Striatal origin of the pathologic beta oscillations in Parkinson's disease
Enhanced oscillations at beta frequencies (8–30 Hz) are a signature neural dynamic pathology in the basal ganglia and cortex of Parkinson’s disease patients (1). Improvement of bradykinesia correlates with a decrease in the enhanced beta frequency oscillations in the subthalamic nucleus (STN) and cortex of Parkinson’s disease patients (2). However, the function of the beta oscillations in parkinsonian pathology remains elusive, and the source of these oscillations is unknown. Two predominating theories exist about the origin of these enhanced beta rhythms. One hypothesis proposes that oscillations arise from the interaction of the STN and the external segment of the globus pallidus (GPi); this is known as the STN/GPi pacemaker hypothesis (3). Evidence for the STN/GPi pacemaker hypothesis comes from a study showing that GPi and STN are able to generate synchronized oscillatory bursting activity in the range of 0.4–1.8 Hz in organotypic cultures of cortex-striatum-STN-GPi (4). A second theory of beta generation in Parkinson’s disease entails cortical patterning of the STN (3). This theory derives partly from studies on anesthetized rats showing that, with dopamine depletion, oscillatory spiking activity in STN and GPi is correlated with and largely dependent on cortical slow-wave (~1 Hz) activity (5). For the cortical patterning, but not the STN/GPi hypothesis, there is experimental evidence for intrinsic production of the beta rhythm (6).

Degeneration of dopaminergic neurons that project to the striatum is a hallmark of Parkinson’s disease pathology (7), but the striatum has been largely ignored as a possible source of beta frequency rhythms in Parkinsson’s disease. There are several reasons for ignoring the striatum. The striatum consists of an almost entirely inhibitory network of cells (99.7% are GABAergic and 0.3% are cholinergic in the rat neostriatum) (8). Furthermore, the predominant striatal cell type, the medium spiny neuron (MSN), rarely spikes (average rate of 1.1 ± 0.18 Hz in freely moving rats) (9). It is not evident that this largely inactive and inhibitory network can generate any independent activity, much less rhythmic activity at a relatively high frequency (8–30 Hz). However, mathematical modeling informs us that inhibition can create neuronal excitation under certain conditions. Specifically, the activation or inactivation of slow currents can lead to the phenomenon of postinhibitory rebound spiking. Moreover, inhibitory networks of such neurons can produce rhythmic activity (10, 11). The frequency of the rhythm largely depends on the time constant of the slow current. At least one slow current is known to have a time constant of decay that allows postinhibitory rebound spiking to occur with a lag appropriate for the formation of beta frequency spiking: the M-current (10). The M-current is a nonactivating potassium current, and GABAa inhibition can temporarily reduce this current, leaving the neuronal membrane in a more excited state (12). The MSNs of the striatum have an M-current and receive inhibition from each other through GABAa synapses. Therefore, the MSNs of the striatum contain both the cellular and the network properties to support the generation of beta frequency oscillations.

We construct mathematical models of networks of MSNs and find that such networks are capable of generating robust beta oscillations. Perturbations that enhance the interaction between MSN M-current and GABAa current increase the power of the population beta rhythm. An interesting prediction is that increases in either MSN background excitation (I_{sec}) or MSN GABAergic inhibition will increase the power of beta oscillations in the model MSN network. Because increasing MSN excitation necessarily increases MSN to MSN inhibition, these seemingly opposite effects, in fact, work synergistically to create beta frequency oscillations in our model MSN networks.

Striatopallidal MSNs have high levels of dopamine D2 receptors and increase their excitability in response to loss of dopamine (13). Furthermore, loss of dopamine has been shown to increase beta oscillations in striatum (14). This finding is consistent with our model results, which predict enhanced beta oscillations in the subnetwork of striatopallidal MSNs after loss of striatal dopamine. MSN excitability can also be increased by other neuromodulators, most notably by the action of acetylcholine (ACH) on M1 receptors (13). This is important in the context of Parkinson’s disease, because dopamine modulates ACh levels. Dopamine tonically inhibits ACh release in the striatum under normal physiological conditions by its action on D2 receptors (15). Moreover, antagonists of either D1 or D2 receptors as well as loss of striatal dopamine in the 6-hydroxydopamine (6-OHDA) rat, an animal model of Parkinson’s disease, result in increased striatal levels of ACh (16).

