The subthalamic nucleus contributes causally to perceptual decision-making in monkeys

Kathryn Rogers, Joshua I. Gold, Long Ding*

Department of Neuroscience, University of Pennsylvania, Philadelphia, PA 19104

^{*} Corresponding author (lding@pennmedicine.upenn.edu)

Abstract

1

13

14

15

- 2 The subthalamic nucleus (STN) plays critical roles in the motor and cognitive function of the
- 3 basal ganglia (BG), but the exact nature of these roles is not fully understood, especially in the
- 4 context of decision-making based on uncertain evidence. Guided by theoretical predictions of
- 5 specific STN contributions, we used single-unit recording and electrical microstimulation in the
- 6 STN of healthy monkeys to assess its causal, computational roles in visual-saccadic decisions
- based on noisy evidence. The recordings identified subpopulations of STN neurons with distinct
- 8 task-related activity patterns that related to different theoretically predicted functions.
- 9 Microstimulation caused changes in behavioral choices and response times that reflected
- multiple contributions to an "accumulate-to-bound"-like decision process, including modulation
- of decision bounds and evidence accumulation, and to non-perceptual processes. These results
- provide new insights into the multiple ways that the STN can support higher brain function.

Introduction

- The subthalamic nucleus (STN) is a critical junction in both the indirect and hyperdirect
- pathways of the basal ganglia (BG). It receives inputs from the external segment of the globus
- pallidum (GPe) and cortex and sends diffuse excitation to pallidal output nuclei of the BG. The
- 19 STN has well-recognized functions in movement control. For example, in humans and monkeys,
- lesions of the STN cause involuntary movements of contralateral body parts (Martin, 1927;
- 21 Martin and Alcock, 1934; Whittier and Mettler, 1949; Carpenter et al., 1950). In monkeys with
- 22 experimentally induced Parkinsonism, STN lesions and inactivation can reverse abnormal BG
- output activity and alleviate both akinesia and rigidity (Bergman et al., 1990, 1994; Wichmann et
- 24 al., 1994a). In Parkinsonian human patients, deep brain stimulation (DBS) of the STN has
- become a common treatment option to alleviate movement abnormalities (DeLong and
- 26 Wichmann, 2001).
- 27 Recognizing that motor symptoms associated with STN damage are often accompanied by
- 28 emotional and cognitive deficits, recent work has also begun to examine the roles of the STN in
- 29 cognition. For example, the STN has been shown to contribute to cued, goal-driven action
- inhibition (Baunez et al., 2001; Desbonnet et al., 2004; Witt et al., 2004; Aron and Poldrack,
- 2006; Frank et al., 2007; Isoda and Hikosaka, 2008; Schmidt et al., 2013; Pasquereau and Turner,
- 32 2017). STN activity can also be sensitive to task complexity and decision conflict, as measured
- in imaging studies and human patients undergoing DBS (Lehericy et al., 2004; Aron et al., 2007;
- Fumagalli et al., 2011; Brittain et al., 2012; Zaghloul et al., 2012; Zavala et al., 2017). These
- 35 findings have led to the idea that STN may also contribute to resolving difficult decisions based
- on uncertain evidence. This idea has been formalized in several computational models, which
- posit three, not mutually exclusive, functions for STN: 1) through its interaction with GPe, STN
- 38 computes a normalization signal to calibrate how the available, alternative options are assessed
- 39 (Bogacz and Gurney, 2007; Coulthard et al., 2012; Green et al., 2013); 2) in coordination with
- 40 the medial prefrontal cortex, STN adjusts decision bounds (i.e., thresholds on accumulated

evidence that govern decision termination and commitment) to control impulsivity in responding (Frank, 2006; Cavanagh et al., 2011; Ratcliff and Frank, 2012; Zavala et al., 2014; Herz et al., 2016, 2017; Pote et al., 2016); and 3) by maintaining the balance between the direct and indirect pathways of the BG, STN helps to implement a nonlinear computation that improves the efficacy with which the BG adjusts decision bounds (Lo and Wang, 2006; Wei et al., 2015).

Guided by predictions of these models (Figure 1B), we assessed the role of the STN in decisions made by monkeys performing a random-dot visual motion direction discrimination task (Figure. 1A). We recorded from individual STN neurons while monkeys performed the task and found activity patterns that were highly heterogenous across neurons. Nevertheless, these patterns could be sorted into three prominent clusters with functional properties that, in principle, could support each of the three theoretically predicted STN functions from previous modeling studies. In addition, we tested STN's causal contribution to the decision process using electrical microstimulation. These perturbations of STN activity affected both choice and reaction time (RT) performance in multiple ways that could be ascribed to particular computational components of an "accumulate-to-bound" decision process. As detailed below, these results show that STN can play multiple, causal roles in the formation of a deliberative perceptual decision,

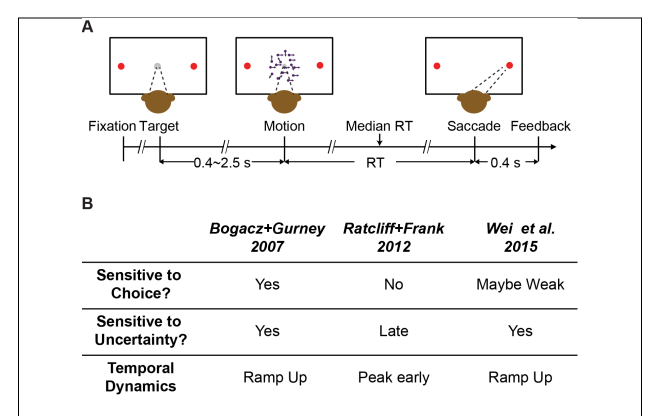


Figure 1. Behavioral task and model predictions. A, Behavioral task. The monkey was required to report the perceived motion direction of the random-dot stimulus by making a saccade towards the corresponding choice target at a self-determined time. B, Three previous models predicted different patterns of STN activity. Sensitive to choice: differential responses for trials ending with different choices. Sensitive to uncertainty: differential responses for trials with different evidence strength.

- 57 likely reflecting its diverse contributions to the many cognitive and motor functions that depend
- on the BG.

60

Results

STN neurons show diverse response profiles

- We recorded 203 neurons while the monkeys were performing a random-dot motion
- discrimination task (n = 115 and 88 for monkeys C and F, respectively). The behavioral
- performance of both monkeys has been documented extensively (Ding and Gold, 2010, 2012a;
- Fan et al., 2018). Their performance in three example sessions are shown in Figure 4A–C (black
- data points). In general, both monkeys made more contralateral choices with increasing signed
- 66 motion strength (positive for motion toward the contralateral target) and had lower RTs (i.e.,
- 67 faster responses) for higher absolute motion strength.
- 68 STN neurons showed diverse response profiles. Figure 2A shows average activity patterns of
- 69 three example neurons. The top neuron showed an initial suppression of activity after motion
- onset and became active, in a choice-dependent manner, before saccade onset. The middle
- 71 neuron showed choice- and motion coherence-dependent activation during the motion-viewing
- 72 period before saccade onset. The bottom neuron exhibited activation after motion onset that was
- 73 similar for both choices and all coherence levels, which then decayed in a choice- and coherence-
- 74 dependent manner around saccade onset.
- 75 The diversity of response profiles can be seen in the summary heatmaps for the population
- 76 (Figure 2B). When activity was averaged across all trial types, STN neurons can become
- activated or suppressed (warm vs. cool colors, respectively), relative to pre-stimulus baseline,
- during motion viewing and around saccade onset. The timing of peak modulation also spanned
- 79 the entire motion-viewing period and extended beyond saccade generation, including a
- substantial fraction of neurons that also responded to target onset before the motion stimulus
- appeared. These diverse spatiotemporal response profiles suggest that the STN as a whole may
- serve multiple functions in perceptual decision-making.
- Across the population, a substantial fraction of neurons was sensitive to choice, motion
- coherence, and RT (Figure 2C-E, Supplementary Figure 1). We performed multiple linear
- regressions, separately for coherence and RT (Eqs. 1 and 2), for each neuron and used the
- 86 regression coefficients to measure these decision-related sensitivities. For choice sensitivity
- 87 (Figure 2C, first row), both contralateral and ipsilateral preferences were commonly observed.
- 88 The overall fraction of neurons showing choice sensitivity increased after motion onset and
- 89 peaked at saccade onset (Figure 2D). For coherence sensitivity, modulations were observed for
- 90 trials with contralateral or ipsilateral choices and with similar tendencies for positive and
- 91 negative coefficients (Figure 2C, rows 2 and 3). The fraction of neurons showing reliable
- 92 coherence sensitivity was also higher around saccade onset (Figure 2D).
- Despite the diverse distributions of regression coefficients, there were systematic patterns in
- 94 when and how these forms of selectivity were evident in the neural responses. Notably, neurons
- showing choice sensitivity were more likely to show coherence modulation during early motion

viewing, especially for trials when the monkey chose the neuron's preferred choice (Figure 2E, purple). In contrast, coherence modulation emerged later for neurons that did not show choice sensitivity (Figure 2E, gray lines). These systematic interactions in modulation types suggest that the STN population does not simply reflect a random mix of selectivity for decision-related quantities. Instead, there appears to exist subpopulations with distinct decision-related modulation patterns, which we detail below.

