Synaptically driven state transitions in distal dendrites of striatal spiny neurons

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Abstract

Striatal spiny neurons (SPNs) associate a diverse array of cortically processed information to regulate action selection. But how this is done by SPNs is poorly understood. A key step in this process is the transition of SPNs from a hyperpolarized ‘down-state’ to a sustained, depolarized ‘up-state’. These transitions are thought to reflect a sustained synaptic barrage, involving the coordination of hundreds of pyramidal neurons. Indeed, in mice simulation of cortical input by glutamate uncaging on proximal dendritic spines produced potential changes in SPNs that tracked input time course. However, brief glutamate uncaging at spines on distal dendrites evoked somatic up-states lasting hundreds of milliseconds. These regenerative events depended upon both NMDA receptors and voltage-dependent Ca$^{2+}$ channels. Moreover, they were bidirectionally regulated by dopamine receptor signaling. This capacity not only changes our model of how up-state are generated in SPNs, it has fundamental implications for the associative process underlying action selection.

Introduction

Striatal spiny neurons (SPNs) are key elements in the basal ganglia circuitry controlling contextually appropriate action selection. SPNs integrate cortically processed information about the environment, internal body state and past experience to facilitate actions that maximize positive and minimize negative outcomes. Dopamine plays a critical role in shaping the activity of SPNs, providing feedback about the outcome of choosing a particular action.

The integrative mechanisms underlying the capacity of SPNs to differentiate one context from another in the process of promoting an appropriate action are poorly defined. A potentially important clue as to how this is done comes from in vivo recordings from SPNs showing that their somatic membrane rapidly moves between a stable hyperpolarized state...
near the K⁺ equilibrium potential (‘down-state’) to a depolarized state near spike threshold (‘up-state’). The 15–20 mV transition from the down- to up-state depends upon a temporally convergent barrage of excitatory, glutamatergic synaptic input from the cerebral cortex.

Because the connectivity between any one cortical pyramidal neuron and an individual SPN is sparse, consisting of a few en passant synapses, a successful transition has been hypothesized to require the synchronization of a great many cortical pyramidal neurons, creating a coincidence threshold that could be used to discriminate context.

Although induced by cortical synaptic input, up-states have a number of features — a stereotyped voltage trajectory and durations of hundreds of milliseconds or more — that suggest they are the product of an intrinsic bistability. However, electrophysiological studies using somatic electrodes have failed to find evidence of bistability in SPNs. This conclusion is consistent with recent work showing that backpropagation of somatic action potentials into dendrites is decrementing and lacks any obvious intrinsic boosting by ion channels carrying inward, depolarizing current. If up-state initiation and maintenance is dependent upon sustained synaptic depolarization, then typical up-states lasting hundreds of milliseconds must require the coordination of hundreds, if not thousands, of cortical pyramidal neurons. This hypothetical feature of synaptic integration in SPNs clearly constrains the corticostriatal network engaged in action selection and limits its pattern recognition capacity.

One of the limitations of the pertinent studies to date is that they have not been able to directly interrogate dendrites, particularly distal dendrites. SPN dendrites are too small to record from with electrodes. To overcome this obstacle, a combination of optical approaches — two photon laser scanning microscopy (2PLSM) and two photon laser uncaging (2PLU) of glutamate — were used in combination with somatic patch clamp methods to study SPNs in brain slices from transgenic mice in which D₁ and D₂ dopamine receptor expression was reported by green fluorescent protein (GFP) — allowing the identification of direct pathway (d)SPNs and indirect pathway (i)SPNs. In agreement with previous studies, the proximal dendritic membrane exhibited little evidence of intrinsic bistability that might contribute to state transitions. However, in distal dendrites of both populations of SPNs, uncaging glutamate at about ten adjacent spines on a single dendrite was capable of evoking a local regenerative event that produced a somatic up-state lasting hundreds of milliseconds. These up-states were dependent upon activation of NMDA receptors and voltage-dependent Ca²⁺ channels. Furthermore, both adenosine and dopamine receptor signaling modulated up-state duration.

**Results**

**Somatic up-states were triggered by distal synaptic input**

Previous studies have shown that SPN up-states are not a product of an intrinsic bistability in the soma and proximal dendrites. However, because of their small size, distal dendrites, which can be electrotonically remote from the soma, have not been directly examined using patch clamp techniques. To interrogate these regions, SPNs were visualized by using a somatic patch electrode containing the dye Alexa 568. This allowed both proximal (40–60 μm from the soma) and distal (>100 μm from the soma) dendrites and
spines to be seen using 2PLSM. 2PLU of bath applied MNI-caged-glutamate was used to activate glutamate receptors at the heads of spines, which are the sites of corticostriatal and thalamostriatal synapses. The uncaging laser power was adjusted to produce simulated excitatory postsynaptic potentials (uEPSPs) at the soma resembling those evoked by stimulation of glutamatergic fibers (Supplementary Fig. 1). Neighboring spines were activated in rapid succession (2 ms inter-event time) to mimic convergent cortical input (Fig. 1a). When performed at distal dendrites (115 ± 6 μm from soma, n=8, Fig. 1b–d), this stimulation paradigm evoked a rapid, stereotyped somatic depolarization after which the membrane potential plateaued, extending well beyond the end of the stimulation, often lasting for hundreds of milliseconds. The somatic membrane potential during these distally evoked plateaus closely resembled that found in vivo during up-states (Fig. 1b). The threshold for evoking a sustained somatic depolarization was about 11 spines (11.9 ± 0.4, n=39) within a 20 μm stretch of distal dendrite (21 ± 1 μm, n=39, Fig. 2d) greater than 100 μm from the soma (113 ± 2 μm, n=39). The duration of the resulting membrane potential plateau was not dependent upon the direction (toward or away from the soma) of spine stimulation (Supplementary Fig. 2).

