Widespread temporal coding of cognitive control in the human prefrontal cortex

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When making decisions we often face the need to adjudicate between conflicting strategies or courses of action. Our ability to understand the neuronal processes underlying conflict processing is limited on the one hand by the spatiotemporal resolution of functional MRI and, on the other hand, by imperfect cross-species homologies in animal model systems. Here we examine the responses of single neurons and local field potentials in human neurosurgical patients in two prefrontal regions critical to controlled decision-making, the dorsal anterior cingulate cortex (dACC) and dorsolateral prefrontal cortex (dlPFC). While we observe typical modest conflict-related firing rate effects, we find a widespread effect of conflict on spike-phase coupling in the dACC and on driving spike-field coherence in the dlPFC. These results support the hypothesis that a cross-areal rhythmic neuronal coordination is intrinsic to cognitive control in response to conflict, and provide new evidence to support the hypothesis that conflict processing involves modulation of the dlPFC by the dACC.

As we navigate through our daily lives, we are often confronted with choices involving competing or conflicting potential courses of action. To resolve this cognitive conflict, we must summon additional cognitive resources—that is, we must both monitor and resolve the conflict. For example, imagine driving through a busy city center area and seeing the next traffic light turn green—normally a signal to go—but a slow-moving pedestrian is on the pedestrian crossing. To make the optimal decision as to whether to brake or swerve, the driver must summon additional cognitive resources to assess the pedestrian’s speed, estimate the trajectory of the car, and determine whether other cars are following closely behind. This capacity to monitor conflict and implement control is a vital, yet poorly understood, element of the repertoire of flexibly intelligent organisms. Understanding its mechanisms is essential for developing treatments for neuropsychiatric disorders associated with impaired conflict processing, and, since conflict is a useful model for other types of executive control, in general.

Much research has implicated the dorsal prefrontal cortex, especially the dorsal anterior cingulate cortex (dACC) and dorsolateral prefrontal cortex (dlPFC), in the monitoring and resolution of cognitive conflict. For example, a large amount of imaging and electrophysiology data from humans and non-human animals supports the hypothesis that these regions play a critical role in one or both processes. Moreover, lesions to these regions lead to a variety of impairments in control functions. Prominent computational and theoretical models ascribe to these regions several features that are critical to a control process: monitoring goal-relevant variables, evaluating relative costs and benefits of choice alternatives, maintaining rule sets and goal-related information in working memory, and producing adaptive biases towards more successful behavior. This work dovetails with careful computational studies on how systems with limited cognitive capacity can trade off between multiple competing sources of guidance. That work in turn results in specific circuit-level predictions about how cognitive control can be implemented in response to conflict. In particular, the study suggests that the dACC serves as a conflict monitor and that the dlPFC, located downstream of the dACC, serves to implement conflict-related changes. The idea that these two regions are both involved in conflict processing but have distinct roles has been difficult to test given the paucity of studies that directly compare the activities of these two regions using neuron-level recording methods.

On the one hand, human neuroimaging studies find clear correlates of conflict but lack the temporal resolution to identify circuit-level correlates. On the other hand, primate neurophysiology studies, which have high temporal and spatial resolution, have generally failed to find clear correlates of cognitive conflict. Rodents do not have a clear homolog of either the dACC or dlPFC, making interpretation of data from rodent studies challenging. The reasons for the disconnect between non-human primate electrode recording studies and human neuroimaging are not clear. One reason may be that animal studies require a very large number of trials; the resulting overtraining may systematically alter conflict processing. Another likely factor is that functional magnetic resonance imaging (fMRI) and single unit activity (SUA) do not have clear one-to-one mapping, so there may be other features of brain responses that predict conflict other than average firing rate responses.

Control, like many other aspects of executive function, requires coordination of activity across regions. This has led to the hypothesis that the major signatures of control would be observed in relation to the local field potential (LFP), which is thought to provide a mechanism for coordination across multiple brain structures. LFP is distinct from unit activity and seems to be a better predictor of...
hemodynamic responses than average unit firing in some cases\textsuperscript{28,29}. Furthermore, an extensive electroencephalography (EEG) literature provides reason to believe that oscillations contain information critically related to conflict processing. For example, variations in theta-range (4–8 Hz) oscillations over frontal EEG contacts, known as frontal midline theta, have repeatedly been shown to reflect conflict and error processing, attentional control, and reinforcement-learning signals\textsuperscript{30,31}. Such findings have been extended and refined with the few existing studies using human intracranial EEG\textsuperscript{32,33}. This literature has led to an increasing number of theories positioning a fundamental role for oscillatory activity in decision-making and cognition more generally\textsuperscript{34,35}. An increasingly substantial body of evidence has demonstrated the importance of neuronal spiking activity as it relates to the background LFP. This alignment of spiking with LFPs, or spike-field coherence (SFC), is a potent channel for coordination of neuronal populations, both locally and across brain regions\textsuperscript{36,37}. For these reasons, our analyses focused on the SFC within the dACC and dlPFC.