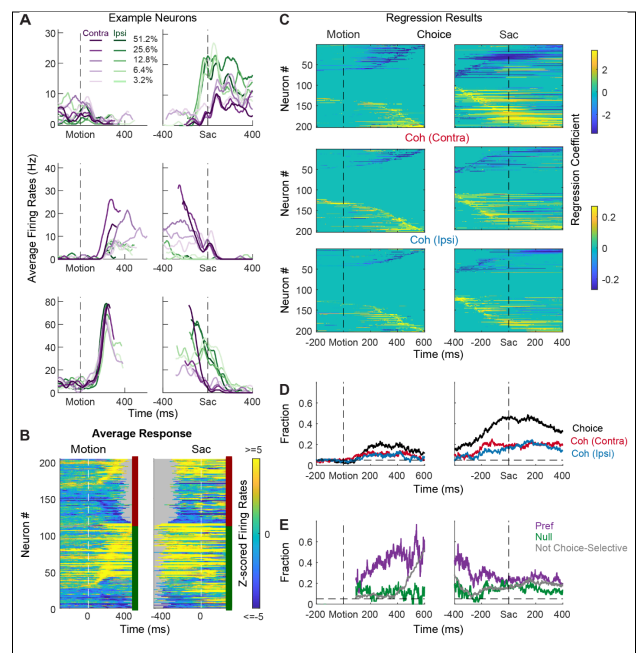


Figure 2. STN neurons have diverse response profiles. A, Activity of three STN neurons (rows) aligned to motion (left) and saccade (right) onsets and grouped by choice *x* motion coherence (see legend). For motion-onset alignment, activity was truncated at 100 ms before saccade onset. For saccade-onset alignment, activity was truncated before 200 ms after

motion onset. B, Summary of average activity patterns. Each row represents the activity of a neuron, z-scored by baseline activity in a 300 ms window before target onset and averaged across all trial conditions. Rows are grouped by monkey (red and green shown to the right of each panel: monkeys C and F, respectively) and sorted by the time of peak values relative to motion onset. Only correct trials were included. C, Heatmaps of linear regression coefficients for choice (top), coherence for trials with contralateral choices (middle), and coherence for trials with ipsilateral choices (bottom), for activity aligned to motion (left) and saccade (right) onsets. Regression was performed in running windows of 300 ms. Regression coefficients that were not significantly different from zero (t-test, p>0.05) were set to zero (green) for display purposes. Neurons were sorted in rows by the time of peak coefficient magnitude. Only correct trials were included. D. Time courses of the fractions of regression coefficients that were significantly different from zero (t-test, p<0.05), for choice (black), coherence for trials with contralateral choices (red), and coherence for trials with ipsilateral choices (blue). Dashed line indicates chance level. E, Time courses of the fractions of non-zero regression coefficients for coherence. Separate fractions were calculated for trials with the preferred (purple) and null (green) choices from choice-selectivity activity and for all trials from activity that was not choice selective (gray). Only time points after motion onset with fractions > 0.05 for choice-selective activity were included. Dashed line: chance level.

STN subpopulations can support previously theorized functions

102

103

104

105106

107

108

109

110

111

112113

114

115116

117118

119 120

121

122

123

124125

Using two forms of cluster analysis, we identified three subpopulations of neurons in the STN with distinct activity patterns that conform to predictions of each of the three previously published sets of models. For the first analysis, we represented a neuron's activity pattern with a 30-dimension vector, consisting of normalized average activity associated with two choices, five coherence levels, and three task epochs. We generated three artificial vectors based on the predicted activity patterns of each model, as follows (Figure 3A). Bogacz and Gurney (2007) posited that STN neurons, through their reciprocal connections with the external segment of globus pallidum, pool and normalize evidence-related signals, leading to the prediction of choice and coherence-modulated activity during motion viewing (Figure 3A, top; based on simulations using equations in their Appendix B). Ratcliff and Frank (2012) posited that STN neurons, through their direct innervation by cortical regions, provide an early signal to suppress immature choices, leading to the prediction of a choice-independent signal that appears soon after motion onset and dissipates over time (Figure 3A, middle; based on their Figure 5). Wei and colleagues (2015) posited that the STN balances evidence-related signals in the GPe until near decision time, leading to the prediction of coherence-dependent ramping activity with no or weak choice selectivity (Figure 3A, bottom; based on their Figure 2D). We performed k-means clustering using these three vectors and another arbitrary vector as the seeds to group the population into four clusters.

Figure 3B shows the average activity from each of the resulted clusters. Consistent with the design of this analysis, the first cluster tends to show choice- and coherence-dependent activity

that also ramps up during motion viewing (Figure 3B, first row,; Supplementary Figure 2). The

second cluster tends to show an early, sharper rise in activity during motion viewing and this

activity gradually decreases toward saccade onset (Figure 3B, second row). The third cluster tends to show ramping activity during motion viewing with similar coherence modulation for both choices and a short burst of activity for one choice just before saccade onset (Figure 3B, third row; Supplementary Figure 2). The last cluster shows mixed and, on average, weak task-

related modulation (Figure 3B, bottom row; Supplementary Figure 2B). The first three clusters

126 127

128 129

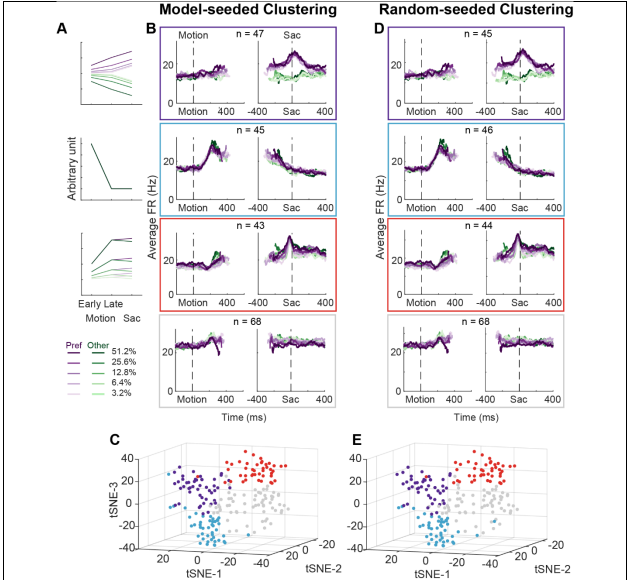


Figure 3. STN contains distinct subpopulations. A, Three activity vectors that were constructed based on theoretical predictions in Figure 1B and used as seeds for *k*-means clustering (see Methods). B, Each panel shows the average activity of neurons in a cluster, same format as Figure 2A. The numbers indicate the cluster size. C, Visualization of the clusters using the *t*-distributed stochastic neighbor embedding (*t*-SNE) dimension-reduction method. D, Average activity of clusters identified using random-seeded *k*-means clustering. Same format as Figure 3B. E, Visualization of the random-seeded clusters in the same tSNE space.

contained similar numbers of neurons. When visualized using the T-distributed stochastic neighbor embedding technique, these clusters did not form a single continuum but instead reflected separable features between clusters (Figure 3C). In other words, the clustering did not

131132

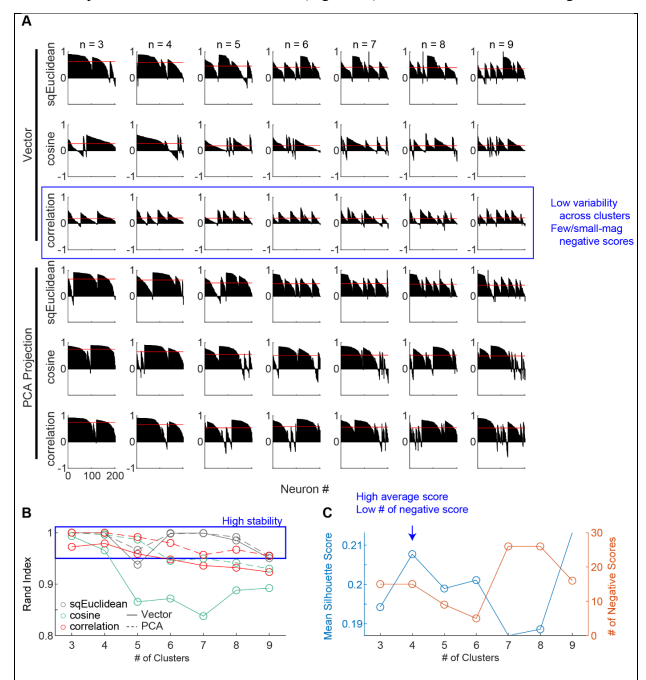


Figure 4. Clustering parameters. A, Silhouette plots for clustering results using different combinations of settings. Silhouette scores for neurons are grouped by clusters and sorted. Red lines indicate the mean scores. Yellow shaded box indicates the chosen setting for results in Figure 3. B, Average Rand indices for different clustering settings. For each setting, the k-means algorithm was run 50 times, each time picking the best clusters out of 100 repetitions. Higher Rand index indicates greater cluster stability across different runs. Blue

box indicates settings with Rand indices > 0.95. C, Mean silhouette scores and the number of negative scores as a function of number of clusters, using the firing rate vectors and correlation distance. Higher mean score and fewer negative scores indicate better clustering.

simply force a uniform distribution with random-mixed selectivity into four groups.

For the second cluster analysis, we used random seeds without considering any of the model predictions and obtained almost identical clusters. As detailed in Methods, we explored a wide range of settings for clustering, including: 1) using directly the 30-D vectors or their principal component projections, 2) basing the clustering on three different distance metrics, and 3) varying the number of presumed clusters. To identify the best setting, we assessed the goodness of clustering using the silhouette score and the stability of clustering using the Rand index (Rand, 1971) (Figure 4). The silhouette score quantifies for each member the relative distance between its average within-cluster distance and distance to those in its closest neighboring cluster (a higher score indicates better cluster separation). The silhouette plots favored the combination of using the 30-D vector directly and correlation distance (Figure 4A, third row), which generated less variability across clusters and few/small-magnitude negative silhouette scores (negative scores indicate that a member is closer to its neighboring cluster than its own cluster).

The Rand index measures how consistently two members are assigned to the same clusters from different iterations of clustering (a high index indicates greater stability). The Rand index was generally high (above 0.95 out of a max of 1; blue box) except for the combination of using the 30-D vector and cosine distance (Figure 4B). Finally, using the raw vector-correlation combination, an assumption of four clusters resulted in the highest average Rand index and assuming 4-6 clusters generally resulted in higher mean silhouette score and lower number of negative scores (Figure 4C; blue arrow). We thus considered that the raw-vector-correlation combination and an assumption of four clusters produced the most stable and plausible results.

As shown in Figure 3D and E, the four clusters thus identified closely matched those obtained using model predicted seeds, in terms of the average activity, the cluster sizes, and their locations in the tSNE space. Increasing the assumed number of clusters caused changes mostly in the gray cluster, with some changes in the blue cluster, and little effects on the red and purple clusters (Supplementary Figures 3 and 4). Together these results suggest that, absent the ground truth on the number of subpopulations in STN, there exist at least three subpopulations that each corresponds to the predictions of one of three previous published models. As a consequence, STN appears in principle to be able to support multiple decision-related functions.