Once the laser power and spine number required to induce a state transition were determined by stimulating distal dendrites, the identical laser power was then used to stimulate the same number of spines in a proximal dendrite of the same cell (49 ± 1 μm from soma, n=8). Somatic uEPSPs generated by proximal uncaging of glutamate were similar in amplitude, kinetics and evoked spine Ca²⁺ transients (as measured using the high affinity Ca²⁺ indicator Fluo 4) to those evoked distally (Supplementary Fig. 3). Although somewhat surprising, the similarity in uEPSPs was attributable to anatomical features of the SPN dendrite (Supplementary Fig. 3). This synaptic scaling translated to endogenously generated spontaneous EPSPs, as locally increasing spontaneous neurotransmitter release to distal and proximal dendrites (via local application of high osmolarity ACSF) yielded identical somatic EPSP amplitude distributions (Supplementary Fig. 4). Stimulation of clustered spines on proximal dendrites produced a rapid somatic depolarization, but unlike the situation when distal spines were stimulated, the membrane potential rapidly decayed back to the resting membrane potential when the stimulation ended (Fig. 1c,d). This relationship was observed at both room and elevated temperatures (Supplementary Fig. 5).

Though it was not feasible to directly measure dendritic voltage, changes in dendritic Ca²⁺ concentration produced by glutamate uncaging were measurable, providing an indirect assay of dendritic voltage. Robust Ca²⁺ transients were observed in distal dendritic spines and shafts following glutamate uncaging that led to up-state generation. Ca²⁺ transients were measured using the low affinity calcium indicator Fluo 4FF, to avoid dye saturation (Fig. 1e). These experiments revealed a strong correlation between the duration of the somatic up-state and the duration of the distal Ca²⁺ signal (Fig. 1f).

Although uEPSPs of a millivolt or more were used to generate state transitions in the experiments described thus far, smaller, microvolt uEPSPs were also capable of triggering these events (Fig. 2). There was no correlation between uEPSP amplitude and up-state duration once a threshold number of spines had been stimulated (Fig. 2b). However, these experiments were difficult to conduct with the radially oriented dendrites of SPNs because
they typically required 15 or more co-planar spines. As a consequence, subsequent studies were performed with uncaging events that produced 1–3 mV uEPSPs (e.g., Fig. 2d).

There are significant differences between dSPNs and iSPNs in the propagation of somatic spikes into dendrites. However, both cell types responded to distal synaptic stimulation in much the same way (Fig. 3a,b). There were no differences in up- and down-state somatic membrane potentials of the two SPNs (Fig. 3c). Nor were there any differences in the density or spatial distribution of uncaging sites required for up-state generation (Fig. 3d,e). Moreover, in neither cell type was there a relationship between the number of stimulated spines and up-state duration (Fig. 3f), just as one would expect of a non-linear process with a threshold.

**Up-states required NMDA receptors and calcium channels**

Because previous studies in pyramidal neurons have implicated NMDA receptors in synaptically evoked regenerative dendritic events, our working hypothesis was that the state transitions evoked by distal glutamate uncaging in SPNs had similar determinants. To test this hypothesis, the NMDA receptor antagonist ((2R)-amino-5-phosphonovaleric acid (AP5, 100 μM) was bath applied. AP5 dramatically shortened somatic up-state duration in response to glutamate uncaging at distal spines (Fig. 4a).

Voltage-dependent ion channels also are potential contributors to regenerative events in dendrites. In the apical dendrite of pyramidal neurons, voltage-dependent Na\(^+\) channels contribute to spike propagation. This role is more modest in SPNs and bath application of the Nav1 Na\(^+\) channel antagonist tetrodotoxin (TTX, 1–2 μM) had no effect on up-states evoked by distal uncaging (Supplementary Fig. 6a). Voltage-dependent Ca\(^{2+}\) channels also contribute to dendritic electrogenesis. In SPNs, L-type Ca\(^{2+}\) channels prolong the depolarization produced by a brief somatic current pulse, and these channels are found in SPN dendrites and spines. However, the regenerative responses to uncaging glutamate were unaffected by bath application of the Cav1.2/1.3 L-type channel antagonist isradipine (5 μM; Supplementary Fig. 6b) or the mamba toxin calciseptine (0.4 μM; Supplementary Fig. 6c). Hence, while Cav1.3 L-type Ca\(^{2+}\) channels are certainly activated near the somatic up-state potential, they do not appear to be necessary for the maintenance of distally generated plateau potentials. Although L-type Ca\(^{2+}\) channels were not necessary, other voltage-dependent Ca\(^{2+}\) channels did appear to be involved as the inorganic Ca\(^{2+}\) channel antagonist Ni\(^{2+}\) (50 μM) effectively reduced plateau duration (Fig. 4b). SPNs express Cav3.2 and Cav3.3 mRNAs that code for the pore forming subunit of low-threshold T-type Ca\(^{2+}\) channels. Although not present in the somatic and proximal dendritic membrane, these channels are found in more distal dendrites and spines. Consistent with their participation in the dendritic regenerative event, the T-type channel antagonist mibefradil (20 μM) significantly decreased the duration (but not amplitude) of somatic up-states evoked by uncaging (Fig. 4c). Another voltage-dependent Ca\(^{2+}\) channel found in the dendrites of SPNs is the R-type Ca\(^{2+}\) channel Cav2.3. The tarantula toxin SNX-482 (0.3 μM), a selective R-type channel antagonist, significantly decreased the duration of somatic up-states evoked by uncaging (Fig. 4d). Thus, T- and R-type Ca\(^{2+}\) channels participate in the maintenance of distally initiated up-states.
The results presented thus far suggest that distal SPN dendrites are capable of regenerative events that depend upon glutamate-bound NMDA receptors. If this were true, then prolonging the opening of NMDA receptors should increase up-state duration and possibly up-state amplitude. To test this hypothesis, NMDA and glycine (10 μM) were added to the bath to sustain NMDA receptor agonism. In the absence of uncaged glutamate, this manipulation increased synaptic noise but did not significantly affect the average somatic membrane potential (Fig. 5a; $V_{\text{rest}} = -78.9 \pm 0.54$ (control), $-77.9 \pm 0.44$ mV (+NMDA), n=5, p>0.05, Mann-Whitney rank sum test). Moreover, this manipulation in and of itself did not render SPNs more responsive to somatic current injection (Fig. 5b). However, it did enhance the response to stimulation of distal synapses. The duration of up-states triggered by distal glutamate uncaging was significantly increased (Fig. 5c). In contrast, the amplitude of the somatic up-state was not significantly altered ($-66.5 \pm 2.9$ (control), $-57.9 \pm 4.5$ mV (+NMDA), n=5, p>0.05, Mann-Whitney rank sum test). Because bath application of NMDA could induce sustained cortical activity that might contribute to lengthening SPN up-states (Supplementary Fig. 7), these experiments were repeated in the presence of TTX (1 μM) to prevent this activity from being propagated to the striatum. The addition of TTX did not prevent bath application of NMDA from increasing up-state duration (Fig. 5d). This outcome also excludes the potential involvement of striatal interneurons and argues that increased opening of NMDA receptors was responsible for the lengthening. To provide an alternative test of this hypothesis, glutamate was repetitively uncaged at spines on distal dendrites to see if up-states could be extended. Indeed, repetitive stimulation readily extended up-state duration (Fig. 5e), suggesting that up-state duration was controlled by the time course of synaptic input and NMDA receptor activation, in agreement with previous work 5-7.