As a first test of our model, we inject the muscarinic acetylcholine receptor agonist carbachol into the striatum of normal awake mice. As predicted, cholinergic agonists in the striatum induce enhanced beta frequency oscillations in the striatal local field potential (LFP). Neither striatal nor cortical LFPs show
increased beta oscillations after infusion of carbachol into the cortex, showing that striatal LFP beta is not the result of diffusion of carbachol into the neighboring cortex. Our combined computational and experimental results suggest that increased MSN excitation, which produces increased MSN to MSN inhibitory interactions, is a potential source of the enhanced beta rhythms in Parkinson’s disease. Our modeling results also suggest that loss of dopamine and increased striatal ACh work in parallel to amplify beta oscillations in striatal networks in Parkinson’s disease.

**Results from Mathematical Models of Normal and Parkinsonian Networks of MSNs**

**Networks of MSNs Alone Produce Beta Oscillations.** A raster plot of our 100-neuron striatal model (see Computational Methods) using weak all-to-all connections displays no readily observable pattern of activity among MSNs (Fig. 1A). However, analysis of the model LFP reveals a distinct peak in the low beta frequency range centered around 12.1 ± 0.7 Hz (Fig. 1B). The beta frequency oscillations wax and wane over time (Fig. 1C). The spiking of individual MSNs is irregular (Fig. 1D). The average spiking rate of the neurons in this network is 0.96 ± 0.03 Hz, consistent with the low average MSN spiking rate in vivo (9). The trace of the model LFP also contains waxing and waning beta frequency oscillations (Fig. 1E).

**Beta Power and MSN Spiking Frequency Increase in Parkinson’s Disease Model Striatum.** We simulate the loss of dopamine in the parkinsonian striatum indirectly through ACh-induced reduction of the M-current (see Computational Methods). Decreasing the maximal M-current conductance by 7.7% has pronounced effects on the striatal network dynamics [the percent decrease of the maximal M-current conductance \( g_m \) depends on various parameters; for example, Fig. S1 shows a qualitatively similar model in which the M-current changes 41.2% between the normal and parkinsonian states]. MSN spiking becomes more obviously patterned into a beta frequency population rhythm (Fig. 1F). The LFP power peaks at a higher frequency (17.1 ± 0.32 Hz), and the peak power is higher than in the non-parkinsonian condition (Fig. 1G). This beta oscillation is persistent in contrast to the waxing and waning of the oscillation in the non-parkinsonian condition (Fig. 1H). Individual MSNs show more prominent subthreshold beta frequency oscillations because of the increased number of MSNs participating in the population beta rhythm, which provides individual MSNs with increased beta frequency synaptic input (Fig. 1I). MSNs spike more frequently than in the non-parkinsonian condition, with an average spiking rate of 4.9 ± 0.15 Hz. The animal literature is unclear; some studies show an increase in striatal activity with parkinsonism (9, 17), whereas others show a decrease (18). The model LFP contains a more pronounced and persistent oscillatory component in the beta frequency range (Fig. 1J). We note that the exact frequency range of oscillations found experimentally within the basal ganglia differs among papers; some describe 15–30 Hz oscillations, whereas others extend this range down to 8 Hz (19, 20).

**Properties of the Mathematical Model**

**GABAa-Current and M-Current Conductance Strengths and Background Excitation \( I_{app} \) Modulate LFP Beta Power and Peak Beta Frequency.** Here, we are not looking at normal and parkinsonian states but merely at the change in beta as we change a parameter. The power and frequency of the beta oscillation tend to change nonmonotonically as the GABAa conductance is increased. (Fig. S2 and Fig. S3A and B). Decreasing the M-current conductance or increasing \( I_{app} \) tends to increase both the power and frequency of the beta oscillation (Fig. S3 C–F). In the absence of M-current, the MSN neurons resemble fast-spiking (FS) interneurons. We find that it is possible to obtain beta oscillations in networks of striatal FS interneurons alone but not in a regime supported by the experimental literature (Fig. S4).