Perturbation of STN activity affects choice and RT

To better understand STN's functional roles in the decision process, we perturbed STN activity using electrical microstimulation while monkeys performed the task. Specifically, we applied a train of current pulses at identified STN sites during decision formation, lasting from motion onset to saccade onset. Figure 5 shows microstimulation effects on choices and RTs in three example sessions. In the first example session, STN microstimulation caused a leftward

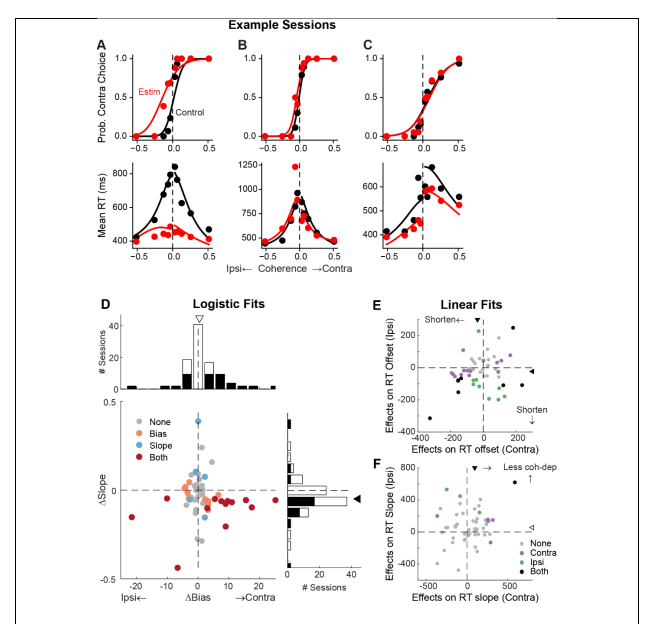


Figure 5. STN microstimulation affects monkeys' choice and RT. A-C, Monkey's choice (top) and RT (bottom) performance for trials with (red) and without (black) microstimulation for three example sessions (A,B: monkey C; C: monkey F). Lines: DDM fits. D, Distributions of microstimulation effects on bias and slope terms of the logistic function. Filled bars in histograms indicate sessions with significant modulation of the specific term (bootstrap method). Triangles indicate the median values. Filled triangle: Wilcoxon sign-rank test for H_0 : median=0, p < 0.05. E and F, Summary of microstimulation effects on the offset (E) and slope (F) terms of a linear regression fit to RT data. Two separate linear regressions were performed for the two choices (lpsi/Contra, as indicated). Triangles indicate the median values. Filled triangles: Wilcoxon sign-rank test, p < 0.05.

horizontal shift (more contralateral choices) and slope reduction (more variable choices) in the psychometric curve (Figure 5A, top), as well as a substantial flattening of RT curves (faster

170

172 responses that depended less on motion coherence) for both choices (bottom). In the second

example session, STN microstimulation induced a minor leftward shift in the psychometric curve

and asymmetric changes in RT for the two choices (Figure 5B). In the third example session,

STN microstimulation did not change the psychometric curve but caused reductions in RT for

both choices (Figure 5C).

174

180

182

183

204

205

207

211

Across 54 different STN sites, microstimulation caused variable choice biases and tended to

178 reduce the dependence of choice on motion strength. We fitted the choice data to a logistic

function and measured choice bias (horizontal shift) and motion strength-dependence (slope). In

23 sessions, microstimulation induced a reliable choice bias (Figure 5D). The induced bias was

toward the contralateral or ipsilateral choice in 15 and 8 sessions, respectively, and the median

value for bias was not significantly different from zero (Wilcoxon sign-rank test, p = 0.15). In 18

sessions, microstimulation induced a change in the slope. The slope was reduced in 15 sessions

and the median value was negative (p = 0.008). These tendencies were robust across different

variants of logistic functions, with or without lapse terms to capture errors independent of motion

strength (Supplementary Figure 5). Of the sessions where inclusion of lapse terms for the control

and microstimulation trials produced lower AICs, very few showed significant microstimulation-

induced changes in lapses (2 sessions each for the "Symmetric Lapse" and "Asymmetric Lapse"

variants). Thus, based on fitting results using logistic functions, STN microstimulation most

consistently reduced the choice dependence on motion strength, caused session-specific choice

biases, and had minimal effects on lapses.

Microstimulation also tended to reduce RT. We fitted linear functions separately for RTs

associated with the two choices, in which offset and slope terms measure coherence-independent

and -dependent changes in RTs, respectively. Microstimulation caused changes in RT offsets in

25 sessions for contralateral choices (18 were reductions in RT, with a median change across all

sessions of -36 ms; Wilcoxon sign-rank test for H_0 : median change=0, p < 0.0001) and 18

sessions for ipsilateral choices (15 reductions, mean change = -23 ms, p<0.0001; Figure 5E).

Microstimulation caused changes in RT slopes in 6 sessions for contralateral choices (5 were

positive, implying a weaker coherence dependence; p < 0.0001) and 4 sessions for ipsilateral

200 choices (all 4 were positive; Figure 5F). Thus, based on fitting results using linear functions,

201 STN microstimulation can induce choice-specific changes in RT, with overall tendencies to

reduce both the coherence-independent component and the RT's dependence on coherence for

the contralateral choice.

Microstimulation effects reflected changes in multiple computational components

To infer STN's computational roles in the decision process, we examined the microstimulation

effects using a drift-diffusion model (DDM) framework. This framework has been widely used in

studies of perceptual decision-making and can provide a unified, computational account of both

209 choice and RT (Gold and Shadlen, 2007). It assumes that noisy evidence is accumulated over

210 time and a decision is made when the accumulated evidence reaches a certain decision bound.

The overall RT is the sum of the time needed to reach the bound and non-decision times

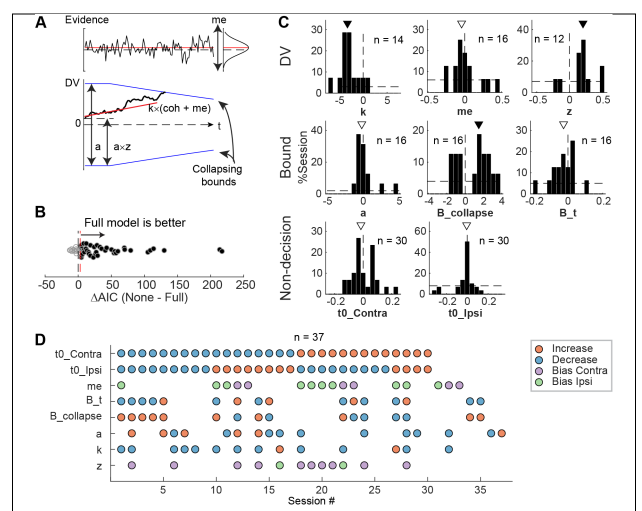


Figure 6. STN microstimulation affected multiple computational components in the **DDM.** A, Illustration of the DDM. Red/black lines represent across-trial mean/single-trial example of the evidence (top) and drift rate (bottom). Blue lines represent the collapsing decision bounds. B, Distribution of the difference in AIC between the None and Full models. Red dashed line indicates the criterion for choosing the full model: AIC difference = 3. C, Histograms of microstimulation effects on DDM parameters. Each histogram included only sessions in which the Full model outperformed the corresponding reduced model (e.g., the histogram for parameter *a* included only sessions in which AIC_{NoA} – AIC_{Full} > 3 and AIC_{None} – AIC_{Full} > 3). Triangles indicate median values. Filled triangles: Wilcoxon sign rank test, p < 0.05. D, Summary of microstimulation effects on all parameters, for sessions in which at least one significant effect was present. Sessions were sorted by the prevalence and sign of the effects.

reflecting perceptual and motor latencies. Previous theoretical models also made predictions about the effects of perturbing STN activity that can be interpreted in the DDM framework. The model by Bogacz and Gurney (2007) predicted that the perturbation would reduce the effect of task difficulty on decision performance by eliminating a nonlinear transformation that is needed for appropriate evidence accumulation (Green et al., 2013). The model by Ratcliff and Frank

212

213

214

215

(2012) predicted that the perturbation, by causing changes in the STN's influence onto the 217 substantia nigra pars reticulata (SNr), would change temporal dynamics of the decision bound 218 219 and influence non-decision time. The model by Wei and colleagues (2015) predicted that the 220 perturbation would result in a reduction in the decision bound. To test whether these predictions, and/or other effects, were present in our microstimulation data, 221 we fitted a DDM to choice and RT data simultaneously (Figure 6A). We performed AIC-based 222 model selection and found that, in 40 of 54 sessions, the Full model, which included 223 224 microstimulation effects on any model parameters, outperformed the None model, which assumed that there was no microstimulation effect on any parameters (Figure 6B). This result 225 implies that, in these sessions, STN microstimulation affected one or more computational 226 227 components of the decision process. To better characterize these effects, we compared AICs between the Full model and six reduced models to identify sessions with reliable 228 microstimulation-induced changes in particular model parameters (Supplementary Figure 6A). 229 We found that STN microstimulation resulted in reliable changes in several model parameters 230 231 over different subsets of sessions (Figure 6C and D). Consistent with model predictions from 232 Bogacz and Gurney (2007), microstimulation reduced the scale factor for evidence accumulation, k, in 14 sessions. This effect contributed to a decreased motion coherence 233 dependence of choice and RT (Figure 6C, first histogram; Wilcoxon sign-rank test for H_0 : zero 234 median effect, p = 0.021). Consistent with model predictions from Ratcliff and Frank (2012) and 235 Wei and colleagues (2015), microstimulation affected parameters that controlled the decision 236 bound (a, B collapse, B t) in 16 sessions each (not necessarily in the same sessions for each 237 parameter, see Figure 6D). The changes in the maximal decision bound (a) were variable across 238 sessions (p = 0.68). The changes in the collapsing bound dynamics (B collapse, B t) tended to 239 indicate faster and earlier decreases in bounds (p = 0.039 and 0.088, respectively). Consistent 240 with model predictions from Ratcliff and Frank (2012), microstimulation caused changes in non-241 decision times in 30 sessions (t0 Contra and t0 Ipsi). These changes varied from session to 242 session (p = 0.28 and 0.75, respectively). Statistical tests on fitted parameters of all sessions, 243 regardless of whether a microstimulation effect was necessary to account for the behavioral data, 244 showed similar trends (Supplementary Figure 6B). 245 STN microstimulation had two additional effects beyond those predicted by the previous 246 modeling studies. First, consistent with the above-demonstrated microstimulation-induced choice 247 biases (Figure 5), microstimulation induced offsets in momentary (me; n = 16 sessions; p = 0.61) 248 and accumulated (z; n = 12 sessions; p = 0.016) evidence. Second, the microstimulation effects 249 involved changes in more than one model parameter in the majority of sessions (Figure 6D). We 250 did not observe any dominant combinations of effects. These results suggest that the STN is 251

Distribution of microstimulation effects reflected intermingled neuron activity patterns

causally involved in multiple decision-related functions, including those mediating the

dependence on evidence, choice biases, and bound dynamics.