The data presented to this point suggests that distal dendrites are capable of regenerative activity. Because these dendritic regions are not accessible to direct interrogation and manipulation, a NEURON simulation was used to better understand the phenomenon. In the model, sequential activation of ten glutamatergic synapses along a stretch of distal dendrite generated a local plateau potential (Fig. 6a,b). Moreover, the dendritic plateau depolarized the somatic membrane in a way that resembled that seen experimentally (Fig. 6b). Removal of either NMDA receptors or T- or R-type Ca$^{2+}$ channels reduced the duration of the plateau potential in the dendrites and the up-state in the soma (Fig. 6c–e), as seen experimentally. Interestingly, the simulations suggested that the sub-cellular positioning of Ca$^{2+}$ channels was important to plateau generation: placement of T-type channels in the dendritic shafts and R-type channels in spines was critical to the state transitions (Fig. 6d–e). Closer examination of spine and shaft voltage trajectories at the site of up-state initiation provided an explanation. In spine heads, the depolarization produced by synaptic stimulation was much greater than in the shaft, reaching levels needed for opening the high-threshold, R-type channels. Shaft placement of low-threshold, T-type channels provided a means of boosting the depolarization created by currents generated in spines.

Given the close correspondence between the model and experimental observations, we used the model to address why dendritic plateaus were restricted to distal dendrites. One contributing factor appeared to be the distal distribution of T-type Ca$^{2+}$ channels. But this alone was not enough (data not shown). Another factor could be that the high input...
impedance of fine distal dendrites allows synaptic input to produce the stronger local 
depolarization needed for a regenerative event. SPN dendrites taper and spine density 
decreases with distance from the soma; these features created a distance independence to 
uEPSPs in the dendritic regions examined here, consistent with the view that rising input 
impedance could be a factor (Supplemental Fig. 3). To directly test the hypothesis, a non-
selective ‘leak’ conductance was placed in tertiary dendrites and the effect on the response 
to synaptic stimulation mapped. As the leak increased and input impedance fell, the local 
response lost its regenerative features and the somatic membrane potential shift became 
smaller and shorter (Fig. 6f).

Dopamine and adenosine receptors modulated up-states

Dopamine is a key modulator of SPN intrinsic excitability and corticostriatal glutamatergic 
synaptic transmission. As outlined above, the two principal types of SPNs are 
distinguished by their expression of D₁ and D₂ dopamine receptors. Postsynaptic 
NMDA receptors are major targets of these receptors. To determine if D₁ dopamine 
receptor signaling could affect up-states, glutamate was uncaged on distal spines of D₁ 
receptor-expressing dSPNs and the D₁ receptor agonist ±-6-chloro-PB hydrobromide (6-
CPB; 5 μM) applied. 6-CPB consistently lengthened up-states evoked in these SPNs (Fig. 7a). It was recently reported that D₂ receptor signaling decreased NMDA receptor opening 
in dendritic spines of iSPNs. In agreement with this report, in D₂ receptor-expressing 
iSPNs the D₂ receptor agonist quinpirole (20 μM) consistently shortened up-states evoked 
by distal glutamate uncaging (Fig. 7b). This effect was independent of concomitant A₂a 
receptor stimulation as it was found in the presence of the A₂a receptor antagonist 
SCH58261 (200 nM). Although D₁ receptors are not present in iSPNs, A₂a receptors are, 
which like D₁ receptors are positively coupled to adenylyl cyclase and activation of protein 
kinase A (PKA). Bath application of the A₂a adenosine receptor agonist CGS21680 (200 
nM) consistently lengthened up-state duration in iSPNs, just as the D₁ receptor agonist did 
in dSPNs (Fig. 7c). Application of the A₂a antagonist SCH58261 had no effect on up-state 
duration by itself, suggesting that there was little basal A₂a receptor activity in our 
superfused brain slices (Fig. 7c).