**Model Results Are Largely Invariant to Changes in Network Connectivity, Network Size, and Heterogeneity.** Model striatal networks with either nearest neighbor connections or random network connections give qualitatively similar results to the network with all-to-all connections (SI Note 4: Model results are largely invariant to changes in network connectivity, network size, and heterogeneity; Fig. S5A). Increasing the number of MSNs to 400 or introducing heterogeneity by giving individual MSNs different maximal GABAa conductances leads to qualitatively similar behavior to the networks with 100 MSNs and homogeneous maximal GABAa conductances. Heterogeneity also helps us distinguish between the models of MSNs with and without M-current, because

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**Fig. 1.** Beta oscillations emerge in the model striatum under normal conditions and become enhanced under parkinsonian conditions. A–E are taken from the same 7-s simulation under normal conditions, and F–I are from the same 7-s simulation under parkinsonian conditions. (A) Raster plot of 100 reciprocally connected medium spiking neurons (MSNs) under normal, non-parkinsonian conditions. (B) Power spectral density of the model LFP. (C) Spectrogram of the model LFP. (D) Membrane voltage fluctuations from one MSN in the network. (E) Waxing and waning of the model LFP trace under normal conditions. (F) Raster plot of 100 reciprocally connected MSNs under parkinsonian conditions. (G) Power spectral density of the model LFP. (H) Spectrogram of the model LFP. (I) Membrane voltage fluctuations from one MSN in the parkinsonian network. (J) The model LFP trace under parkinsonian conditions.
Experimental Results

Increase in Striatal Cholinergic Activity Leads to Increase in Striatal Beta Frequency Oscillations. We tested our model experimentally by recording the striatal LFP while infusing the cholinergic agonist carbachol into the striatum of normal awake, head-fixed mice. Carbachol (0.5–1 mM and 1–2 μL) infusion directly into the dorsolateral striatum (stereotactic coordinate: AP (anterior posterior) = −0.2 to 0.8, ML (medial lateral) = 2.0–2.5, and DV (dorsal ventral) = 3.0–3.5) induced prominent beta frequency oscillations in simultaneously recorded LFP (Fig. 2A–E). Often, the observed increase in beta power was preceded with a brief decrease in beta power (Fig. 2A). Episodes of prominent increase in beta oscillations often last several hundred milliseconds to several seconds, and they often presented a slight decrease in frequencies from high to low beta during each episode (Fig. 2D and E). Upon striatal carbachol infusion, a tremor-like movement was observed on the contralateral side. We refer to this movement as tremor-like to indicate that the movement has a rhythmic component. We do not assume that these movements represent the parkinsonian tremor, which occurs at a higher frequency. These tremor-like movements often interrupt the enhanced beta oscillation episodes before each movement onset. This observation is consistent with evidence that desynchronization of beta frequency oscillations occurs in both non-Parkinson’s and Parkinson’s disease patients before movement, although the latency to premovement desynchronization is delayed in Parkinson’s disease patients in the absence of L-dopa (21–23).

Of the six mice tested, it took 442 ± 119 ms (mean ± SD) for beta power to increase after infusion onset. Because the infusion site is typically positioned 300–700 μm away from the recording electrode, diffusion of carbachol from the infusion site to the recording electrode may account for a significant portion of this observed delay. After the increase in beta power, beta power often dropped below the baseline and stayed low for an extended period even at the end of the recording session, typically 1 h after the end of infusion (Fig. 2F).

To eliminate the possibility that the observed LFP change was caused by the pressure induced by drug infusion, which may activate mecanosensitive mechanisms in the striatum, we infused carbachol at a lower concentration (0.1–0.2 mM, n = 5; three mice infused with 1–2 μL and two mice infused with 4–5 μL). Infusion of low-concentration carbachol failed to induce any change in the LFP, even with up to 5 μL infusion. Beta power remained stable throughout the entire infusion period (Fig. 2G). Furthermore, we did not observe any reduction in beta power at the end of these recording sessions. Thus, the reduction in beta oscillations following augmentation as induced with high-concentration carbachol infusion cannot be explained by prolonged recording, confirming that the observed beta power reduction following beta increase is due to neural network mechanisms (Fig. 2F). In addition, in mice infused with low-concentration carbachol, no prominent tremor-like movement was observed.