252

253

254

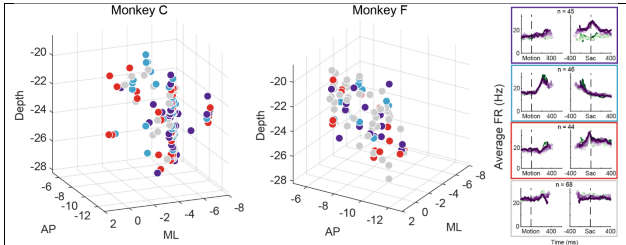


Figure 7. Different STN subpopulations are intermingled. Locations of STN neurons, color-coded by clusters based on random-seed clustering (same as Figure 3D). The Medial-Lateral (ML) values were jittered for better visualization of neurons recorded along the same track and at similar depths. Anterior-Posterior (AP) levels were relative to the anterior commissure. ML and depth levels were relative to the center of the recording chambers.

The multi-faceted microstimulation effects, combined with the fact that the kind of microstimulation we used tends to activate not just one neuron, but rather groups of neurons near the tip of the electrode (Tehovnik, 1996), suggested that STN neurons with different functional

roles are located close to one another. Consistent with this idea, neurons that were classified as belonging to different clusters tended to be intermingled (Figure 7). We did not observe any consistent topographical organization patterns within or between the two monkeys. At certain locations, neurons belonging to different clusters were recorded using the same electrode. We calculated silhouette scores to quantify whether the activity pattern-based neuron clusters also formed clusters in the 3D physical space. The mean values were -0.09 and -0.11 for the two monkeys, respectively, indicating that neurons were often closer to others from a different cluster than those within the same cluster. In other words, STN subpopulations did not segregate from each other and instead tended to be intermingled, and thus microstimulation likely activated multiple neurons with different functional properties.

Although the intermingled organization of STN subpopulations, defined based on their task-related activity patterns (Figure 3), made it challenging to relate a specific microstimulation effect to a specific subpopulation, we did observe certain trends that could contribute to the site-specific microstimulation effects. We assigned the single or multi-unit activity at the stimulation sites according to the clusters identified using the random-seeded clustering (Figure 8A). We then grouped the sites by neuron clusters (Figure 8C). When neurons of different clusters were recorded at the same site, the same microstimulation effects were assigned to each cluster. We found that the second cluster was associated with lower overall likelihood of observing microstimulation effects compared to other clusters (Figure 8B; Chi-square test, H_0 : the likelihood is the same for the first cluster and the other clusters; p = 0.003), while the third cluster had higher overall likelihood (p = 0.035). For the first three clusters, no microstimulation

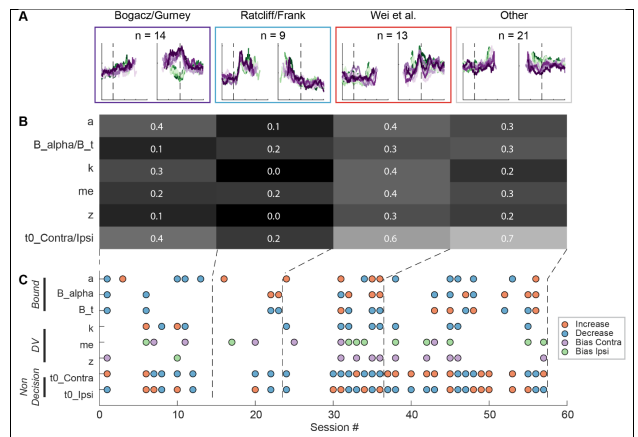


Figure 7. STN microstimulation effects depend partially on neural clusters. A, Average activity at stimulation sites, grouped by four clusters that were identified in Figure 3D. B, Fractions of significant microstimulation effects for sites with the presence of each neuron cluster. Significance was based on AIC comparison between reduced and Full models. C, Microstimulation effects grouped by neuron cluster. Same format as Figure 5D. D, Fractions of significant microstimulation effects, grouped by effects reflecting changes in bound, decision variable computation, and non-decision processes.

effect dominated (p > 0.3 for all), whereas it was more likely to observe effects on non-decision times for the fourth cluster (i.e., with neural activity patterns not related to the three models; p = 0.001).

The sign of microstimulation effects depended weakly on neuron clusters. For example, it was more likely to observe an increase in maximal bound height ("a") for the third neuron cluster (Chi-square test, H_0 : same fractions of increase/decrease for all clusters; p = 0.073; Chi-square test, H_0 : equal fractions of increase/decrease within the cluster; p = 0.036). Microstimulation decreased the scale factor ("k") for the third and fourth clusters but caused variable changes for the first cluster (p = 0.070 and 0.021, respectively). Microstimulation effects on the non-decision time for the contralateral choices were dominated by increases for the fourth cluster (p = 0.04 and 0.007, respectively). Together, these results suggested that microstimulation effects reflected multiple contributions of intermingled STN subpopulations to decision- and non-decision-related processes.

Heterogeneous activity patterns and microstimulation effects cannot be explained by

variations in motivational state

- Another potential source of heterogeneity in our data may reflect variations in the monkeys'
- 296 motivational state across sessions. In two sets of analyses, we did not observe any significant
- influence of motivational state on the recording or microstimulation data. For these analyses, we
- used the rate of fixation break, overall error rate, and mean RT as indices of motivational state.
- None of these measurements differed among sessions when different subpopulations were
- encountered (Supplemental Figure 7), suggesting that the motivational state cannot predict which
- 301 type of activity pattern would be observed. Similarly, none of these measurements significantly
- 302 correlated with the microstimulation effects in any DDM component (Supplemental Table 1),
- suggesting that the motivational state did not modulate the magnitude of microstimulation
- effects. Together, these results suggest that the diverse activity patterns and microstimulation
- effects cannot be accounted for by variations in the monkeys' task engagement.

Discussion

293

294

- We provide the first characterization of single-unit recordings and electrical microstimulation in
- 308 the STN of monkeys performing a demanding perceptual-decision task. We show that: 1) STN
- neurons are heterogeneous in their response profiles; 2) different STN subpopulations, with
- distinct decision-related activity modulation patterns and intermingled within the region, can
- support previously-theorized functions; and 3) electrical microstimulation in STN causes
- changes in choice and RT behaviors, reflecting effects on multiple computational components of
- an accumulate-to-bound decision process. These results indicate that the STN plays important
- and complex roles in perceptual decision formation, both supporting and extending existing
- views of STN function.
- Our study was motivated by the differing predictions of STN activity patterns from several
- 317 theoretical studies that were based on STN cellular physiology, connectivity, and/or response
- patterns in non-perceptual decision-making contexts (Bogacz and Gurney, 2007; Ratcliff and
- Frank, 2012; Wei et al., 2015). Remarkably, we found three clusters of STN activity that are
- 320 consistent with each of these predictions. The three clusters were robust and stable, emerging
- when we used two different clustering methods (one with model-based seeds, the other with
- random seeds). Interestingly, Zavala and colleagues (2017) have reported two types of STN
- responses in human patients performing a flanker task. The "early" response they identified may
- 324 correspond to our second cluster, while the "late" response may reflect a combination of our first
- and third clusters. Together these results suggest that the primate STN contains distinct
- subpopulations with different functional roles. Combined with the microstimulation results, the
- presence of these subpopulations suggests that the STN can both contribute to the conversion of
- sensory evidence into an appropriately formatted/calibrated decision variable and modulate the
- dynamics of decision bound. Future studies of BG function should strive to better understand
- how these subpopulations interact with each other, as well as with other neurons in the BG and
- the larger decision network to support decision making.

334

335

336337

338

339

340

341

342

343 344

345

346

347

348

349

350

351

352

353

354

355 356

357

358

359

360

361

362

363

364

365

366

367

368

369

370

371

372

373

Despite the general agreements between our observations and previous theoretical predictions, there were also differences that could be informative for developing future BG models. Most notably, previous models focused on the period of evidence accumulation and less on neural activity patterns at or after decision commitment. In contrast, our data show interesting modulations around saccade onset (see Figure 3) that raise several intriguing possibilities for STN's contributions to the decision process. In particular, one subpopulation showed a broad peak with strong choice modulation and little coherence modulation. These modulations may reflect bound-crossing in an accumulate-to-bound process (but see below). A second subpopulation returned to the baseline level before saccade onset. The relatively constant trajectory of this modulation may reflect a collapsing bound or urgency signal that is dependent only on elapsed time and not the sensory evidence. A third subpopulation maintained coherencedependent activity until very close to saccade onset and showed a sharp peak with little choice or coherence modulation. This sharp peak may signal the end of decision deliberation, without specifying which decision is made, to direct the network to a post-decision state for decision evaluation. The diverse activity patterns and their intermingled distribution in the STN underscore the challenge of identifying specific, causal contributions of a particular neural subpopulation. In many sessions, we observed effects that have been predicted theoretically and observed experimentally in human PD patients undergoing DBS. These effects included a reduction in RT, a weaker dependence on evidence, and changes in the maximal value and trajectories of the decision bound (Frank et al., 2007; Cavanagh et al., 2011; Coulthard et al., 2012; Green et al., 2013; Zavala et al., 2014; Herz et al., 2016; Pote et al., 2016). In addition to these previously observed effects, microstimulation also changed choice biases, measured as horizontal shifts of psychometric functions and as two different types of biases in the DDM framework. This departure from previous DBS studies may arise from different task designs (button press versus eye movement), health status of the subjects, and experience level (minimally versus extensively trained). The lateralized bias suggests that the STN may be involved in flexible decision processes that adapt to environments with asymmetric prior probability and/or reward outcomes for different alternatives, in addition to modulating speed-accuracy tradeoff. Consistent with this idea, DBS can affect the threshold for deliberations over uncertain sensory inputs or motivational factors such as reward and effort (Pagnier et al., 2024), suggesting that the STN may be part of a general selection machinery that can incorporate sensory evidence with information about the task environment (Redgrave et al., 1999). The intermingled subpopulations may appear at odds with the conventional idea of topography in how the STN is organized. For example, the "tripartite model" suggests that STN is segregated by motor, associative, and limbic functions (Parent and Hazrati, 1995); afferents from motor cortices and neurons related to different types of movements are largely somatotopically organized in the STN (DeLong et al., 1985; Nambu et al., 1996); and certain molecular markers are expressed in an orderly pattern in the STN (reviewed in Prasad and Wallén-Mackenzie, 2024). Because we focused on STN neurons that were responsive on a single oculomotor decision task, our sampling was likely biased toward STN subdivisions related to associative function and oculomotor movements. As such, our results do not preclude the presence of