Discussion

Our studies revealed that uncaging glutamate on distal dendritic spines of SPNs, mimicking 
activation of corticostriatal synapses, triggered a local regenerative plateau potential that 
was effectively propagated to the soma, producing a potential change that strongly 
resembled an up-state in vivo. The dendritic plateau potential required the rapid activation of 
only about 10 neighboring synapses. Inward current through both NMDA and voltage-
dependent T- and R-type Ca²⁺ channels contributed to the plateau, shaping its amplitude and duration. In keeping with previous work examining the modulation of SPN NMDA 
receptors, the duration of the somatic up-states was increased by activation of D₁ 
dopamine receptors in dSPNs and shortened by activation of D₂ dopamine receptors in 
iSPNs. Taken together, these results argue that the distal dendrites of SPNs are capable of 
sustaining regenerative plateau potentials that depend upon the release of glutamate and 
engagement of NMDA receptors. The existence of distally generated regenerative events not

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only reconciles the stereotyped features of state transitions in SPNs with the available physiological literature, it fundamentally shifts our model of how SPNs recognize patterns of cortical activity and use this recognition to gate action selection.

Why distal dendrites?

Synaptic stimulation of distal dendrites was much more effective in evoking dendritic plateau potentials and somatic up-states than stimulation of proximal dendrites. This difference was not attributable to a higher density of NMDA receptors at distal synapses. Rather, two other factors appeared to be important. One was the fine caliber and resulting high input impedance of distal dendritic shafts. Because these fine dendrites restrict ion flow away from the site of stimulation, more of the current flowing through synaptically located AMPA receptors is available to discharge the membrane capacitance and depolarize the neighboring membrane, leading to more effective recruitment of NMDA receptors that become unblocked by Mg\(^{2+}\) at depolarized potentials. Simulations of the SPN plateau potentials corroborated this inference. A similar scenario has been described in the fine basal and oblique dendrites of pyramidal neurons where NMDA receptor dependent spikes are seen. Because tertiary SPN dendritic diameter tapers gradually, the capacity to trigger plateau potentials should also increase gradually with distance from the soma, if this factor dominates the process.

Another factor that might contribute to the distal generation of plateau potentials was the investment of these dendrites with T-type Ca\(^{2+}\) channels. SPNs robustly express Cav3.2 and Cav3.3 mRNAs that code for a slowing gating, pore-forming subunit of the T-type Ca\(^{2+}\) channel. These channels are not present to any significant extent in the soma and proximal dendritic membrane of SPNs, but are present in more distal dendrites and spines. The participation of T-type channels in dendritic plateaus was inferred from the ability of the T-/L-type blocker mibefradil to shorten somatic up-states and the failure of the selective L-type antagonists isradipine and calciseptine to do the same. Simulations provided corroboration for these results showing that T-type Ca\(^{2+}\) channels in dendritic shafts boosted synaptically generated depolarization. A similar situation has recently been described in thalamic reticular nucleus neurons. These simulations also suggested the placement of R-type Ca\(^{2+}\) channels in the spine head, rather than the dendritic shaft, was an important boosting mechanism helping to link synaptic depolarization to dendritic plateau potentials.

Are these ‘real’ up-states?

Although our studies show that temporally convergent glutamatergic input to distal dendrites can evoke somatic voltage changes that closely resemble up-states seen in vivo, it was not proven that this is how they are generated in vivo. Nevertheless, there is substantial circumstantial support for this proposition. First, SPN up-states seen in anesthetized rodents depend upon activation of corticostriatal synapses and NMDA receptors, as described here. Second, in this model, the duration of the cortical synaptic barrage dictates the duration of the up-state in vivo, as here. The only discrepancy is the inferred precision of this dependence. In our model, up-state duration is dependent upon locally induced regenerative events created by activation of NMDA receptors, rather than an ongoing synaptic barrage that holds the membrane potential up. Thus, the depolarized plateau potential can outlast the...
synaptic input by hundreds of milliseconds, depending upon glutamate clearance rates and NMDA receptor gating kinetics. A third reason for drawing a connection is that SPN up-states in vivo, as inferred from the duration of episodic spiking, are lengthened by D₁ dopamine receptor activation and shortened by D₂ dopamine receptor activation, as were up-states described here.

The most substantial discrepancy between our model and the prevailing one is the number of synaptic inputs required to drive an up-state. Rather than requiring hundreds of temporally convergent inputs to drive an up-state and hundreds more to sustain it, our data suggests that an order of magnitude fewer glutamatergic inputs are necessary because the distal dendrites of SPNs, like dendrites found in a wide variety of other neurons, are active and capable of regenerative activity. The restriction of the regenerative activity initiation to distal dendrites during synaptic stimulation explains why previous studies have failed to see evidence for it using somatic electrodes. When taken together with the stimulation densities used to evoke up-states, the high spatial convergence (approximately 20 μm of a distal dendrite) described in this study suggests a re-envisioned “synchrony threshold” for up-state induction in SPNs. Fewer highly coupled corticostriatal inputs (as mimicked by adjusting the stimulation strength to induce somatic EPSPs greater than 1 mV) are needed (range = 7–45, median = 25) to induce up-states than weakly coupled inputs (range = 16–64, median = 57). Taking coupling strength into account, and considering a corticostriatal release probability of 0.42, we calculate that about 59 (17–107) strongly coupled or 135 (38–152) weakly coupled synaptic units are required to induce an up-state. Presuming active corticostriatal pyramidal neurons to fire bursts of 4–5 spikes at up to 50 Hz in vivo, it could take the synchronous firing of only 12–15 corticostriatal pyramidal neurons that are strongly coupled to a SPN to induce an up-state. At the other extreme it may take considerably more cortical pyramidal neurons to fire synchronously (27–34) if they are weakly coupled to the SPN.

