dioxo-1,4-dihydrobenzo[f]quinoxaline-7-sulfonamide (NBQX). Under these conditions, glutamate uncaging evoked an average (n = 30) excitatory postsynaptic current (uEPSC, $13.0 \pm 1.4 \, pA$) and associated Ca²⁺ transient in the spine head $(\Delta G_{sp}/G_{sat})$ = $26.5 \pm 3.0\%$; Fig. 4a). Activation of D2Rs produced no change in the average (n = 27) NMDAR-mediated uEPSC (11.1 \pm 1.3 pA, P = 0.31), but significantly decreased the Ca²⁺ transient ($\Delta G_{sp}/G_{sat}$ = $17.0 \pm 2.1\%$, P < 0.05; Fig. 4a). We performed a similar analysis for nonNMDAR-mediated responses using ACSF with control Mg²⁺ concentration and in the presence of the NMDAR antagonist CPP. The average (n = 19) nonNMDAR-mediated uEPSC (26.0 \pm 3.6 pA) and associated Ca²⁺ transient ($\Delta G_{\rm sp}/G_{\rm sat}=0.7\pm0.1\%$) were not altered in the presence of quinpirole $(n = 16; 30.2 \pm 2.9 \text{ pA},$ P = 0.39; $\Delta G_{\rm sp}/G_{\rm sat} = 0.8 \pm 0.1\%$, P = 0.62; **Fig. 4b**), indicating that D2Rs selectively reduce Ca2+ accumulation resulting from activation of NMDARs.

We determined the actions of D2Rs on VGCCs by examining the modulation of Ca²⁺ transients evoked by back-propagating action potentials (bAPs; **Fig. 5a**). On average (n=30), a single bAP evoked by brief somatic current injection resulted in Ca²⁺ influx in the spine head ($\Delta G_{\rm sp}/G_{\rm sat}=8.8\pm0.01\%$) and neighboring dendrite ($\Delta G_{\rm den}/G_{\rm sat}=6.7\pm0.4\%$; **Fig. 5b**). During a brief burst of three bAPs, Ca²⁺ was further increased in both compartments

Figure 4 Activation of D2Rs modulates NMDAR-mediated, but not nonNMDAR-mediated, synaptic responses. (a) Uncaging-evoked NMDAR-mediated uEPSCs (left) and $\Delta Ca_{\rm sp}$ (right) recorded in voltage clamp in the presence of nominally zero extracellular Mg, the AMPAR antagonist NBQX and a cocktail of VGCC blockers. Mean (solid lines) and mean \pm s.e.m. (shaded regions) are shown for responses in control conditions (black, n=30) and in the presence of quinpirole (gray, n=27). (b) Uncaging-evoked nonNMDAR-mediated uEPSCs (left) and $\Delta Ca_{\rm sp}$ (right) recorded in the presence of normal extracellular Mg, the NMDAR antagonist CPP and a cocktail of VGCC blockers. Mean (solid lines) and mean \pm s.e.m. (shaded regions) are shown for responses in control conditions (black, n=19) and in the presence of quinpirole (gray, n=16).

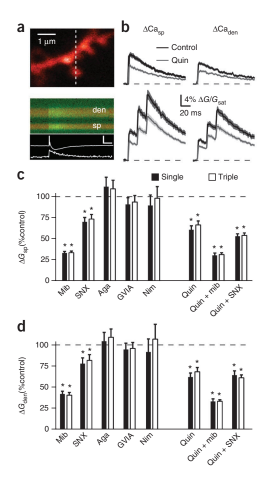
Figure 5 Activation of D2Rs modulates R-type VGCCs. (a) Top, twophoton laser-scanning microscopy image of a striatopallidal MSN dendrite filled with Alexa Fluor 594 and Fluo 5F. Bottom, red and green fluorescence in the spine head (sp) and neighboring dendrite (den) measured in line scan over the region indicated by the dashed line in the upper image. The Ca²⁺-dependent increase in green fluorescence was evoked by a single action potential generated by somatic current injection. The somatic action potential and associated $\Delta \text{Ca}_{\text{sp}}$ are shown below (white traces). Inset scale bars represent 10 mV, 8% $\Delta G/G_{\rm sat}$, 50 ms. (b) bAP-evoked $\Delta Ca_{\rm sp}$ (left) and $\Delta Ca_{\rm den}$ (right) produced by either single or triple (50 Hz) bAPs are shown for control conditions (black, n = 30) or in the presence of quinpirole (gray, n = 34). Mean (solid lines) and mean \pm s.e.m. (shaded regions) are shown. (c) Population data for ΔCa_{sp} , expressed as a percentage of the control value, for single and triple bAP-evoked responses measured in the presence of drugs as indicated: mibefradil (Mib, n = 24), SNX-482 (SNX, n = 23), ω -agatoxin IVA (Aga, n = 23), ω -conotoxin GVIA (GVIA, n = 22), nimodipine (Nim, n = 20), quinpirole alone (n = 34), quinpirole + mibefradil (n = 25) and quinpirole + SNX-482 (n = 23). (d) Data are presented as in c for ΔCa_{den} .