- topography at a larger scale. Rather, our results underscore the importance of activity pattern-
- based analysis, in addition to anatomy-based analysis, for understanding the functional
- 376 organization of the STN.
- Our findings also suggest that STN's role in decision formation differs in important ways from
- other oculomotor regions that have been examined under similar conditions. First, in the frontal
- eye field (FEF), lateral intraparietal area (LIP), and superior colliculus (SC), decision-related
- neural activity is dominated by a choice- and coherence-dependent "ramp-to-bound" pattern
- 381 (Roitman and Shadlen, 2002; Ding and Gold, 2012a; Crapse et al., 2018; Cho et al., 2021; Jun et
- al., 2021; Stine et al., 2023), with additional multiplexing of decision-irrelevant signals (Meister
- et al., 2013). In contrast, different STN subpopulations can carry distinct signals that may all be
- relevant to decision formation. Moreover, these signals include patterns not evident in the other
- regions, such as a choice- and coherence-independent activation in early motion viewing (blue
- cluster in Figure 3B and D), that may signal a unique role for the STN.
- 387 Second, choice-selective ramping activity has been identified in LIP, FEF, SC, the caudate
- nucleus, and now two STN subpopulations (Ding and Gold, 2010; Fan et al., 2020). However,
- such activity differs among these oculomotor regions just before saccade onset for the preferred
- choice. In LIP, FEF, and SC, when the ramping activity is aligned to saccade onset, it shows
- 391 negative coherence modulation and positive RT modulation before converging to a common,
- 392 higher level, consistent with an accumulate-to-bound process. In the caudate nucleus, the
- ramping activity does not converge to a common, higher level. For the first STN subpopulation
- 394 (Figure 3), the ramping activity showed on average positive coherence modulation and negative
- 395 RT modulation (opposite to predictions of an accumulate-to-bound process) before converging to
- a common, higher level. The second STN subpopulation did not show choice-selective activity
- before saccade onset. These differences suggest that the caudate and STN neurons participate in
- decision deliberation but do not directly mediate decision termination (bound crossing). It is also
- 399 possible that the ramping activity reflects some roles for the STN in the evaluation of the
- decision process, the tracking of elapsed time, or both. How these possible roles relate to those of
- 401 caudate neurons awaits further investigation (Fan et al., 2024).
- 402 Third, whereas unilateral perturbations in LIP and SC tend to induce contralateral choice biases
- 403 (Hanks et al., 2006; Jun et al., 2021; Jeurissen et al., 2022; Stine et al., 2023), unilateral STN
- 404 (and caudate) microstimulation can induce both contralateral and ipsilateral choice biases,
- depending on the stimulation site (Ding and Gold, 2012b; Doi et al., 2020). At many sites, STN
- 406 microstimulation effects on RT were often bilateral and of the same polarity. Moreover, STN
- 407 microstimulation seems to have a particularly strong effect on the overall dependence of choice
- and RT on evidence, which was not the case for other oculomotor regions. These differences
- suggest that the STN has unique roles in choice-independent computations, potentially including
- 410 those involving evidence pooled for all alternatives or general bound dynamics (Bogacz and
- 411 Gurney, 2007; Ratcliff and Frank, 2012).
- In summary, we characterized single-neuron activity and the effects of local perturbations in the
- STN of monkeys performing a deliberative visual-oculomotor decision task. Our results
- validated key aspects of previous theoretical predictions, providing experimental evidence for the

- 415 multiple involvement in modulating decision deliberation and commitment. Our results also
- 416 identified other features of decision-related processing in STN that differ from both theoretical
- 417 predictions and known properties of other brain areas that contribute to these kinds of decisions.
- These differences can help guide future investigations that aim to delineate how cortical-
- subcortical interactions in general, and interactions involving the STN in particular, support
- decision-making and other aspects of higher brain function.

Methods

421

422

- 423 For this study, we used two adult male rhesus monkeys (*Macaca mulatta*) that have been
- extensively trained on the direction-discrimination (dots) task. All training, surgery, and
- experimental procedures were in accordance with the National Institutes of Health Guide for the
- 426 Care and Use of Laboratory Animals and were approved by the University of Pennsylvania
- 427 Institutional Animal Care and Use Committee (protocol # 804726).
- 428 <u>Task design and electrophysiology</u>
- The behavioral task (Figure 1A), general surgical procedure, and data acquisition methods have
- been described in detail previously (Ding and Gold, 2010, 2012b). Briefly, the monkey was
- required to report the perceived motion direction of the random-dot stimulus with a saccade at a
- self-determined time. Trials with different motion coherences (drawn from five levels) and
- directions were interleaved randomly. The monkey's eye position was monitored with a video-
- based eye tracker and provided reward/error feedback online based on comparisons between the
- monkey's eye position and task-relevant locations. Saccade reaction time (RT) was measured
- offline with established velocity and acceleration criteria. Neural activity was recorded using
- glass-coated tungsten electrodes (Alpha-Omega) or polyamide-coated tungsten electrodes (FHC,
- Inc.), using a grid system through a recording chamber with access to the STN. For
- microstimulation sessions, lower-impedance FHC electrodes were used to record and stimulate at
- the same sites. Single units were identified by offline spike sorting (Offline Sorter, Plexon, Inc.).
- 441 Electrical microstimulation was delivered using Grass S88 stimulator as a train of negative-
- leading bipolar current pulses (250 us pulse duration, 200 Hz) from motion onset to saccade
- onset. For most sessions, a current intensity of 50 µA was used. In other sessions, we lowered the
- intensity to ensure that microstimulation did not abolish the monkey's ability to complete the
- 445 trials. We randomly interleaved trials with and without microstimulation at a 1:1 ratio.

Localizing the STN

- We obtained structural MRI scans using T1- MPRAGE and/or T2-SPACE sequences. We
- estimated the likely chamber coordinates with access to the STN from these images (and 3D
- reconstruction using BrainSight from Rogue Research, Inc) and mapped the surrounding areas
- electrophysiologically. Specifically, we identified several putative landmark regions, including 1)
- 451 thalamus, which showed characteristic bursts of activity in a low-firing background while the
- monkey dozed off; 2) reticular nucleus of the thalamus, where neurons exhibited high baseline
- 453 firing rates (with bursts sometimes > 100Hz); 3) zona incerta, where neurons exhibited low, tonic

- baseline firing and briefly paused their activity around saccades (Ma, 1996); 4) substantia nigra,
- pars reticulata, where some neurons showed high baseline firing rates and suppression in activity
- around visual stimulus or saccade onset (Hikosaka and Wurtz, 1983); and 5) substantia nigra,
- pars compacta, where neurons showed low baseline firing and responded to unexpected reward.
- Based on a macaque brain atlas (Saleem and Logothetis, 2007) and previously reported STN
- activity patterns (Matsumura et al., 1992; Wichmann et al., 1994b; Isoda and Hikosaka, 2008),
- we defined STN as the area that: 1) was surrounded by these landmark regions, 2) was separated
- 461 from them by gaps with minimal activity (white matter), and 3) exhibited irregular firing patterns
- with occasional short bursts. The baseline firing rate, measured within 50 ms before fixation
- point onset, had a mean±SD magnitude of 15.4±12.4 spikes/s in our sample.

464 Neural-activity analysis

- We measured the firing rates for each neuron and trial condition in running windows (300 ms)
- aligned to motion and saccade onsets. To visualize the overall activation/suppression, we
- averaged the firing rates across trial conditions and computed the z-scores using a 300 ms
- 468 window before motion onset as the baseline. To visualize the overall choice preferences, we
- averaged the firing rates for each choice, computed the difference between choices, and z-scored
- 470 the difference using the same baseline window. To quantitatively measure each neuron's task-
- 471 related modulation, we performed two multiple linear regressions for each running window,
- separately for coherence and RT because monkeys' RT strongly depends on coherence on our
- 473 task:
- Spike count = $\beta_0 + \beta_{Choice} \times I_{Choice} + \beta_{Coh-Contra} \times I_{Coh-Contra} + \beta_{Coh-Ipsi} \times I_{Coh-Ipsi}$
- 475 (Eq. 1)
- Spike count = $\beta_0 + \beta_{Choice} \times I_{Choice} + \beta_{RT-Contra} \times I_{RT-Contra} + \beta_{RT-Ipsi} \times I_{RT-Ipsi}$ (Eq.
- 477 2)
- where $I_{Choice} = \{1 \text{ for contralateral choice}, -1 \text{ for ipsilateral choice} \}$,
- 479 $I_{Coh-Contra} = \{\text{coherence for contralateral choice}, 0 \text{ for ipsilateral choice}\},$
- 480 $I_{Coh-Ipsi} = \{0 \text{ for contralateral choice, coherence for ipsilateral choice}\}.$
- 481 $I_{RT-Contra} = \{RT \text{ for contralateral choice}, 0 \text{ for ipsilateral choice}\},$
- 482 $I_{RT-lpsi} = \{0 \text{ for contralateral choice}, RT \text{ for ipsilateral choice}\}.$
- 483 Contralateral/ipsilateral choices refer to saccades toward the target contralateral/ipsilateral to the
- 484 recording sites. Significance of non-zero coefficients was assessed using a t-test (criterion: p =
- 485 0.05).