Here we examined simultaneously collected single-neuron responses and LFPs in the dACC and dlPFC in human neurosurgical patients. We found modest neuronal firing rate correlates of conflict in the dACC and negligible ones in the dlPFC. By contrast, we found robust effects of conflict on the preferred phase of spiking in the dACC (but not in the dlPFC), but greater conflict-related SFC changes in the dlPFC than in the dACC. These results support the hypothesis that processing cognitive conflict requires cross-areal coordination mediated, at least in part, by oscillatory activity. Moreover, they are consistent with the hypothesis that conflict processing involves monitoring by the dACC and control by the dlPFC\textsuperscript{38}. Finally, our results show how neural codes that combine information in the spiking and oscillatory frequency (LFP) domains can facilitate the types of coordinated computation needed for executive control and provide important constraints on computational models of conflict processing.

Results
Limited, mixed firing rate code for conflict in the dACC. We simultaneously recorded firing rates of single neurons in the dACC, and LFPs in both the dACC and dlPFC, in six human patients with medically refractory epilepsy undergoing intracranial recordings to detect seizures (Fig. 1a and Supplementary Tables 1–3). Participants performed the multi-source interference task (MSIT), a task that independently manipulates both spatial (Simon) and flanker (Eriksen) types of decision conflict (Fig. 1b). Participants had an overall low error rate (average of 1.4 ± 2.7% per session) and exhibited the hallmark behavioral signature of tasks demanding cognitive control: increasing reaction time with increasing level of decision conflict (Fig. 1c).

We recorded firing rates from 136 well-isolated dACC neurons (Fig. 2a). To classify neurons on the basis of firing rate coding of task-relevant variables, we used a sliding-window generalized linear model (GLM) incorporating three factors corresponding to the three main task variables: decision conflict, response identity, and feedback valence. We defined decision conflict as the sum of the two binary variables corresponding to the two forms of conflict.
We focused our analysis on the period after cue presentation and before the behavioral response. We therefore chose a time window 250–750 ms after cue presentation for the subsequent analyses. In our sample, 10.3% of dACC neurons (n = 14 of 136) were selective for decision conflict (Fig. 2b), 8.8% (n = 12 of 136) were selective for response identity (Supplementary Fig. 1a), and 8.1% (n = 11 of 136) were selective for feedback valence (Supplementary Fig. 1b). There was very little overlap among the different neuron classes, suggesting that these forms of conflict are encoded in largely disjointed sets of cells (Fig. 2c). The majority of dACC neurons recorded (72.8%, n = 99 of 136) did not exhibit selectivity for any of the task variables based on firing rate. Nonetheless, it is clear that some human dACC neurons do use a detectable firing rate-coding scheme for task-relevant features, although these neurons are fairly uncommon.

**Temporal code for decision conflict in the dACC.** We next tested the hypothesis that PFC neurons use a temporal-coding strategy in the context of controlled decision-making. Of the three main task features, we focused our analysis on decision conflict, as this was the primary task manipulation (analyses of other features are presented in Supplementary Fig. 2). Specifically, we hypothesized that temporal patterns of neuronal spiking relate to population oscillatory activity, and that this relationship is modulated in a conflict level-dependent manner.

We measured the SFC for each neuron in relation to the LFP in the dACC. We observed significant increases in the SFC in the beta and theta ranges. These findings indicate that dACC spike timing could be predicted from population-level oscillations in these two frequency ranges (henceforth we refer to neurons exhibiting this property as beta- and theta-coherent, respectively, Fig. 3). Beta-coherent neurons (n = 50, 36.8%) showed significant increases in coherence in the frequency range between 16.1 and 24.4 Hz (SFC permutation test, P < 0.05). Theta-coherent neurons (n = 43, 31.6%) showed cue-evoked changes in coherence between 2.9 and 9.2 Hz (SFC permutation test, P < 0.05). The observed SFC results were not driven simply by variations in LFP power (Supplementary Fig. 3).