 $(\Delta G_{\rm sp}/G_{\rm sat}=22.0\pm1.6\%,\,\Delta G_{\rm den}/G_{\rm sat}=17.4\pm1.2\%;\,$ Fig. 5b and Supplementary Fig. 2). Activation of D2Rs reduced the average (n = 34) bAP-evoked Ca²⁺ transient for both single $(\Delta G_{sp}/G_{sat} =$ $5.4 \pm 0.5\%$, P < 0.0001; $\Delta G_{\rm den}/G_{\rm sat} = 4.2 \pm 0.4\%$, P < 0.0001) and triple $(\Delta G_{\rm sp}/G_{\rm sat} = 15.0 \pm 1.2\%, P < 0.001; \Delta G_{\rm den}/G_{\rm sat} = 11.9 \pm 0.9\%,$ P < 0.001; **Fig. 5b-d**) stimuli, suggesting that VGCCs are targets of D2R modulation. Quinpirole did not alter the temporal summation (Supplementary Fig. 2) or exponential decay time constants of the bAP-evoked spine Ca²⁺ transients (75.6 \pm 6.3 ms versus 82.6 \pm 9.4 ms for control and quinpirole, respectively, P = 0.55), suggesting a lack of effect on Ca²⁺ clearance.

In the presence of mibefradil, the average (n = 24) single bAPevoked Ca²⁺ signal was reduced to $33.4 \pm 2.8\%$ (P < 0.0001) and $42.6 \pm$ 3.9% (P < 0.0001) of control values for spines and dendrites, respectively, indicating a large contribution of R-, T- and/or L-type channels (Fig. 5c,d). The selective R-type blocker SNX-482 also reduced single bAP-evoked Ca²⁺ influx to 70.0 \pm 5.6% (n = 23, P < 0.0001) and 77.8 \pm 7.3% (P < 0.01) of control values for spines and dendrites, respectively (Fig. 5c,d). In contrast, there was no change in Ca²⁺ influx in the presence of ω -agatoxin IVA (n = 23), ω -conotoxin-GVIA (n = 22) or 3 μ M nimodipine (n = 20) (Fig. 5c,d). Similar results were seen following bursts of three bAPs (Fig. 5c,d), and none of the VGCC blockers altered the temporal summation of bAP-evoked Ca²⁺ signals (Supplementary Fig. 2). These results suggest that R-type as well as T- and/or dihydropyridine-resistant L-type channels contribute to bAP-evoked Ca²⁺ influx.

We next examined the ability of mibefradil and SNX-482 to mimic and occlude the actions of quinpirole (Fig. 5c,d). A one-way ANOVA revealed significant differences between the effects of mibefradil, SNX-482, quinpirole alone and each of the VGCC blockers in the presence of quinpirole (F = 13.26, P < 0.0001 and F = 11.30, P < 0.0001 for spines and dendrites, respectively) on single bAPevoked Ca²⁺ transients. Tukey's multiple post hoc comparisons revealed that SNX-482 mimicked and occluded the actions of quinpirole, while mibefradil produced a significantly larger reduction in Ca²⁺ influx than either SNX-482 or quinpirole alone (comparisons significant for P < 0.05). Similar results were obtained for the triple bAP stimuli, arguing that D2Rs modulate R-type VGCCs, with minimal effect on T- or L-type channels.

Previous studies have linked dopaminergic regulation of Ca_v1.3 L-type channels with MSN spine stability³¹. To examine whether these channels were present, but either inactive or insensitive to dihydropyridine block³⁰, during brief synaptic or bAP stimuli, we measured