486

487

Cluster analysis

- We converted each neuron's activity into a 30-D vector consisting of the average firing rate
- within three 200-ms windows for all trial conditions (i.e., 2 choices x 5 coherence levels). The
- windows were selected as early motion viewing (100 300 ms) after motion onset), late motion
- viewing (300 500 ms after motion onset), and peri-saccade (100 ms before to after saccade)

onset). The choice identity was designated as either "preferred" and "other", based on the

relative average activity in the peri-saccade window. Note that this designation was used so that

- neurons with similar general modulation patterns except for the polarity of their choice
- selectivity would be grouped together. This designation was not based on any statistical test and
- did not imply that the peri-saccade activity was reliably choice selective. The average firing rate
- for each neuron was then z-scored based on baseline rates measured in a 300 ms window ending
- 498 at motion onset.
- We explored multiple method variations using k-means clustering and present results from the
- variation with the highest stability. These variations included: 1) whether or not the vectors were
- projected onto 11 principal components that together explained at least 95% of total variance;
- and 2) calculation of vector distance, including squared Euclidean, cosine, and correlation
- metrics. We determined the best settings using: 1) the Rand index (Rand, 1971), which quantifies
- the stability of clusters in repeated clustering; 2) Silhouette scores, which quantifies the quality
- of grouping and separation between clusters; and 3) visual inspection of clustering results in
- terms of both cluster distribution in a t-SNE space and average activity of the clusters. To
- 507 compute the Rand index, we performed 50 runs of clustering, assuming 3-9 clusters, for each
- 508 combination of variations. The Rand index was computed as the fraction of consistent grouping
- between a pair of units between two clustering runs. For two runs of clustering results, Rand
- index = $\frac{N_{same-same} + N_{different-different}}{N_{all\ pairs}}$, where $N_{same-same}$ counts the number of neuron pairs that
- share clusters in both runs, $N_{different-different}$ counts the number of neuron pairs that do not
- share clusters in either run, and $N_{all\ pairs}$ counts the total number of neuron pairs. To compute
- 513 the Silhouette scores, we chose the best of 100 repetitions of clustering for each combination of
- variations. For each neuron, Silhouette score = $\frac{\max(D_{inter-cluster}, D_{intra-cluster})}{D_{inter-cluster}, \text{ where}}$, where
- $D_{inter-cluster}$ is the average distance to the neuron's nearest neighboring cluster, and
- $D_{intra-cluster}$ is the average distance to other neurons in the same cluster. A positive score
- 517 implies that, for the given neuron, its activity was more similar to other neurons within the same
- cluster than those in its nearest neighboring cluster. A negative score implies that the neuron's
- activity was more similar to those outside its own cluster.
- To classify activity recorded at a microstimulation site, we calculated the correlation between its
- 30-D vector and the centroids from random-seeded clustering. The centroid with the highest
- 522 correlation value determined the cluster identity of the activity.
- 523 <u>Microstimulation-effects analysis</u>
- We analyzed microstimulation effects in several ways. To characterize the effects without
- assumptions about the underlying decision process, we fitted logistic functions to the choice data
- and linear functions to the RT data. We used three variants of the logistic functions that differed
- in their use of lapse rates, which measure the probability of errors independent of motion
- 528 strength:
- No Lapse: p (contralateral choice) = $\frac{1}{1+e^{-(Slope_0+Slope_{estim})\times(Coh+Bias_0+Bias_{estim})}}$ (Eq. 3)

Symmetric Lapse: p (contralateral choice) = $\lambda_0 + \lambda_{estim} + \frac{1-2\times(\lambda_0+\lambda_{estim})}{1-2}$

531
$$\frac{1-2\times(\lambda_0+\lambda_{estim})}{1+e^{-(Slope_0+Slope_{estim})\times(Coh+Bias_0+Bias_{estim})}}$$
 (Eq. 4)

Asymmetric Lapse: p (contralateral choice) = $\lambda_{Ipsi0} + \lambda_{Ipsi-estim} +$

$$\frac{1 - \lambda_{Ipsio} - \lambda_{Ipsi-estim} - \lambda_{Contrao} - \lambda_{Contra-estim}}{1 + e^{-(Slope_0 + Slope_{estim}) \times (Coh + Bias_0 + Bias_{estim})}}$$
(Eq. 5)

- where Coh is the signed coherence (positive/negative for motion toward the
- contralateral/ipsilateral choice). Contralateral/ipsilateral choices refer to saccades toward the
- targets contralateral/ipsilateral to the microstimulation sites, respectively. To assess the
- significance of the "estim" terms, we used bootstrap methods. Specifically, we generated 200
- sets of data by shuffling the microstimulation status of trials within each session. We fitted these
- artificial data using the same logistic functions to estimate null distributions for each parameter
- and performed a one-tailed test to determine if the actual fit value exceeded chance (criterion, p
- 541 < 0.05).
- We fitted linear functions to the RT data, separately for the two choices:

$$RT = Offest_0 + Offset_{estim} + (Slope_0 + Slope_{estim}) \times Coh_{unsigned}$$
 (Eq. 6)

- We assessed significance using *t*-tests (criterion, p < 0.05).
- To infer microstimulation effects on decision-related computations, we fitted drift-diffusion
- models to choice and RT data simultaneously. We used DDM variants with collapsing bounds
- 547 (DDM; Figure 7A), following previously established procedures (Fan et al., 2018; Doi et al.,
- 548 2020). Briefly, the DDM assumes that motion evidence is accumulated over time into a decision
- variable (DV), which is compared to two collapsing choice bounds. A choice is made when the
- DV crosses either bound, such that the time of crossing determines the decision time and the
- identity of the bound determines the choice identity. The model has eight basic parameters
- (presented here in six groups): 1) a, the maximal bound height; 2) B collapse and B t, the decay
- speed and onset specifying the time course of the bound "collapse"; 3) k, a scale factor governing
- the rate of evidence accumulation; 4) me, an offset specifying a bias in the rate of evidence
- accumulation; 5) z, an offset specifying a bias in the DV, or equivalently, asymmetric offsets of
- equal magnitude for the two choice bounds; and 6) $t0_{contra}$ and $t0_{insi}$, non-decision times for
- the two choices that capture RT components that do not depend on evidence accumulation (e.g.,
- visual latency and motor delay).
- We used 8 variants of DDM. In the Full model, all eight parameters were allowed to change with
- microstimulation. In the None model, all eight parameters did not change with microstimulation.
- In six reduced models (NoA, NoCollapse, NoK, NoME, NoZ, NoT), the corresponding group of
- parameters (specified above) were fixed while the other parameters were allowed to change with
- microstimulation. We fitted each model using the maximum a posteriori estimate method and
- previously established prior distributions (Wiecki et al., 2013). We performed five runs for each
- 565 fit and used the best run (highest likelihood) for analyses here. We used the Akaike Information
- 566 Criterion (AIC) for model selection. We considered an AIC difference >3 to indicate that the
- smaller-AIC model significantly outperformed the larger-AIC model. For a given sessions, if the

Full model outperformed a reduced model and the None model, we considered that session to show significant microstimulation effect(s) on the corresponding model parameter(s). For example, we considered STN microstimulation to induce significant changes in *k* if the Full model outperformed both None and NoK models for a given session.

Acknowledgements

We thank Jean Zweigle for outstanding animal care and training, Lowell Thompson and Kara McGaughey for comments on the manuscript, and Michael Suplick for machine shop support (NIH National Eye Institute Core Grant P30 EY001583). This work was supported by NIH National Eye Institute (R01-EY022411; LD and JIG).

Author contributions

Conceptualization, LD and JIG; methodology, KR and LD; investigation, KR and LD; visualization, LD and JIG; funding acquisition, LD and JIG; project administration, LD; supervision, LD; writing – original draft, LD; writing – review & editing, KR, JIG, and LD.

Declaration of interests

The authors declare no competing financial interests.

Figure Legends

- Figure 1. Behavioral task and model predictions. A, Behavioral task. The monkey was
- required to report the perceived motion direction of the random-dot stimulus by making a
- saccade towards the corresponding choice target at a self-determined time. B, Three previous
- models predicted different patterns of STN activity. Sensitive to choice: differential responses for
- trials ending with different choices. Sensitive to uncertainty: differential responses for trials with
- 592 different evidence strength.
- Figure 2. STN neurons have diverse response profiles. A, Activity of three STN neurons
- (rows) aligned to motion (left) and saccade (right) onsets and grouped by choice x motion
- coherence (see legend). For motion-onset alignment, activity was truncated at 100 ms before
- saccade onset. For saccade-onset alignment, activity was truncated before 200 ms after motion
- onset. B, Summary of average activity patterns. Each row represents the activity of a neuron, z-
- scored by baseline activity in a 300 ms window before target onset and averaged across all trial
- conditions. Rows are grouped by monkey (red and green shown to the right of each panel:
- monkeys C and F, respectively) and sorted by the time of peak values relative to motion onset.
- Only correct trials were included. C, Heatmaps of linear regression coefficients for choice (top),
- 602 coherence for trials with contralateral choices (middle), and coherence for trials with ipsilateral
- choices (bottom), for activity aligned to motion (left) and saccade (right) onsets. Regression was
- performed in running windows of 300 ms. Regression coefficients that were not significantly
- different from zero (t-test, p>0.05) were set to zero (green) for display purposes. Neurons were
- sorted in rows by the time of peak coefficient magnitude. Only correct trials were included. D,
- Time courses of the fractions of regression coefficients that were significantly different from zero
- 608 (t-test, p<0.05), for choice (black), coherence for trials with contralateral choices (red), and
- 609 coherence for trials with ipsilateral choices (blue). Dashed line indicates chance level. E, Time
- 610 courses of the fractions of non-zero regression coefficients for coherence. Separate fractions
- were calculated for trials with the preferred (purple) and null (green) choices from choice-
- selectivity activity and for all trials from activity that was not choice selective (gray). Only time
- points after motion onset with fractions > 0.05 for choice-selective activity were included.
- Dashed line indicates chance level.
- Figure 3. STN contains distinct subpopulations. A, Three activity vectors that were
- constructed based on theoretical predictions in Figure 1B and used as seeds for k-means
- clustering (see Methods). B, Each panel shows the average activity of neurons in a cluster, same
- 618 format as Figure 2A. The numbers indicate the cluster size. C, Visualization of the clusters using
- the *t*-distributed stochastic neighbor embedding (*t*-SNE) dimension-reduction method. D,
- Average activity of clusters identified using random-seeded k-means clustering. Same format as
- Figure 3B. E, Visualization of the random-seeded clusters in the same tSNE space.
- 622 Figure 4. Clustering parameters. A. Silhouette plots for clustering results using different
- 623 combinations of settings. Silhouette scores for neurons are grouped by clusters and sorted. Red
- lines indicate the mean scores. Yellow shaded box indicates the chosen setting for results in
- Figure 3. B, Average Rand indices for different clustering settings. For each setting, the k-means
- algorithm was run 50 times, each time picking the best clusters out of 100 repetitions. Higher