Striatum Contains Sufficient Neural Network Components to Initiate Enhanced Beta Oscillations upon Cholinergic Activation. To examine whether the observed increase in beta oscillations is initiated in the striatum but not due to the diffusion of carbachol into the adjacent cortical regions, we infused carbachol (1 mM) into the cortex directly above the dorsolateral striatum where the striatal
infusion sites were located. We did not observe any increase in beta oscillations either locally in the cortex adjacent to the cortical infusion site or in the striatum directly below (Fig. 2 H and I). When advancing the infusion apparatus into the dorsolateral striatum and infusing carbachol (1 mM) in the striatal sites, we reliably induced an increase in beta oscillations in the striatum, confirming that the lack of beta increase on infusion in the cortical areas is due to the local network properties at the cortical infusion sites. These results demonstrate that carbachol diffusion into the cortical regions cannot explain the increase in striatal beta oscillations upon striatal infusion, thus suggesting that striatum is sufficient to induce enhanced beta oscillations.

**Discussion**

Our results indicate that the interaction between the MSN GABAa current and the M-current generates beta frequency oscillations within striatal networks of MSNs. Increased MSN excitation promotes additional cellular-level interaction between these two currents, resulting in increased beta frequency power in the model LFP. Our experimental work verifies that increasing MSN excitability through amplification of striatal cholinergic tone, a condition relevant to Parkinson’s disease, is sufficient to engender robust beta frequency rhythms in the striatum.

Although it is not surprising that a muscarinic agonist may increase MSN spiking rate (13), it is surprising that a muscarinic agonist can induce rhythmic activity in the beta frequency range in striatum. Our mathematical model of striatum predicts that the M-current plays an important role in establishing the beta frequency of the rhythm. The mechanism by which reciprocally connected GABAergic neurons with an M-current generate beta frequency population spiking is described in McCarthy et al. (10). The GABAa current reduces the M-current by bringing the membrane potential closer to the potassium reversal potential. Reduction of the M-current depolarizes the neuron, potentially allowing the neuron to rebound spike. The dynamical interaction involving the slow time constants of decay of each current allows rebound spiking to occur with lags appropriate to the formation of a beta rhythm. An unintuitive consequence of this interaction is that increasing the GABAa conductance can cause a greater reduction of the M-current and provide additional membrane excitation. As a result of increasing GABAa conductance, the neuron may be able to rebound spike in the absence of MSN synchrony; that is, without the necessary neural network components to augment beta oscillations upon excitation of MSNs and that network interactions between MSNs alone, independent of striatal GABAergic interneurons, may be capable of creating beta frequency oscillations. Furthermore, the ability of our model to produce beta and to increase that beta in the parkinsonian state are both largely invariant to the pattern of network connectivity. This is an important result, suggesting that the beta rhythms in striatum may be quite robust to plastic changes in network connectivity.

Because MSN to MSN connections are very weak, their function in the striatal network is unclear (24). Our results here suggest that these weak connections structure the spiking of MSNs so that the latter is more likely to spike at the same phase of the beta cycle. Furthermore, any modulator of the GABAa current will influence the magnitude and peak frequency of the MSN network beta oscillation. Increasing the GABAa conductance increases the peak frequency of the beta oscillation. Because elevated MSN spiking rates increase MSN to MSN GABAergic inhibition, any potentiator of MSN excitability increases the beta frequency oscillation in our model MSN network. Thus, the weak functional synapses between MSNs generate a highly modulable system in which both the power and the frequency of the beta oscillations are regulated by modulators of either MSN excitability or the GABAa current. Although the functional physiological significance of both normal and pathological beta rhythms remains to be elucidated, a high degree of modulation provides this system with the flexibility to adapt to a range of behavioral conditions. Additional study of this system will reveal the extent to which modulators of GABAa and the M-current can create beta rhythms in the MSN (SI Note 8: Other currents in MSNs has more discussion of this topic).