spine Ca2+ influx during 300-ms voltage steps from -70 to 0 mV (Fig. 6a). In the presence of blockers for R-, N- and P/Q-type VGCCs, we observed a large Ca²⁺ transient that rapidly reached steady state and decayed following return of the holding potential to baseline. Nimodipine (3 μ M) reduced the step-evoked transients from 60.2 \pm 2.4% (n = 22) to 43.7 \pm 2.2% (n = 21) (**Fig. 6b**). Activation of D2Rs decreased the amplitude to $36.7 \pm 1.4\%$ (n = 20), whereas coapplication of quinpirole and nimodipine produced an additional reduction to $21.4 \pm 2.6\%$ (n = 20; Fig. 6b). A one-way ANOVA comparing control, quinpirole, nimodipine and both quinpirole and nimodipine revealed significant differences between these groups (F = 54.6, P < 0.0001). Post hoc analysis revealed that quinpirole and nimodipine both significantly reduced Ca²⁺ influx relative to control and that the two together produced significantly greater reduction than either alone (comparisons significant for P < 0.05). However, calculation of the nimodipine-sensitive component (the arithmetic difference between pre- and post-nimodipine values) indicated that the contribution of L-type VGCCs was unaffected by quinpirole (16.5 \pm 2.3% versus 15.3 \pm 2.1% $\Delta G/G_{sat}$, in control and quinpirole, respectively, P > 0.05). In summary, our data indicate that R-, L- and T-type VGCCs are present in MSN spines, although R-types are the primary channel regulated by D2Rs.

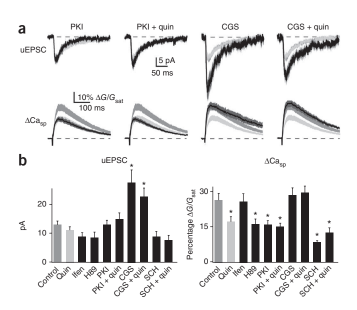
D2Rs act via PKA-dependent and PKA-independent pathways

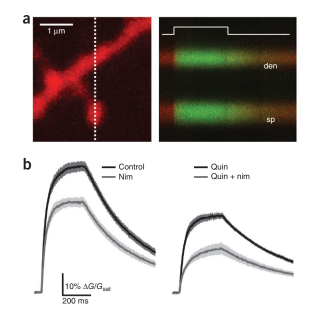
Studies in hippocampal neurons have indicated that a PKA-dependent mechanism augments Ca²⁺ influx through NMDARs without altering total current flow³². We therefore examined whether D2R-mediated inhibition of PKA is responsible for the reduction in NMDARmediated Ca2+ influx. We isolated NMDAR-mediated responses as

Figure 6 Voltage-steps reveal L-type Ca^{2+} channels in MSN spines. (a) Left, two-photon laser-scanning microscopy image of a striatopallidal MSN dendrite filled with Alexa Fluor 594 and Fluo 5F. Right, red and green fluorescence in the spine head (sp) and neighboring dendrite (den) measured in line scan over the region indicated by the dashed line at left. The increase in Ca^{2+} -dependent green fluorescence was evoked by a 300-ms voltage step from -70 mV to 0 mV at the time indicated by the white line. (b) Left, mean \pm s.e.m. (shaded traces) are shown for responses evoked under control conditions (n = 22, black) and in the presence of nimodipine (n = 21, gray). Right, mean \pm s.e.m. for responses evoked in the presence of quinpirole (n = 20, black) or quinpirole + nimodipine (n = 20, gray).

described above and tested the ability of PKI(14-22), a membranepermeable specific antagonist of PKA, to mimic and occlude the actions of quinpirole. In the presence of PKI(14-22), the average (n = 21) NMDAR-mediated uEPSC was 13.0 ± 1.6 pA and the associated spine Ca²⁺ transient was $15.8 \pm 2.0\%$ and $6.9 \pm 1.2\%$ (Fig. 7a,b). Similar results were found using the alternate PKA antagonist H89 (n = 19; Fig. 7b). Following coapplication of PKI(14-22) and quinpirole (n = 20), the uEPSC was 14.9 ± 2.3 pA and the Ca²⁺ signal was $15.1 \pm$ 1.7% (Fig. 7a,b). A one-way ANOVA comparing control, PKI(14-22), quinpirole and both PKI(14-22) and quinpirole revealed significant differences for Ca^{2+} transients between these groups (F = 6.14, P < 0.01). Post hoc comparisons showed significant differences between control and PKI(14-22), as well as between control and both PKI(14-22) and quinpirole, whereas there was no difference between quinpirole alone and in combination with PKI(14-22) (comparisons significant for P < 0.05). Thus, antagonism of PKA is sufficient to mimic and occlude the actions of D2Rs on NMDAR-mediated Ca²⁺ transients.