- Rand index indicates greater cluster stability across different runs. Blue box indicates settings
- with Rand indices > 0.95. C, Mean silhouette scores and the number of negative scores as a
- function of number of clusters, using the firing rate vectors and correlation distance. Higher
- 630 mean score and fewer negative scores indicate better clustering.
- Figure 5. STN microstimulation affects monkeys' choice and RT. A-C, Monkey's choice (top)
- and RT (bottom) performance for trials with (red) and without (black) microstimulation for three
- example sessions (A,B: two sites in monkey C; C: monkey F). Lines: DDM fits. D, Distributions
- of microstimulation effects on bias and slope terms of the logistic function. Filled bars in
- 635 histograms indicate sessions with significant modulation of the specific term (bootstrap method).
- Triangles indicate the median values. Filled triangle: Wilcoxon sign-rank test for H_0 : median=0,
- 637 p < 0.05. E and F, Summary of microstimulation effects on the offset (E) and slope (F) terms of a
- 638 linear regression fit to RT data. Two separate linear regressions were performed for the two
- choices (Ipsi/Contra, as indicated). Triangles indicate the median values. Filled triangles:
- 640 Wilcoxon sign-rank test, p < 0.05.
- Figure 6. STN microstimulation affected multiple computational components in the DDM.
- A, Illustration of the DDM. Red/black lines represent across-trial mean/single-trial example of
- the evidence (top) and drift rate (bottom). Blue lines represent the collapsing decision bounds. B,
- Distribution of the difference in AIC between the None and Full models. Red dashed line
- indicates the criterion for choosing the full model: AIC difference = 3. C, Histograms of
- 646 microstimulation effects on DDM parameters. Each histogram included only sessions in which
- the Full model outperformed the corresponding reduced model (e.g., the histogram for parameter
- 648 *a* included only sessions in which $AIC_{NoA} AIC_{Full} > 3$ and $AIC_{None} AIC_{Full} > 3$). Triangles
- indicate median values. Filled triangles: Wilcoxon sign rank test, p < 0.05. D, Summary of
- 650 microstimulation effects on all parameters, for sessions in which at least one significant effect
- was present. Sessions were sorted by the prevalence and sign of the effects.
- Figure 7. Different STN subpopulations are intermingled. Locations of STN neurons, color-
- 653 coded by clusters based on random-seed clustering (same as Figure 3D). The Medial-Lateral
- 654 (ML) values were jittered for better visualization of neurons recorded along the same track and at
- similar depths. Anterior-Posterior (AP) levels were relative to the anterior commissure. ML and
- depth levels were relative to the center of the recording chambers.
- 657 Figure 8. STN microstimulation effects depend partially on neural clusters. A. Average
- activity at stimulation sites, grouped by four clusters based on the clusters in Figure 3D. B,
- Fractions of significant microstimulation effects for sites with the presence of each neuron
- cluster. Significance was based on AIC comparison between reduced and Full models. C,
- Microstimulation effects grouped by neuron cluster. Same format as Figure 5D. D, Fractions of
- significant microstimulation effects, grouped by effects reflecting changes in bound, decision
- variable computation, and non-decision processes.
 - **Supplemental Information**

- Suppl. Figure 1. STN activity is modulated by choice and RT. Same format as Figure 2,
- except using choice and RT as regressors.

- 667 Suppl. Figure 2. Summary of regression results, separated for different subpopulations.
- A: fractions of neurons in each category that showed significant modulation (t-test,
- p<0.05) at each time window by choice (top), coherence for trials with contralateral choices
- 670 (middle), and coherence for trials with ipsilateral choices (bottom). Dashed horizontal lines
- indicate the 5% chance level. Colors indicate categories as in Figure 3. B: median values
- of regression coefficients for choice and coherence as a function of time.
- 673 Suppl. Figure 3. Clustering results using alternative numbers of clusters, visualized in tSNE
- **space.** Same format as Figure 3E.
- 675 Suppl. Figure 4. Clustering results using alternative numbers of clusters, visualized as
- average firing rates for each cluster. Same format as Figure 3B and D.
- 677 Suppl. Figure 5. Comparison of different logistic models. A, The No Lapse model was
- associated with the lowest AIC for most sessions. The Symmetric Lapse model was associated
- with lower AICs for 12 sessions. The Asymmetric Lapse model was associated with lower AICs
- 680 for 8 sessions. B, Histograms of microstimulation effects on bias, slope, and lapse terms in the
- 681 Symmetric Lapse model. C, Histograms of microstimulation effects on bias, slope, and two lapse
- (for each choice) terms in the Asymmetric Lapse model. Same format as the histograms in Figure
- 683 5D.
- 684 **Suppl. Figure 6.** A, Differences in AIC between reduced and Full models. Filled circles indicate
- sessions for which $AIC_{Reduced} AIC_{Full} > 3$ (red line). Note that for three sessions, the Full model
- outperformed the None model but not any of the reduced models. B, Histograms of difference in
- DDM parameters between trials with and without microstimulation. Filled bars represent
- sessions considered to show significant microstimulation effects on the given parameter, based
- on AIC comparisons. Triangles indicate median values. Filled triangles: Wilcoxon sign-rank test,
- 690 p < 0.05.

- 691 Suppl. Figure 7. Indices of motivational state did not differ among sessions with different
- 692 **neuron subpopulations.** Panels show the summary of rate of fixation break (left), overall error
- rate (middle) and mean RT (right) for the four categories identified in Figure 3. All indices were
- 594 z-scored across sessions for each monkey. Red lines indicate median values. The bottom and top
- edges of the box indicate the 25th and 75th percentiles, respectively. The whiskers extend to the
- 696 most extreme data points not considered outliers, and the outliers are plotted individually as dots.
- ANOVA, p = 0.06, 0.91, and 0.29, respectively. No significant difference was observed for each
- 698 monkey separately (p > 0.08 and 0.12 for all indices for monkeys C and F, respectively).

Suppl Table 1. Indices of motivational state did not correlate with microstimulation effects.

P values are raw values from Pearson correlation, not corrected for multiple testing.

DDM components	Fixation Break		Error Rate		Mean RT	
	Correlation	p Value	Correlation	p Value	Correlation	p Value
B_d	-0.018	0.897	0.098	0.481	0.134	0.335
B_alpha	-0.070	0.615	0.075	0.591	-0.314	0.021
a	-0.049	0.723	-0.046	0.739	0.013	0.927
k	0.236	0.085	0.149	0.283	0.223	0.105
me	0.057	0.680	-0.027	0.844	0.035	0.802
z	-0.260	0.057	-0.031	0.822	0.041	0.768
T0_ipsi	0.205	0.138	0.049	0.724	0.166	0.231
T0_contra	0.060	0.669	-0.010	0.941	0.270	0.048

705 706

707

708

709

710 711

712

713

714

715

716

717

718

719 720

721

722

723

724

725

726 727

728

729

730

731

732

733

734

735 736 References Aron AR, Behrens TE, Smith S, Frank MJ, Poldrack RA (2007) Triangulating a cognitive control network using diffusion-weighted magnetic resonance imaging (MRI) and functional MRI. J Neurosci Off J Soc Neurosci 27:3743–3752. Aron AR, Poldrack RA (2006) Cortical and subcortical contributions to Stop signal response inhibition: role of the subthalamic nucleus. J Neurosci 26:2424–2433. Baunez C, Humby T, Eagle DM, Ryan LJ, Dunnett SB, Robbins TW (2001) Effects of STN lesions on simple vs choice reaction time tasks in the rat: preserved motor readiness, but impaired response selection. Eur J Neurosci 13:1609–1616. Bergman H, Wichmann T, DeLong MR (1990) Reversal of experimental parkinsonism by lesions of the subthalamic nucleus. Science 249:1436–1438. Bergman H, Wichmann T, Karmon B, DeLong MR (1994) The primate subthalamic nucleus. II. Neuronal activity in the MPTP model of parkinsonism. J Neurophysiol 72:507–520. Bogacz R, Gurney K (2007) The basal ganglia and cortex implement optimal decision making between alternative actions. Neural Comput 19:442–477. Brittain J-S, Watkins KE, Joundi RA, Ray NJ, Holland P, Green AL, Aziz TZ, Jenkinson N (2012) A role for the subthalamic nucleus in response inhibition during conflict. J Neurosci 32:13396-13401. Carpenter MB, Whittier JR, Mettler FA (1950) Analysis of choreoid hyperkinesia in the Rhesus monkey; surgical and pharmacological analysis of hyperkinesia resulting from lesions in the subthalamic nucleus of Luys. J Comp Neurol 92:293-331. Cavanagh JF, Wiecki TV, Cohen MX, Figueroa CM, Samanta J, Sherman SJ, Frank MJ (2011) Subthalamic nucleus stimulation reverses mediofrontal influence over decision threshold. Nat Neurosci 14:1462-1467. Cho S-H, Crapse T, Grimaldi P, Lau H, Basso MA (2021) Variable Statistical Structure of Neuronal Spike Trains in Monkey Superior Colliculus. J Neurosci 41:3234–3253. Coulthard EJ, Bogacz R, Javed S, Mooney LK, Murphy G, Keeley S, Whone AL (2012) Distinct roles of dopamine and subthalamic nucleus in learning and probabilistic decision making. Brain. Crapse TB, Lau H, Basso MA (2018) A Role for the Superior Colliculus in Decision Criteria. Neuron 97:181-194.e6. DeLong MR, Crutcher MD, Georgopoulos AP (1985) Primate globus pallidus and subthalamic nucleus: functional organization. J Neurophysiol 53:530–543.