Our models predict increased MSN spiking in the parkinsonian state. Such increased spiking activity of MSNs has been reported in animal models of Parkinson’s disease (9, 27). Our models also predict an increased number of MSNs spiking at the same phase of each cycle of the beta rhythm under parkinsonian conditions. Because MSNs are the only output neurons of the striatum, increasing the number of synchronously spiking MSNs at each cycle of the beta oscillation has important implications concerning the transmission of this rhythmicity to downstream structures. Substantial convergence of input is thought to occur between the striatum and the globus pallidus (GP) because of the much larger number of neurons in striatum compared with the GP (28). Thus, increasing MSN synchrony should lead to stronger GABAergic input onto GP neurons. Because GP neurons can phase reset in response to GABAa inhibitory input (29) and in particular, to transient striatal input (30), the GP neurons receiving the synchronous MSN input may be patterned into the beta frequency oscillation. Additionally, in Parkinson’s disease, D1 and D2 MSNs are thought to decrease and increase their excitability, respectively, in response to loss of dopamine (13, 31). Our model predicts the power of the beta oscillation scales with the level of excitation in the MSN population. Thus, we expect the population of D2 MSNs to increase network beta in the parkinsonian state and the population of D1 MSNs to decrease network beta in the parkinsonian state. Furthermore, because the rate of unilateral connectivity from D1 to D2 MSNs is relatively low (6%) (32), we do not expect significant interaction between these populations. Thus, we expect the mechanism of beta rhythm production that we describe here to be most applicable to the dynamics of neurons that project mainly to GPe (D2 MSNs) in the context of Parkinson’s disease. This is in line with current thinking that suggests that the direct pathway is underactive in Parkinson’s disease and that the indirect pathway is overactive (1).

Our computational model producing striatal beta oscillations is independent of both specific external input and input from intrastriatal FS interneurons. In fact, we expect that striatal FS interneuron input may be diminished in Parkinson’s disease and in our experiments with carbachol, because ACh attenuates FS inhibition of MSNs through presynaptic muscarinic receptors (33). Thus, adding FS interneurons to the model will only affect the results of the normal, non-parkinsonian case. Because we are not trying to replicate all of the LFP patterns of the normal striatum but rather, explain the parkinsonian beta oscillation, we feel that a more complex model of the striatum with FS cells is not necessary. Similarly, we expect glutamatergic input from cortex and thalamus to be reduced, because activation of striatal M1 muscarinic receptors decreases both the probability of glutamate release from presynaptic terminals as well as the potency
of individual glutamatergic synapses onto MSNs (34). However, we expect striatal levels of ACh, which are thought to come mainly from striatal cholinergic interneurons, to be elevated in Parkinson’s disease (15, 16). We note that anticholinergics were the only available pharmacologic treatment for Parkinson’s disease until the introduction of L-dopa in the 1960s (35). Our model predicts that some of their efficacy may reside in their ability to decrease striatal beta oscillations.

The experimental results show additional changes not predicted by the computational models. The model does not predict the lower delta and theta frequency rhythms seen in the absence of carbachol. This may be due to the presence of other striatal cell types such as FS neurons not present in our model. Tremor-like activity is not a prediction of our model; beta oscillations in Parkinson’s disease do not necessarily correlate with tremor (2). Even in the present study, however, the tremor can be viewed as interrupting the beta oscillation. The mouse beta oscillations are interrupted by periods of decreased beta before and after the tremor movement. We attribute this to some in vivo compensatory change in the response to increased beta in the striatum, possibly modulated by thalamic input, as there is evidence that parkinsonian tremor is modulated by the thalamus (36). Also, in the mice, there is a drifting in the frequency of beta, which attains its highest values postmovement, drifts to lower values, and disappears before the tremor movement. Our model provides several potential explanations for frequency drift, including changes to the GABAa or M-current conductances or changes in the background excitation. However, it is unclear which of these mechanisms might be contributing to the frequency drift in the mouse striatum. We also cannot account for the decreases in beta seen before the carbachol-induced rise in beta, which might represent a homeostatic response to the carbachol infusion. However, the decrease in beta that follows the rise in beta may be due to overrecovery of the M-current, in which increases in the M-current follow its suppression (37). Differences between our study and other published studies as well as other models of striatal MSNs are discussed in SI Note 9: Other Related Studies.