Striatopallidal MSNs also express A2ARs, which are positively coupled to PKA via $G\alpha_s^7$ and oppose the actions of D2Rs on long-term synaptic plasticity in the striatum¹⁹. We examined whether these two modulatory pathways also interact in the control of NMDAR-mediated Ca^{2+} influx. Application of the specific A2AR agonist CGS-21680 (n=16) resulted in a uEPSC magnitude of 27.8 \pm 4.5 pA and a spine Ca^{2+} transient of 28.5 \pm 3.2% (**Fig. 7a,b**). Coapplication of CGS-21680 and quinpirole (n=15) resulted in a uEPSC magnitude of 22.7 \pm 3.0 pA and spine Ca^{2+} transient of 29.5 \pm 3.0% (**Fig. 7a,b**). A one-way ANOVA comparing control, quinpirole, CGS-21680 and both CGS-21680 and quinpirole revealed significant differences between these groups (F=3.9, P<0.05). Post hoc analyses indicated that the uEPSC





magnitudes for A2AR and combined A2AR and D2R activation were significantly increased, whereas spine Ca^{2+} transients did not differ from control values (significant for P < 0.05).

We next examined whether basal activation of A2ARs influences Ca²⁺ influx through NMDARs. Application of the specific A2AR antagonist SCH-58261 (n = 19) resulted in a uEPSC magnitude of 8.8 \pm 1.8 pA and a spine Ca^{2+} transient of 8.5 \pm 0.7% (**Fig. 7b**). Coapplication of SCH-58261 and quinpirole (n = 18) resulted in a uEPSC magnitude of 7.6 \pm 1.7 pA and a spine Ca²⁺ transient of 12.7 \pm 2.3% (Fig. 7b). A one-way ANOVA with *post hoc* tests comparing control, quinpirole, SCH-58261 and both SCH-58261 and quinpirole revealed no significant change in uEPSC magnitude (F = 2.521, P = 0.063). However, A2AR antagonism alone or in combination with quinpirole significantly reduced synaptic Ca²⁺ influx, whereas there was no difference between these two groups (significant for P < 0.05). In summary, basal levels of A2AR activation enhance Ca2+ influx through NMDARs and reduction of this activity is sufficient to mimic and occlude the effects of D2R activation. Furthermore, exogenous activation of A2ARs can counteract the D2R-mediated, PKA-dependent reduction in synaptic Ca²⁺ influx, suggesting that modulatory control of PKA exerts a bidirectional influence on NMDAR Ca²⁺ signaling.

In the hippocampus, NMDARs containing the NR2B-type subunit contribute to particularly large Ca²⁺ influx³³, and removal of these subunits might explain the reduction in synaptic Ca²⁺ influx. However, application of the selective NR2B-containing NMDAR

Figure 7 Push-pull modulation of NMDARs by D2Rs and A2ARs is dependent on PKA activity. (a) Uncaging-evoked NMDAR-mediated uEPSCs (upper traces) and Δ Ca_{sp} (lower traces), recorded as in **Figure 4a**. Mean (black solid lines) and mean ± s.e.m. (dark gray shaded regions) are shown for responses evoked in the presence of the PKA antagonist PKI(14-22) alone (first column, n = 21), PKI(14-22) + quinpirole (second column, n = 20), the A2AR agonist CGS-21680 alone (third column, n = 16) or CGS-21680 + quinpirole (fourth column, n = 15). Medium gray and light gray shaded regions indicate the mean ± s.e.m. responses under control and quinpirole conditions, respectively, reproduced from **Figure 4a** for comparison. (b) Population data for peak uEPSC (left) and Δ Ca_{sp} (right) are shown for the conditions illustrated in **a**. Also shown are the population data for the PKA antagonist H89, the NR2B subunit–containing NMDAR antagonist ifenprodil, the A2AR antagonist SCH-58261, and SCH-58261 + quinpirole. *P < 0.05.

Figure 8 Modulation of VGCCs by D2Rs is independent of A2AR and PKA activity. (a) bAP-evoked ΔCa_{sp} (upper traces) and ΔCa_{den} (lower traces) recorded as in **Figure 5b**. Mean (black solid lines) and mean \pm s.e.m. (dark gray shaded regions) are shown for responses evoked in the presence of PKI(14-22) alone (left, n=20), PKI(14-22) + quinpirole (middle, n=17) or CGS-21680 + quinpirole (right, n=11). Medium gray and light gray shaded regions indicate the mean \pm s.e.m. responses under control and quinpirole conditions, respectively, reproduced from **Figure 5b** for comparison. (b) Population data for ΔCa_{sp} (left) and ΔCa_{den} (right) are shown for the conditions illustrated in **a**. *P < 0.05.

antagonist ifenprodil (n = 18) revealed no significant change in uEPSC or Ca²⁺ transient magnitude (P < 0.05; **Fig. 7b**), suggesting that this subgroup of receptors does not contribute substantially to Ca²⁺ influx under control conditions.