Given that there are about 30 glutamatergic cortical synapses within the requisite 20 μm stretch of distal dendrite, this convergence requirement does not seem difficult to reach. This number is likely an overestimate for at least two reasons. First, corticostriatal inputs show marked paired pulse inhibition, which will in effect increase the weight of subsequent spikes in a burst. Second, corticostriatal pyramidal neurons are not quiescent in vivo, which will be reflected in the SPN resting membrane potential, and potentially place the SPN in an environment even more conducive to up-state generation. Indeed, our simulations show that mimicking disperse depolarizing inputs to a SPN (by depolarizing the soma ~4 mV) lowers the number of glutamatergic inputs required to induce an up-state (Supplementary Fig. 8).

The full partnership of the post-synaptic membrane in up-state generation and the reliance upon a small number of spatially convergent inputs also helps to explain why residence time in the up-state increases in SPNs with cortical desynchronization and wakefulness. If the convergence requirement for up-state generation is very high, cortical desynchronization should diminish up-state transitions. However, if distal dendrites are active, cortical synchrony is not required for up-state generation. Rather, up-states could depend upon activation of neighboring corticostriatal synapses that represent meaningful patterns of
cortical activity. With wakefulness, the frequency of such meaningful cortical activity should increase, potentially resulting in a sustained up-state, as observed experimentally.

**Implications for pattern recognition and action selection**

The existence of active dendritic regions dramatically increases the pattern recognition capacity of neurons. This increased capacity is achieved by local interactions that confer meaning on synaptic location, as well as synaptic strength. Based upon a clusteron learning rule, the repeated co-activation of neighboring synapses will lead to their stabilization or strengthening as a group, providing local control of plasticity. In SPNs, longer loop feedback also appears to be provided by dopamine, as both D₁ and D₂ receptors modulated the duration of local plateaus in a way consistent with their global effects on synaptic plasticity. That said, this inferred form of local plasticity in distal SPN dendrites sharply contrasts with spike-timing dependent plasticity (STDP), which depends upon the temporal conjunction of local synaptic activity and a ‘global’ somatic spike that back-propagates into the dendrite. Because of this dependence, STDP must normally be limited to proximal dendrites capable of supporting bAP invasion. The fact that the loss of a bAP signal along an SPN dendrite is paralleled by the growth in local regenerative capacity suggests that these two forms of synaptic plasticity are, for the most part, mutually exclusive.

The ability of a small number of distal synapses to induce up-state transitions in SPNs also provides a new framework for understanding what they mean to striatal information processing. Rather than being a consequence of massed cortical activity, up-states could reflect the occurrence of a specific pattern of cortical activity that has been repeated frequently or has been paired with a psychologically meaningful event that altered striatal dopamine release. It is tempting to speculate that these ‘meaningful’ cortical patterns could represent a context that is appropriate for a particular action and that their occurrence triggers an up-state transition in a population of SPNs, making them available or responsive to some other synaptic signal that could evoke spikes. The proposition that inputs triggering spikes and those inducing up-states are different has compelling experimental support. In this scenario, that ‘other’ input could represent possible actions, with those that have been shaped by association with environmental contingencies being the strongest and the most likely to evoke spikes. Linking possible actions to the timing of spikes (rather than up-states) would endow activity in the striatal network with a temporal precision widely perceived as being important for movement control. Proximal synapses that have been shaped by processes like STDP would be the most likely to serve this purpose. In keeping with previous conjectures on this point, our model shows that during an up-state, the effect of a single proximal synapse is significantly enhanced as the electrotonic distance shrinks with closure of dendritic Kir2 K⁺ channels (Supplementary Fig. 9).

If this model of cortical control of SPN activity is correct, it suggests that proximal and distal SPN synapses arise from fundamentally distinct cortical circuits – a conjecture that is not at variance with the anatomy of the corticostriatal projection, but one that will be difficult to test given the lack of a layered striatal cytoarchitecture. One of the few examples of dendritically segregated innervation of SPNs is that of inhibitory circuits, specifically GABAergic interneurons: parvalbumin containing FS interneurons predominantly synapse with...
on SPN soma and proximal dendrites while somatostatin-containing PLTS interneurons predominantly synapse on more distal dendrites. Both interneurons form feed forward corticostriatal circuits. The differences in subcellular targeting of these inputs might have functional consequences. For example, somatic GABAergic input could control the timing of axon initial segment spikes, whereas distal dendritic GABAergic input, by creating a localized shunt that decreases input resistance, could prevent up-state generation by one cortical ensemble, while leaving the capacity of another ensemble to do so intact. GABAergic modulation of state transitions need not be restricted to inputs from interneurons however, as GABAergic SPN axonal collaterals have previously been implicated in playing such a role.

In summary, our studies have revealed that the distal dendrites of SPNs are capable of regenerative activity that produces somatic potential changes resembling up-states in vivo. These regenerative dendritic events were dependent upon the active co-activation of NMDA receptors and voltage-dependent Ca$^{2+}$ channels in small caliber dendrites having high local input impedance. The presence of active dendrites increases the pattern recognition capacity of SPNs. Moreover, it suggests that the associative functions of SPNs – linking context and appropriate action selection – are performed by the integration of two operationally distinct dendritic regions.

**Methods**

**Brain slice preparation**

Para-saggital brain slices (275 μm) were obtained from 21–30 day old BAC D$_1$ or BAC D$_2$ transgenic mice following procedures approved by the Northwestern University Animal Care and Use Committee. The mice were anesthetized with a mixture of ketamine (50 mg kg$^{-1}$) and xylazine (4.5 mg kg$^{-1}$) and perfused transcardially with 5–10 ml ice-cold artificial CSF (aCSF) containing (in mM): 124 NaCl, 3 KCl, 1 CaCl$_2$, 1.5 MgCl$_2$, 26 NaHCO$_3$, 1 NaH$_2$PO$_4$, and 16.66 glucose, continuously bubbled with carbogen (95% O$_2$ and 5% CO$_2$). The slices were then transferred to a holding chamber where they were incubated in aCSF containing (in mM) 2 CaCl$_2$, 1 MgCl$_2$, at 35°C for 60 min, after which they were stored at room temperature until recording.