DeLong MR, Wichmann T (2001) Deep brain stimulation for Parkinson's disease. Ann Neurol 49:142–143.

- 739 Desbonnet L, Temel Y, Visser-Vandewalle V, Blokland A, Hornikx V, Steinbusch HWM (2004)
- Premature responding following bilateral stimulation of the rat subthalamic nucleus is
- amplitude and frequency dependent. Brain Res 1008:198–204.
- Ding L, Gold JI (2010) Caudate encodes multiple computations for perceptual decisions. J
 Neurosci 30:15747–15759.
- Ding L, Gold JI (2012a) Neural correlates of perceptual decision making before, during, and after decision commitment in monkey frontal eye field. Cereb Cortex 22:1052–1067.
- Ding L, Gold JI (2012b) Separate, causal roles of the caudate in saccadic choice and execution in a perceptual decision task. Neuron 75:865–874.
- Doi T, Fan Y, Gold JI, Ding L (2020) The caudate nucleus contributes causally to decisions that balance reward and uncertain visual information. Elife 9:e56694.
- Fan Y, Doi T, Gold JI, Ding L (2024) Neural Representations of Post-Decision Accuracy and Reward Expectation in the Caudate Nucleus and Frontal Eye Field. J Neurosci 44.
- Fan Y, Gold JI, Ding L (2018) Ongoing, rational calibration of reward-driven perceptual biases. Elife 7:e36018.
- Fan Y, Gold JI, Ding L (2020) Frontal eye field and caudate neurons make different contributions to reward-biased perceptual decisions. eLife 9:e60535.
- Frank MJ (2006) Hold your horses: a dynamic computational role for the subthalamic nucleus in decision making. Neural Netw 19:1120–1136.
- Frank MJ, Samanta J, Moustafa AA, Sherman SJ (2007) Hold your horses: impulsivity, deep brain stimulation, and medication in parkinsonism. Science 318:1309–1312.
- Fumagalli M, Giannicola G, Rosa M, Marceglia S, Lucchiari C, Mrakic-Sposta S, Servello D,
- Pacchetti C, Porta M, Sassi M, Zangaglia R, Franzini A, Albanese A, Romito L,
- Piacentini S, Zago S, Pravettoni G, Barbieri S, Priori A (2011) Conflict-dependent
- dynamic of subthalamic nucleus oscillations during moral decisions. Soc Neurosci
- 764 6:243–256.
- Gold JI, Shadlen MN (2007) The neural basis of decision making. Annu Rev Neurosci 30:535–574.
- Green N, Bogacz R, Huebl J, Beyer AK, Kuhn AA, Heekeren HR (2013) Reduction of influence
- of task difficulty on perceptual decision making by STN deep brain stimulation. Curr
- 769 Biol 23:1681–1684.

Hanks TD, Ditterich J, Shadlen MN (2006) Microstimulation of macaque area LIP affects decision-making in a motion discrimination task. Nat Neurosci 9:682–689.

- Herz DM, Tan H, Brittain JS, Fischer P, Cheeran B, Green AL, FitzGerald J, Aziz TZ, Ashkan K,
- Little S, Foltynie T, Limousin P, Zrinzo L, Bogacz R, Brown P (2017) Distinct
- mechanisms mediate speed-accuracy adjustments in cortico-subthalamic networks. Elife
- 775 6.
- Herz DM, Zavala BA, Bogacz R, Brown P (2016) Neural Correlates of Decision Thresholds in the Human Subthalamic Nucleus. Curr Biol 26:916–920.
- Hikosaka O, Wurtz RH (1983) Visual and oculomotor functions of monkey substantia nigra pars
- reticulata. III. Memory-contingent visual and saccade responses. J Neurophysiol
- 780 49:1268–1284.
- Isoda M, Hikosaka O (2008) Role for subthalamic nucleus neurons in switching from automatic to controlled eye movement. J Neurosci 28:7209–7218.
- Jeurissen D, Shushruth S, El-Shamayleh Y, Horwitz GD, Shadlen MN (2022) Deficits in
- decision-making induced by parietal cortex inactivation are compensated at two
- 785 timescales. Neuron 110:1924-1931.e5.
- Jun E, Bautista A, Nunez M, Allen D, Tak J, Alvarez E, Basso M (2021) Causal role for the
- primate superior colliculus in the computation of evidence for perceptual decisions. Nat
- 788 Neurosci 24:1121–1131.
- Lehericy S, Ducros M, Krainik A, Francois C, Van de Moortele PF, Ugurbil K, Kim DS (2004)
- 790 3-D diffusion tensor axonal tracking shows distinct SMA and pre-SMA projections to the
- human striatum. Cereb Cortex 14:1302–1309.
- Lo CC, Wang XJ (2006) Cortico-basal ganglia circuit mechanism for a decision threshold in reaction time tasks. Nat Neurosci 9:956–963.
- Ma TP (1996) Saccade-related omnivectoral pause neurons in the primate zona incerta.
- 795 Neuroreport 7:2713–2716.
- Martin JP (1927) Hemichorea resulting from a local lesion of the brain. (The syndrome of the body of Luys.). Brain 50:637–649.
- Martin JP, Alcock NS (1934) Hemichorea associated with a lesion of the corpus Luysii. Brain 57:504–516.
- Matsumura M, Kojima J, Gardiner TW, Hikosaka O (1992) Visual and oculomotor functions of monkey subthalamic nucleus. J Neurophysiol 67:1615–1632.
- Meister ML, Hennig JA, Huk AC (2013) Signal multiplexing and single-neuron computations in lateral intraparietal area during decision-making. J Neurosci 33:2254–2267.

805

806 807

808

809

810

811

812

813

814

815

816

817

818

819

820

821

822

823

824

825

826

827

828

829

830

831

832

833

834

835 836

837

838

Nambu A, Takada M, Inase M, Tokuno H (1996) Dual somatotopical representations in the primate subthalamic nucleus: evidence for ordered but reversed body-map transformations from the primary motor cortex and the supplementary motor area. J Neurosci 16:2671–2683. Pagnier GJ, Asaad WF, Frank MJ (2024) Double dissociation of dopamine and subthalamic nucleus stimulation on effortful cost/benefit decision making. Curr Biol CB 34:655-660.e3. Parent A, Hazrati LN (1995) Functional anatomy of the basal ganglia. II. The place of subthalamic nucleus and external pallidum in basal ganglia circuitry. Brain Res Brain Res Rev 20:128-154. Pasquereau B, Turner RS (2017) A selective role for ventromedial subthalamic nucleus in inhibitory control. Elife 6. Pote I, Torkamani M, Kefalopoulou ZM, Zrinzo L, Limousin-Dowsey P, Foltynie T, Speekenbrink M, Jahanshahi M (2016) Subthalamic nucleus deep brain stimulation induces impulsive action when patients with Parkinson's disease act under speed pressure. Exp Brain Res 234:1837–1848. Prasad AA, Wallén-Mackenzie Å (2024) Architecture of the subthalamic nucleus. Commun Biol 7:1–14. Rand WM (1971) Objective Criteria for the Evaluation of Clustering Methods. J Am Stat Assoc 66:846-850. Ratcliff R, Frank MJ (2012) Reinforcement-based decision making in corticostriatal circuits: mutual constraints by neurocomputational and diffusion models. Neural Comput 24:1186–1229. Redgrave P, Prescott TJ, Gurney K (1999) The basal ganglia: a vertebrate solution to the selection problem? Neuroscience 89:1009–1023. Roitman JD, Shadlen MN (2002) Response of neurons in the lateral intraparietal area during a combined visual discrimination reaction time task. J Neurosci 22:9475–9489. Saleem KS, Logothetis N (2007) A combined MRI and histology atlas of the rhesus monkey brain in stereotaxic coordinates. London; Burlington, MA: Academic. Schmidt R, Leventhal DK, Mallet N, Chen F, Berke JD (2013) Canceling actions involves a race between basal ganglia pathways. Nat Neurosci 16:1118–1124. Stine GM, Trautmann EM, Jeurissen D, Shadlen MN (2023) A neural mechanism for terminating decisions. Neuron 111:2601-2613.e5. Tehovnik EJ (1996) Electrical stimulation of neural tissue to evoke behavioral responses. J Neurosci Methods 65:1–17.

840

841

842

843

844

845

846

847

848

849

850 851

852

853

854

855

856

857

858 859

860

861

Wei W, Rubin JE, Wang XJ (2015) Role of the indirect pathway of the basal ganglia in perceptual decision making. J Neurosci 35:4052-4064. Whittier JR, Mettler FA (1949) Studies on the subthalamus of the rhesus monkey; hyperkinesia and other physiologic effects of subthalamic lesions; with special reference to the subthalamic nucleus of Luys. J Comp Neurol 90:319–372. Wichmann T, Bergman H, DeLong MR (1994a) The primate subthalamic nucleus. III. Changes in motor behavior and neuronal activity in the internal pallidum induced by subthalamic inactivation in the MPTP model of parkinsonism. J Neurophysiol 72:521–530. Wichmann T, Bergman H, DeLong MR (1994b) The primate subthalamic nucleus. I. Functional properties in intact animals. J Neurophysiol 72:494–506. Witt K, Pulkowski U, Herzog J, Lorenz D, Hamel W, Deuschl G, Krack P (2004) Deep brain stimulation of the subthalamic nucleus improves cognitive flexibility but impairs response inhibition in Parkinson disease. Arch Neurol 61:697–700. Zaghloul KA, Weidemann CT, Lega BC, Jaggi JL, Baltuch GH, Kahana MJ (2012) Neuronal Activity in the Human Subthalamic Nucleus Encodes Decision Conflict during Action Selection. J Neurosci 32:2453-2460. Zavala B, Damera S, Dong JW, Lungu C, Brown P, Zaghloul KA (2017) Human Subthalamic Nucleus Theta and Beta Oscillations Entrain Neuronal Firing During Sensorimotor Conflict. Cereb Cortex 27:496-508. Zavala BA, Tan H, Little S, Ashkan K, Hariz M, Foltynie T, Zrinzo L, Zaghloul KA, Brown P (2014) Midline Frontal Cortex Low-Frequency Activity Drives Subthalamic Nucleus Oscillations during Conflict. J Neurosci 34:7322.