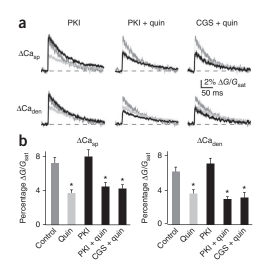
Finally, we examined whether similar PKA-dependent mechanisms underlie the D2R-mediated reduction in bAP-evoked Ca²⁺ signals. A one-way ANOVA revealed significant differences between spine Ca²⁺ transients in control, quinpirole, PKI(14-22) (n = 20), PKI(14-22) and quinpirole (n = 17), and CGS-21680 and quinpirole (n = 11) (F = 10.3, P < 0.0001; **Fig. 8a,b**). Post hoc tests indicated that, in the presence of PKI(14-22), Ca²⁺ influx did not significantly differ from control conditions. Furthermore, coapplication of quinpirole and either PKI(14-22) or CGS-21680 produced a reduction in signal that did not differ from that seen with quinpirole alone (all comparisons significant for P < 0.05). Similar results were obtained for dendrite Ca²⁺ signals (**Fig. 8a,b**). These findings indicate that the actions of D2Rs on bAP-evoked VGCC-mediated Ca²⁺ transients occur independently of PKA and A2AR activation and suggest that D2Rs couple to two independent pathways that converge on a similar endpoint of reduced Ca²⁺ signaling in dendritic spines.

DISCUSSION

Perturbation of D2R-dependent dopaminergic modulation in the striatum is linked to a variety of neuropsychiatric disorders, including Parkinson's disease, obsessive-compulsive disorder and, most notably, schizophrenia^{2,4}. Schizophrenic individuals show evidence of increased density and occupancy of striatal D2Rs34, and most effective antipsychotic agents antagonize D2Rs^{5,35}. Furthermore, overexpression of striatal D2Rs in mice produces behavioral impairments that are thought to model those seen in schizophrenic individuals³⁶. We found that activation of D2Rs reduced glutamate release from cortical afferents and attenuated Ca2+ influx through NMDARs. The regulation of NMDAR Ca2+ signaling occurs via a reduction in PKA activity and can be prevented by coactivation of A2ARs. D2Rs also act via a PKA-independent pathway to reduce Ca²⁺ signaling through SNX-482–sensitive R-type (Ca $_{
m V}$ 2.3) VGCCs. Thus, our results indicate that D2R activation triggers divergent signaling cascades that inhibit multiple synaptic Ca²⁺ sources. Furthermore, we provide evidence of a functional synaptic role for D2R- and A2ARmediated regulation of PKA, implicating this kinase as a target for convergent neuromodulatory control over NMDAR-mediated Ca²⁺ signaling and a likely substrate for control of synaptic plasticity in the striatum.

Regulation of corticostriatal glutamate release

By using mice in which corticostriatal afferents expressed light-activatable ChR2, we were able to selectively characterize the D2R-mediated modulation of cortical inputs to striatopallidal MSNs. Consistent with the results of previous studies, activation of D2Rs reduced the magnitude of evoked postsynaptic potentials and currents¹¹. Coupled with the observation that D2Rs failed to alter



uncaging-evoked synaptic potentials or currents, our data strongly support the hypothesis that D2Rs regulate corticostriatal transmission by reducing glutamate release from afferent terminals^{10,11}. However, we cannot conclude whether pre- or postsynaptic D2Rs, possibly coupled to retrograde endocannabinoid signaling, mediate this phenomenon^{10,11,37}.

In contrast, we found no evidence for direct modulation of postsynaptic nonNMDA-type glutamate receptors. Notably, experiments using electrical stimuli revealed no modulation of glutamatergic responses, suggesting that such stimulation may evoke dopamine release from the terminals of nigrostriatal afferents, thereby occluding the effects of exogenously applied agonists¹¹.

Sources of synaptic Ca2+ influx in striatopallidal MSNs

Our data reveal that, following either single or burst stimulation of an individual synapse, NMDARs contribute the bulk of Ca²⁺ influx, although both Ca²⁺-permeable AMPARs and VGCCs make additional contributions. However, in contrast with hippocampal pyramidal cells³³, we found little contribution of NR2B-containing NMDARs. Similar to hippocampal and cortical pyramidal neurons (reviewed in ref. 38), the major VGCC contributors to synaptic or bAP-evoked spine Ca²⁺ influx are SNX-482- and mibefradil-sensitive channels. Although SNX-482 specifically indicates the presence of R-type (Ca_v2.3) VGCCs, the nonselective mibefradil suggests contributions from T-type (Ca_V3) and/or L-type (Ca_V1.2 and 1.3) channels. Using large voltage steps to directly depolarize the spine, we found a nimodipine-sensitive Ca²⁺ transient, confirming that L-type VGCCs are present in MSN spines³¹. However, these channels either contribute little to synaptic Ca²⁺ influx or are not effectively blocked by nimodipine during brief depolarizations³⁰.