**Electrophysiology**

Patch pipettes were pulled from thick-walled borosilicate glass on a Sutter P-97 puller and fire-polished before use. Pipette resistance was typically 3–4 MΩ when filled with recording solution. The internal recording solution contained (in mM): 135 KMeSO$_4$, 5 KCl, 10 HEPES, 2 ATP-Mg$^{2+}$, 0.5 GTP-Na$^+$, 5 phosphocreatine-tris; 5 phosphocreatine-Na$^+$, 0.1 spermine, 0.2 Fluo-4 pentapotassium salt and 0.05 Alexa Fluor 568 hydrazide Na$^+$ salt (Invitrogen), pH was adjusted to 7.25 with NaOH and osmolarity to 270–280 mOsm l$^{-1}$. In some cases Fluo 4FF (500 μM) was used in place of Fluo 4, as indicated in the text. Slices were continuously perfused with carbogen-bubbled aCSF. Slices were transferred to a submersion-style recording chamber mounted on an Olympus BX51 upright, fixed-stage microscope. Electrophysiological recordings were obtained with a Multiclamp 700B amplifier. Stimulation and display were obtained as previously described (Day et al., 2008).
using the custom-written shareware package WinFluor (John Dempster, Strathclyde University, Glasgow, Scotland, UK), which automates and synchronizes the two-photon imaging and electrophysiological protocols. The amplifier bridge circuit was adjusted to compensate for serial resistance and continuously monitored during recordings. Recordings were performed at room temperature, unless otherwise noted.

2-photon laser scanning microscopy (2PLSM)

Direct and indirect pathway SPNs were identified by somatic eGFP two-photon excited fluorescence using an Ultima Laser Scanning Microscope system (Prairie Technologies). A DODT contrast detector system was used to provide a bright-field transmission image in registration with the fluorescent images. The green GFP signals (490–560 nm) were acquired using 810 nm excitation (Verdi/Mira laser). SPNs were patched using video microscopy with a Hitachi CCD camera and an Olympus 60X/0.9 NA lens. Alexa 568 fluorescence was used for visualization of cell bodies, dendrites, and spines. Following patch rupture, the internal solution was allowed to equilibrate for 15–20 minutes before imaging. Maximum projection images of the soma and dendrites were acquired with 0.36 μm² pixels with 10 μs pixel dwell time; ~80 images are taken with 1 μm focal steps.

Ca²⁺ imaging

Drugs were either bath applied by dissolving them in the external aCSF (for experiments involving NMDA and glycine) or superfused via the perfusion pump used to apply MNI-glutamate described below. In the latter instances drugs were dissolved in HEPES-based aCSF containing (in mM): 124 NaCl, 3 KCl, 2 CaCl₂, 1 MgCl₂, 26 NaHCO₃, 1 NaH₂PO₄, 26 HEPES and 16.66 glucose. Dendritic changes in Ca²⁺ were measured using Fluo-4 as previously described. Ca²⁺ transient were expressed as ΔF/Fo. Green fluorescent line scan signals were acquired at 6 ms per line and 512 pixels per line with 0.08 μm pixels and 10 μs pixel dwell time. The laser-scanned images were acquired with 810 nm light pulsed at 90 MHz (~250 fs pulse duration). Power attenuation was achieved with two Pockels cells electro-optic modulators (models 350–80 and 350–50, Con Optics, Danbury, CT). The two cells were aligned in series to provide an enhanced modulation range for fine control of the excitation dose (0.1% steps over four decades). The line scan was started 200 ms before the stimulation protocol and continued 4 seconds after the stimulation to obtain the background fluorescence and to record the decay of the optical signal after stimulation. To reduce photo-damage and photo-bleaching, the laser was fully attenuated using the second Pockels cell at all times during the scan except for the period directly flanking the evoked state transition.

2PLU

2P uncaging of MNI-glutamate was performed simultaneously with Ca²⁺ imaging using a Chameleon-XR laser system (Coherent Laser Group, Santa Clara, CA). MNI-glutamate (Tocris Cookson, Ellisville, MO) was superfused (0.4 ml/hr) at 5 mM using a system of syringe pumps (WPI, Sarasota, FL) and a multi-barreled perfusion manifold fitted with a small-volume mixing tip that allowed rapid switching between solutions (Cell MicroControls, Norfolk, VA). MNI-glutamate was uncaged using 1ms pulses of 720 nm light independently controlled by a third Pockels cell modulator (model 302, Con Optics, Danbury, CT). Experiments involved uncaging on a single spine to evoke a somatic EPSP
measuring 1–4 mV. The uncaging pulses were 1 ms in duration and typically about 15 mW in strength measured at the sample plane. Photolysis power was adjusted to closely mimic spontaneously occurring EPSPs and tuned to achieve the predetermined somatic EPSP amplitude. We opted to use this strategy of tuning laser power, as opposed to relying on photobleaching, because of our method of MNI glutamate delivery. As this study required same-cell pharmacological manipulations, MNI glutamate was delivered via a superfusion pump rather than a recirculating pump, introducing chemical penetration as an addition variable, making photobleaching an inappropriate measure of total glutamate uncaging. State transitions were evoked by stimulating approximately 10 neighboring coplanar spines in rapid succession (500 Hz), with each spine stimulated 1–4 times. The distance from soma was measured by a straight line drawn from the site of stimulation to the origin of the dendrite at the soma. Simultaneous photolysis and line-scan images were made from shafts/spines. As described above, the custom-written software package WinFluor automated and synchronized the Ca$^{2+}$ imaging with the electrophysiological stimulation and the photolysis.