Other studies suggest that the enhanced beta rhythms in Parkinson’s disease may arise from the interaction of the STN and the GPe or from the cortical patterning of STN (3). Both theories are supported by experimental studies showing very low frequency oscillations (less than 1.8 Hz) developing in structures relevant to Parkinson’s disease (4, 5). In contrast, our studies on striatum show the emergence of robust beta frequency oscillations in both our computational and experimental results. Computational models of the STN-GPe network show that, under certain conditions, the system exhibits rhythmic oscillations within the frequency range appropriate to Parkinson’s disease (3, 38). Interestingly, the critical condition for this to occur is increased inhibitory input from the striatum; increased patterned inhibitory input is a prediction of our model. Furthermore, the failure of cortical infusions of carbachol to induce beta oscillations is evidence against the idea that increasing gain anywhere within the cortico-basal ganglia-thalamic loop is sufficient to produce beta oscillations. Additionally, most of the mechanisms of beta production that we are currently aware of in the cortex are completely different from the mechanism of beta rhythm generation that we describe in our striatal model, and depend on excitatory cells (39, 40). One study using rat brain slices shows that high beta can be elicited from M1 by coapplication of carbachol and kainate (6). This beta rhythm appears to be dependent on networks of GABAergic interneurons as well as on excitatory cell drive similar to our model of striatum. However, Yamawaki et al. (6) think the mechanism of beta in M1 is similar to the mechanism producing persistent gamma oscillations in other cortical areas.

The results presented here highlight the powerful combination of mathematical and experimental approaches in addressing problems in systems neuroscience. Dynamics of biological systems do not readily yield to direct observation using even the most sophisticated experimental approaches. We show here that informed biophysical modeling can be highly predictive of complex biological dynamics. Additionally, these findings also have broad implications in understanding beta oscillations in normal motor function as well as their inappropriate expression in other disorders with striatal involvement.

**Computational Methods**

**MSNs.** We model MSNs using single-compartment models with Hodgkin-Huxley-type dynamics. Membrane currents (I_{mem}) consist of a fast sodium current (I_Na), a fast potassium current (I_K), a leak current (I_L), and an M-current (I_m) (12). We do not model MSN up and down states, which are prevalent during sleep and anesthesia, because these transitions are not prevalent in the awake state (41), which is the focus of our model. Therefore, in our model MSNs, we do not include Kir2 currents, which are active in the MSN down state. The sum of all excitatory input from the cortex and thalamus is modeled using a background excitation term (I_{app}) and Gaussian noise.

**Networks.** MSNs connect primarily to other MSNs through GABAergic synapses (42). The GABAa current (I_{GABAa}) is modeled using a Hodgkin-Huxley-type conductance with weak GABAa maximal conductances. MSNs have extensive local axonal projections that mainly contact other MSNs (42). We construct a striatal model with 100 MSNs and examine several patterns of network connectivity: all-to-all connections, nearest neighbor connections, and 30% random connections, which approximates the rate of connectivity found in both slice and cortex-striatum-substantia nigra organotypic cultures (32, 43).

**Parkinsonian Striatum.** We model the effect of loss of dopamine indirectly through its effect on ACh-induced reduction of the M-current. Thus, we model the parkinsonian striatum by reducing the maximal M-current conductance from 1.3 to 1.2 mS/cm².

**LFP.** MSNs account for 90–95% of all neurons in the rat neostriatum (44). LFPs are modeled as the sum of all MSN to MSN GABAa currents. Stationarity of the network appears in the raster plots after ~700 ms. To ensure stationarity, our LFP is evaluated only after 1,000 ms of simulated time. More details are in SI Computational Methods.

**Experimental Methods**

All procedures were done in accordance with the National Institutes of Health Guide for Laboratory Animals and were approved by the Massachusetts Institute of Technology Animal Care and Use and Biosafety Committees. Adult C57 or Swiss Webster mice were used. Under isoflurane anesthesia, a custom-fabricated plastic head plate was surgically affixed to the skull. Adult mice were head-fixed and recorded awake with glass electrodes filled with saline. LFP was collected with a Multiclamp 700B amplifier, digitized with a Digidata 1440 and acquired with pClamp 10 software at a sampling rate of 20 kHz (Molecular Devices). Recordings and infusions in the striatum were made at the stereotactic coordinate (−0.2 to 0.8, 2.0–2.5, 3.0–3.5) that receives projections from both the motor and sensory cortices (45). Infusion in the cortex directly above the striatal infusion sites were at the stereotactic coordinate (−0.2 to 0.8, 2.0–2.5, 1.2–1.7). Carbamoylcholine chloride (Carbachol; from Sigma) was dissolved in saline and infused at a 0.2 μL/min rate. The infusion cannula was positioned ~300–700 μm away from the recording electrode tip. The LFP power spectrum was obtained with Hilbert transform in Matlab. More details are in SI Experimental Methods.

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