An unexpected finding is that blocking either NMDARs or R-type VGCCs increased uEPSP magnitude, a result mimicked by blocking SK-type Ca²⁺-activated potassium channels. In the hippocampus and amygdala, Ca²⁺ influx through these two sources and subsequent opening of SK channels produces local hyperpolarization and inhibits synaptic responses^{23,27}. Our data suggest that SK channels may have a similar feedback role in MSNs. Notably, quinpirole reduced Ca²⁺ influx through both NMDARs and R-type VGCCs, but did not augment the uEPSP, suggesting that D2Rs may directly regulate SK activation.

Regulation of spine Ca²⁺ signaling by D2Rs

We found that activation of D2Rs reduced synaptically evoked spine Ca²⁺ transients mediated by NMDARs. This effect could result

from a D2R-mediated reduction in the number of NR2B-containing receptors, which exhibit a higher fractional Ca^{2+} permeability³³. However, the failure of ifenprodil to alter synaptic Ca^{2+} transients suggests that these receptors do not contribute substantially under control conditions. Alternatively, D2Rs may regulate NMDAR Ca^{2+} permeability, consistent with previous studies indicating that fractional permeability for divalent ions can be modified by phosphorylation or structural changes^{32,39}. Such mechanisms may also be developmentally regulated, as studies in other brain regions indicate that relative NR2B expression levels and PKA-dependent modulation of NMDAR Ca^{2+} permeability are greater in young animals^{32,40}. As our experiments were performed in juvenile mice, it will be important to determine whether D2R actions on striatal NMDARs are similarly developmentally regulated.

Modulation of NMDAR Ca2+ transients was mimicked and occluded by pharmacologically inhibiting PKA activity, consistent with the biochemical coupling of D2Rs to Gα, and downstream inhibition of cAMP generation and PKA activity⁶. Moreover, blocking the activity of A2ARs, which are positively coupled to cAMP production and PKA activity via $G\alpha_s^7$, similarly reduced NMDAR Ca²⁺ influx and occluded the actions of quinpirole. Furthermore, the actions of D2Rs were prevented by coapplication of an A2AR agonist. These results suggest that basal A2AR tone facilitates NMDARmediated Ca2+ entry, but D2Rs and A2ARs act competitively to control synaptic Ca2+ influx. Activation of A2ARs also boosted the uEPSC amplitude, possibly reflecting enhancement of current through extrasynaptic receptors. Notably, although biochemical experiments have shown that PKA may increase phosphorylation of both AMPARs and NMDARs either directly or indirectly via DARPP-32-dependent inhibition of protein phosphatase 1 (refs. 41,42), our results provide, to the best of our knowledge, the first description of a functional consequence for D2R- or A2ARmediated control of PKA on synaptic transmission. D2R signaling via release of $G_{\beta\gamma}$ subunits is known to produce a membrane-delimited PLC- and calcineurin-dependent inhibition of whole-cell Ca²⁺ currents through VGCCs^{12,13}. This alternative pathway may underlie the previously reported D2R-mediated reduction in bAP-evoked Ca²⁺ influx⁴³ and is consistent with our finding that inhibition of R-type VGCCs was independent of PKA activity. Although reduction in dendritic excitability might also decrease bAP-evoked Ca²⁺ influx through VGCCs by limiting local depolarization⁴³, our analysis of relatively proximal spines (<60 µM from the soma) suggests this mechanism is unlikely to explain our findings. Nevertheless, it is possible that dopaminergic modulation may be different in more distal dendritic regions.

Implications for striatal plasticity

Plasticity of glutamatergic synapses in the striatum is thought to represent an important substrate for both normal and pathological motor learning and is strongly regulated by dopaminergic activity^{2,44}. In striatopallidal neurons, activation of D2Rs during pairing of pre- and postsynaptic activity is sufficient to convert NMDAR-dependent LTP into LTD, and this switch is prevented by coactivation of A2ARs¹⁹. Our finding that D2R and A2AR activities bidirectionally control NMDAR-mediated Ca²⁺ influx provides a potential mechanism for these observations. Notably, striatal LTD, which is dependent on the activation of postsynaptic metabotropic glutamate receptors, Ca²⁺ influx through L-type VGCCs and retrograde endocannabinoid signaling, is disrupted by antagonism of D2Rs^{16,20,21}. In addition, both genetic and pharmacological deletion of D2R signaling converts LTD into LTP^{16,19}. However, the mechanisms underlying D2R control of