**Neuron modeling**

A SPN was modeled using NEURON version 6.0. The presented model is based upon anatomical reconstructions of SPNs and similar to one previously published by our group containing 6 primary dendrites having primary, secondary and tertiary branch points. In addition, 1396 spines were added to 4 tertiary dendrites. The other dendrites were modeled in a way that mimicked the presence of spines, but computationally simplified the model. Soma dendrites were populated with ion channels known to contribute to SPN excitability, with the exception of Nav1 Na$^{+}$ channels, which were shown experimentally in the present study not to be participants in up-state generation; this omission simplified the simulations by eliminating the complication of somatic spikes. Spines were endowed with L-type, Cav3.2 T-type and R-type voltage gated Ca$^{2+}$ channels, K$_{Leak}$ channels and Kir2 potassium channels. Presynaptic glutamate releasing terminals were added to 10 neighboring spines on a single tertiary dendrite (mimicking a distal stimulation site) to simulate a corticostriatal synapse. These 10 corresponding spines were further endowed with AMPA and NMDA receptors. Tertiary dendrite and spine Ra was adjusted to 125 $\Omega$cm. Cav3.2 conductances matching those in spines were added to tertiary dendrites. NMDA/AMPA/Cav3.2/R-type channel conductances were tuned to give somatic voltage responses similar to those observed experimentally. Kv1 and KCNQ potassium conductances were increased in the initial axon segment to favor somatic state transition kinetics similar to those observed experimentally.

**Statistical analysis**

Differences in state transition duration and Ca$^{2+}$ transients were examined using the Mann-Whitney U nonparametric test of significance and the Wilcoxin signed rank test. Differences were considered statistically significant if p<0.05.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.
Acknowledgments

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Figure 1.
State transitions can be triggered by local stimulation of distal dendrites. (A) Low (left) and high (right) magnification maximum intensity projections (MIPs) of a dSPN filled with Alexa Fluor 568. Spines were stimulated with 1 ms uncaging pulses at 500 Hz, and are numbered in the order stimulated. (B) Somatic voltage recording of the cell shown in (A) in response to glutamate uncaging of a single spine (red), or all labeled spines in rapid succession (black). Red and black shaded regions indicate timing of uncaging pulses. An all points histogram is shown in the inset to the upper right. State membrane potentials are shown above as box plots (n=56 cells). For all box plots, lines indicate median, boxes
outline upper and lower quartiles and whiskers are the upper and lower 10% values. Shaded green boxes indicate state membrane potentials reported in a previous in vivo study and are shown for reference. (C–D) State transitions were evoked by uncaging at distal (black) dendritic spines; the same protocol was then used to excite a proximal dendrite (red). Somatic voltage traces aligned to the last uncaging stimulus and normalized show up-states are readily induced by distal but not proximal activation. Box plots indicate time between end of uncaging and 25% voltage fall (n=8, p=0.002, Mann-Whitney rank sum test). (E) Example spine Ca\textsuperscript{2+} transient (top) and corresponding somatic membrane potential (bottom) of an example up state in a dSPN. Ca\textsuperscript{2+} transient is measured using the low affinity Ca\textsuperscript{2+} indicator Fluo 4FF. (F) Experiment presented in 1c repeated using the Ca\textsuperscript{2+} indicator Fluo 4FF (n=9 cells; *p<0.01, Mann-Whitney rank sum test). Up-state duration is plotted against the corresponding local distal dendritic Ca\textsuperscript{2+} transient (measured at the dendritic shaft around which spines were stimulated).
Figure 2.
State transitions generated by proximal and distal dendritic glutamate uncaging are independent of the somatic EPSP size and recording internal used. (A) Example of a state transition induced in a dSPN: recording was made using a K-gluconate based internal: (in mM) 115 K-gluconate, 20 KCl, 10 Na-phosphocreatine, 10 HEPES, 2 Mg-ATP, and 0.3 Na-GTP<sup>47</sup> and Fluo 4FF; laser power was tuned to evoke small (463 μV) somatic EPSPs from a 0.5ms stimulation of a single distal dendritic spine (inset). The corresponding all points histogram for this cell is shown to the right. (B) Once a state transition is achieved, the up-state duration is not correlated with either the somatic EPSP induced by stimulation of a single distal dendritic spine or the recording internal solution used (n=6 cells; internal = KMeSO<sub>4</sub> or K-gluconate with Fluo 4 or Fluo 4FF). (C) Though requiring a higher stimulation density, up-states induced using small amplitude (463–774 μV; n=6 cells) somatic EPSP are similar to those induced by supra-mV EPSP (i.e. figure 1). (D) In one case a modest state transition was produced by the repetitive stimulation of a single distal spine