LTD are unclear. One hypothesis suggests that D2Rs enhance the release of endocannabinoids from spines via a $G_{\beta\gamma}$ -dependent activation of PLC, leading to retrograde inhibition of glutamate release 11,20,45 . This possibility is supported by our observation that D2R-medidated inhibition of VGCCs occurs via a PKA-independent pathway. Thus, we propose a model of striatal long-term plasticity in which NMDAR-dependent LTP and endocannabinoid-dependent LTD compete to determine net synaptic strength. In this model, D2Rs tip the balance in favor of depression by simultaneously reducing NMDAR-mediated Ca^{2+} influx via a PKA-dependent process, reducing VGCC-mediated Ca^{2+} influx and likely enhancing endocannabinoid production via PKA-independent signaling. In summary, our findings highlight the existence of local neuromodulatory actions on synaptic transmission and provide new insight into the role of dopaminergic modulation in the striatum.

METHODS

Methods and any associated references are available in the online version of the paper at http://www.nature.com/natureneuroscience/.

Note: Supplementary information is available on the Nature Neuroscience website.

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AUTHOR CONTRIBUTIONS

M.J.H. conducted the experiments and data analysis. M.J.H. and B.L.S. designed the experiments and wrote the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Slice preparation and pharmacology. All animal handling was performed in accordance with guidelines approved by the Harvard Institutional Animal Care and Use Committee and federal guidelines. Recordings were made from greenfluorescing MSNs in striatal slices taken from male and female postnatal day 15-22 drd2-GFP mice^{15,22} or Emx-Cre/drd2-GFP mice (see below) that were killed under isoflurane anesthesia. Coronal slices (300 µm thick) were cut in ice-cold external solution containing 110 mM choline, 25 mM NaHCO₃, 1.25 mM NaH₂PO₄, 2.5 mM KCl, 7 mM MgCl₂, 0.5 mM CaCl₂, 25 mM glucose, 11.6 mM sodium ascorbate and 3.1 mM sodium pyruvate, bubbled with 95% O_2 and 5% CO₂. Slices were then transferred to ACSF containing 127 mM NaCl 25 mM NaHCO₃, 1.25 mM NaH₂PO₄, 2.5 mM KCl, 1 mM MgCl₂, 2 mM CaCl₂ and 25 mM glucose, bubbled with 95% O₂ and 5% CO₂. After an incubation period of 30-40 min at 34 °C, the slices were maintained at 22-25 °C until use. All experiments were conducted at 32 °C. In all experiments, the ACSF included 50 μM picrotoxin to block $GABA_{A/C}$ receptor–mediated inhibition and 10 μM scopolamine to block confounding actions of muscarinic acetylcholine receptors. For all glutamate uncaging experiments, 10 μM serine was included in the ACSF to reduce NMDAR desensitization. For some experiments, extracellular MgCl₂ was reduced to nominally 0 mM. Finally, in some experiments, one or more of the following drugs were added to the ACSF: 8 μM quinpirole, 10 μM CPP, 10 μM NBQX, 3 µM ifenprodil, 1 µM tetrodotoxin, 10 µM mibefradil, 0.1 µM SNX-482, $0.2\,\mu\text{M}$ agatoxin IVA, $1\,\mu\text{M}$ ω -conotoxin GVIA, 3 or $20\,\mu\text{M}$ nimodipine, $0.1\,\mu\text{M}$ apamin, 1 μM PKI(14-22), 10 μM H89, 0.2 μM CGS-21680 and 0.1 μM SCH-58261. All chemicals were from Tocris, with the exception of SNX-482, agatoxin IVA and ω -conotoxin GVIA which were from Peptides International.

Electrophysiology and imaging. Whole-cell recordings were obtained from D2R-expressing MSNs identified with video-infrared/differential interference contrast and GFP epifluorescence. Two-photon laser-scanning microscopy confirmed the presence of prominent dendritic spines. For current-clamp recordings, glass electrodes (2–3.5 $\mbox{M}\Omega)$ were filled with internal solution containing 135 mM KMeSO₃, 10 mM HEPES, 4 mM MgCl₂, 4 mM Na₂ATP, 0.4 mM NaGTP and 10 mM sodium creatine phosphate, adjusted to pH 7.3 with KOH. For voltageclamp recordings, cesium was substituted for potassium to improve space clamp. For Ca²⁺-imaging experiments, either 150 μM (Figs. 5 and 8), 600 μM (Fig. 6) or 300 µM (all other experiments) of the Ca²⁺-sensitive indicator Fluo-5F and $10\,\mu\text{M}$ Alexa Fluor-594 were added. As previous work has shown that membrane potential influences the Ca²⁺ sources activated during synaptic transmission²⁸, cells were held near -60 mV for current-clamp recordings to approximate the conditions of MSNs in $vivo^{46}$. Voltage-clamp experiments were performed with a holding potential of -70 mV. All recordings were made using an Axopatch 200B amplifier, and data were filtered at 5 kHz and digitized at 10 kHz. For electrical activation of excitatory input fibers, brief (0.2 ms) current injections were applied to a small glass electrode (tip diameter, $2-4\,\mu m$) filled with ACSF and placed at the border between the striatum and the overlying white matter, $100-300 \mu m$ from the recorded cell.