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(16 stimuli, 0.5 ms duration, 500 Hz). Average somatic EPSP evoked by a single stimulus and the corresponding all points histogram are shown to the right.
Figure 3.
State transition generation is similar in dSPN and iSPNs. (A–B) State transitions were generated by glutamate uncaging on distal dendrites of dSPN (left) and iSPNs (right). High magnification MIP images showing the location of glutamate uncaging (A) and the resulting somatic voltage traces (B). Insets show all point histograms of membrane potentials for each trace. (C) Somatic membrane potentials of ‘up’ and ‘down’ states in dSPNs (n=56 cells) and iSPNs (n=18 cells). Data from figure 1b is shown for comparison and highlighted in green in 2c. (D–E) The length of spiny distal dendrite and the stimulation density (number of stimuli per 1 μm dendritic length) required to evoke state transitions is similar in dSPNs (n=39) and iSPNs (n=11). (F) No obvious differences were observed in the relationship between stimulus density and up-state duration in D1 (n=39) or D2 (n=11) SPNs.
Figure 4.
State transition generation requires NMDA receptors and voltage gated Ca\(^{2+}\) channels. (A) State transitions were generated in dSPNs by glutamate uncaging on distal dendritic spines, and then the response to the same uncaging protocol was measured in the presence of the NMDA receptor antagonist AP5 (100 μM). Shaded region indicates timing of glutamate uncaging. Voltage traces are normalized to maximum amplitude and overlain. Traces shown to the right are aligned to the end of the last uncaging pulse and normalized to the maximum amplitude of the state transition, as shown in figure 1d. Box plots indicate time between end of uncaging stimulation and 25% voltage fall. AP5 significantly reduced state transition duration, abolishing it in most cases (n=5, p<0.05 Mann-Whitney rank sum test). (B–D) State transitions were generated in dSPNs by glutamate uncaging on distal dendritic spines, and then the response to the same uncaging protocol was measured in the presence of (B) NiCl\(_2\) (50 μM; n=8), (C) Mibefradil (20 μM; n=8) or (D) SNX-482 (0.3 μM; n=6). Box plots indicate time between end of uncaging stimulation and 25% voltage fall. All three agents significantly reduce the somatic state transition duration (* p<0.05 Mann-Whitney rank sum test).
Figure 5.
NMDA acts at distal dendritic synapses to enable state transition generation. (A) Somatic voltage recordings of spontaneous depolarizations in a dSPN before and after application of 10 μM NMDA + 10 μM glycine + 10 μM gabazine (SR95531) (“NMDA cocktail”, top) shows an increase in excitatory inputs, which is prevented by 1 μM TTX (bottom). (B) Somatic voltage of a dSPN in response to suprathreshold current injection in the presence and absence of 10 μM NMDA/glycine/gabazine. No regenerative changes in somatic voltage were observed, implying both the location of up-state duration and the NMDA receptors involved in their maintenance are located distally. (C) State transitions were evoked by glutamate uncaging on distal dendritic spines of dSPNs, then again in the presence of 10 μM NMDA/glycine/gabazine. Somatic up-state duration was significantly enhanced (n=6, p<0.05 Signed Rank test). Shaded region indicates timing of glutamate uncaging. Traces are normalized to the maximum amplitude. Box plots indicate time between end of uncaging and 25% voltage fall. (D) The increase in up-state duration in dSPNs caused by 10 μM NMDA/glycine/gabazine is not occluded by 1 μM TTX, suggesting the action of NMDA is directly on distal SPN dendrites and does not rely upon action potential dependent glutamate release (n=4). (E) A state transition was induced by stimulation of 10 distal dendritic spines (twice each) in a dSPN. The up-state could be maintained by supplemental stimulation of these same spines (once each) every 50 ms. Uncaging pulses are indicated by shaded regions.
Figure 6.
Neuron modeling supports synergistic action of NMDA receptors and T/R Ca\(^{2+}\) channels in up-state generation and maintenance. (A) NEURON model of a SPN containing 6 primary dendrites, 12 secondary dendrites and 24 tertiary dendrites. 4 of the tertiary dendrites are invested with spines and indicated by thick blue lines (s3°). The other 20 tertiary dendrites do not contain localized spines (ns3°). All tertiary dendrites have diameters tapered from 2.5–0.3 µm. AMPA and NMDA receptors are located in spines only; Cav3.2 channels are located in spines and tertiary dendrites, except where noted in (E); R-type channels are located only in tertiary spines, except where noted in (D). (B) Simulations were run using the same stimulation protocol described experimentally, and the resulting membrane potential is shown for the first stimulated spine, the stimulated spiny 3° dendrite, the corresponding parent 2° dendrites of a stimulated and nonstimulated 3° dendrite, and the soma. (C–E) Simulations were run in ‘control’ conditions and in the absence of (C) functional NMDA receptors, (D) R-type channels or (E) Cav3.2 T-type channels. Voltage traces are of the first stimulated spine (top), the activated tertiary dendrite (middle) and soma (bottom). (F) Simulations were run using the same protocol, gradually increasing the spine ‘leak’ conductance from 0 to \(10^{-3}\) S/cm\(^2\). Increasing the leak conductance, effectively...
decreasing the input impedance, reduces the membrane potential amplitude and plateau duration in both the stimulated spine (top), 3° dendrite (middle) and soma (bottom). Shaded regions indicate timing of stimulation.
Figure 7.
Up-state duration is modulated by dopamine and adenosine receptor signaling. (A) Example normalized traces of an up-state induced by stimulation of distal dendritic spines in a dSPN before and after application of the D<sub>1</sub> agonist 6-CPB (5 μM). Action potentials are truncated to better display state transitions. Shaded region indicates timing of glutamate uncaging. Box plots to the right show time between end of uncaging stimulation and 25% voltage fall. 6-CPB increased up-state duration in all dSPNs examined (n=8; p=0.008 Signed rank test). Before and after data points from the same cells are connected with lines. (B) Example normalized traces of an up-state induced by stimulation of distal dendritic spines in an iSPN before and after application of the D<sub>2</sub> agonist quinpirole (20 μM) and A2a receptor antagonist SCH58261 (200 nM). Shaded region indicates timing of glutamate uncaging. Box
plots (time from end of uncaging stimulation to 25% voltage fall) to the right show that quinpirole/SCH58261 reduces up-state duration in iSPNs (n=6; p<0.05 Signed Rank Test). Before and after data points from the same cells are connected with lines. (C) Box plots showing the percent change in up-state duration in D1 and D2 SPNs by 6-CPB (from (A)), CGS21680 (200 nM; n=6, p<0.05 Signed rank test), SCH58261 (n=5, signed rank test) and quinpirole + SCH58261 (from (B)).