Intracellular Ca^{2+} imaging and glutamate uncaging were accomplished with a custom microscope, as previously described²⁸. The microscope was assembled using components manufactured by the Harvard Medical School Department of Neurobiology machine shop or Mike's Machine Company. Neurons were filled via the patch electrode for 10-15 min before imaging. Fluo-5F (green) and Alexa Fluor-594 (red) were excited using 840-nm light to monitor Ca^{2+} signals and spine morphology, respectively. To measure Ca^{2+} signals, we collected green and red fluorescence during 500-Hz line scans across a spine and a neighboring dendrite. Reference frame scans were taken between each acquisition to correct for small spatial drift of the preparation over time. Ca^{2+} signals were first quantified as increases in green fluorescence from baseline normalized to the average red

fluorescence ($\Delta G/R$). The elevated resting green fluorescence as a result of GFP expression slightly increased shot noise, but did not alter the Ca²⁺-dependent signal magnitude or the linearity of the response. To allow greater generalization of the data across various microscope configurations, we expressed fluorescence changes as the percentage of the G/R ratio measured in saturating Ca²⁺ ($G_{\rm sat}/R$) in a sealed recording pipette. To calculate $G_{\rm sat}/R$, we imaged a 1:1 mixture of internal solution and 1 M CaCl₂ in the specimen plane under conditions identical to those used during recordings.

For two-photon laser uncaging experiments, MNI-glutamate was bath applied at 2.5 mM and glutamate uncaging was achieved using a 0.5-ms pulse of 725-nm light. To achieve standard uncaging power (which translates into a constant amount of glutamate uncaged on each trial), we used photobleaching of Alexa Fluo-594 in the spine of interest, as previously described²³. Bleaching is a function of the laser power and provides a readout of power delivery that is independent of spine depth in the slice and electrophysiological responses.

Viral ChR2 expression and activation. To selectively express a conditional form of the light-activated membrane channel ChR2 (ref. 47) in corticostriatal afferents, we crossed drd2-GFP mice with mice expressing Cre recombinase under control of the *Emx1* promoter⁴⁸. At postnatal day 12, the F1 offspring of these lines were injected intracranially into somatomotor cortex under isoflurane anesthesia with 1.0 μl of a recombinant adenoassociated virus (serotype 2/8, Harvard Gene Therapy Institute) containing a double *loxP*-flanked inverted construct coding for the fusion protein ChR2-mCherry under control of the *Ef1a* promoter⁴⁷. Mice were killed 7–10 d post-injection for slice preparation as described above and exhibited strong labeling of descending cortical fibers that arborized throughout the dorsal striatum (**Fig. 1b**). To activate ChR2 -positive fibers, we focused light from a 473-nm laser (Optoengine) on to the back aperture of the microscope objective, producing a wide-field exposure around the recorded MSN of 10–20 mW mm⁻². Brief (1–2 ms) pulses of light were delivered under control of the acquisition software.

Data acquisition and analysis. Imaging and physiology data were acquired using National Instruments data acquisition boards and custom software written in MATLAB (Mathworks) 49 . Off-line analysis was performed using custom routines written in MATLAB and Igor Pro (Wavemetrics). The amplitudes of all EPSPs and nonNMDAR-mediated EPSCs were calculated by averaging over a 3-ms window around the peak, whereas a 10-ms window was used to calculate amplitudes of NMDAR-mediated EPSCs. The amplitudes of triplet burst-evoked uEPSPs were calculated around the peak response following the third stimulus. For imaging experiments, measurements of $\Delta G/G_{\rm sat}$ were calculated by taking the average of the signal over a 150-ms post-stimulus window.

In sections describing optical or uncaging responses measured from individual spines, the stated n indicates the number of spines analyzed. In sections describing electrically evoked synaptic responses, the stated n indicates the number of cells analyzed. All summary graphs and data are expressed as mean \pm s.e.m. and comparisons were made using a two-tailed Student's t test unless otherwise stated. Differences were judged to be statistically significant for P < 0.05.

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Erratum: Competitive regulation of synaptic Ca²⁺ influx by D2 dopamine and A2A adenosine receptors

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The sizing of Figure 6 has been modified in the print, PDF and HTML versions of this article.