Moreover, we observed a prominent phase code for decision conflict in these temporally coherent neurons. We measured the mean phase at which neurons fired in relation to the population LFP for neurons with a significance cluster that overlapped with the time window used to assess firing rate selectivity. We found a clear pattern of dependence on the amount of decision conflict in both theta- and beta-coherent neurons (Fig. 3c,g). Neurons generally fired before the LFP trough during high-conflict trials (that is, theta-coherent neurons fired before the theta trough and beta-coherent neurons fired before the beta trough) and after the LFP trough during low-conflict trials (Fig. 3d,h). The number of neurons that showed a phase-specific temporal code for decision conflict was larger than the number that showed a conventional rate code (McNemar’s Test, P < 10^{-3}, \chi^2 = 13.5, Fig. 3i). Furthermore, the phase code and rate code were largely independent from each other: rate-coding neurons seldom showed a phase code, and vice versa (correlation between firing rate code and theta temporal code: \rho = 0.03, P = 0.76; correlation between firing rate code and beta temporal code \rho = -0.02, P = 0.74; Fig. 3j,k). These results show robust and widespread temporal coding of decision conflict in the dACC, particularly evident as a phase code, and independent of a co-existing and less prevalent rate code.

**Relationship between dACC spiking activity and broader network LFP activity.** We next addressed the question of how these largely distinct neuronal populations with different information-encoding strategies differentially participate in the implementation of controlled decisions. To do so, we examined their interactions with the broader network that has previously been implicated in cognitive control, which includes the dorsolateral prefrontal cortex (dlPFC). We examined the spike-triggered LFP (stLFP) of dACC neurons to determine the relationship between dACC spiking activity and the local dACC LFP as well as the distant dlPFC LFP (for example, Fig. 1a, recording locations for each patient are shown in Supplementary Fig. 4).

The average stLFP of dACC spikes with dACC LFP (dACC–dACC stLFP) showed a consistent deflection starting immediately after the spike and becoming maximally negative around 100 ms post-spike (Fig. 4, all neuron categories shown in Supplementary Fig. 5). The stLFP amplitude for the rate-coding neurons was significantly greater than that for the beta- or theta-coherent neurons.
which in turn was significantly greater than that for the non-coding neurons and null distribution (Fig. 4a–c).

Spiking activity in the dACC also interacted with LFP in the dlPFC. SFC analysis between dACC spikes and dlPFC LFPs again revealed prominent temporal coding, with both beta- and theta-coding neuronal populations (Supplementary Fig. 6). dACC spikes were also associated with significant deflections in dlPFC LFP (dACC–dlPFC stLFP) (Fig. 4d,e). These stLFPs followed a similar pattern to the dACC–dACC stLFPs, with the greatest stLFP in rate-coding neurons, a lesser but still significant stLFP in beta and theta phase-coding neurons, and non-significant stLFPs in non-coding neurons (Fig. 4f). More dACC units were coherent with dlPFC theta (49 units, 36.8%) and coherent with dlPFC beta (43 units, 31.6%)
Fig. 4 | dACC neuronal interactions within a broader control network. a, stLFP waveforms evoked by dACC neurons on dACC LFP (dACC–dACC) for conflict-selective rate-coding neurons (pink, n = 2,370 spikes), non-coding neurons (gray, n = 10,292 spikes), and null distribution (black). b, dACC–dACC stLFP waveforms for temporal-coding neurons: beta-coherent (green, n = 8,398 spikes), theta-coherent (blue, n = 7,581 spikes), and null distribution (black). c, Distributions of dACC–dACC stLFP amplitudes for each neuron group in a and b. dACC–dACC stLFP amplitudes were decorrelated by their covariance matrices and z-scored. d–f, stLFP waveforms evoked by dACC neurons on dlPFC LFP (dACC–dlPFC), dACC–dlPFC waveforms for temporal-coding neurons: beta-coherent (green, n = 2,370 spikes), non-coding neurons (gray, n = 2,370 spikes), conflict-selective rate-coding neurons (pink, n = 750 ms after cue presentation) as in the dACC analysis above. Using the same sliding GLM approach to test for firing rate selectivity, we found that a small proportion of neurons encoded conflict (n = 15, 4.1%). This proportion was not significantly greater than chance (exact test; P > 0.05) and, not surprisingly, was significantly smaller than the proportion found in the dACC (χ² test, χ² = 8.2, P = 4.3 × 10⁻³). Additionally, 18 neurons were selective for response identity (4.9%, Supplementary Fig. 8a) and 24 were selective for feedback valence (6.5%, Supplementary Fig. 8b, Fig. 5b).

SFC analysis revealed prominent theta-range SFC in the majority of recorded neurons (n = 238, 64.9%). The strength of coherence between spike timing and theta oscillations increased with higher levels of conflict (Fig. 5d–f). This effect was prominent enough to be visible across the entire population of recorded neurons (Fig. 5g) and was significant at the individual neuron level in the majority of cells (n = 191, 52.0%). Furthermore, we observed this pattern in neurons from each of the nine participants (mean ± standard deviation = 67.3% ± 19.9% of neurons from each participant). In contrast to its prominence in the dlPFC, only a small number of dACC neurons exhibited similar conflict-modulated scaling in SFC amplitude (Fisher’s exact test, P < 10⁻³). By contrast, the prominent phase-coding motif evident in the dACC was not apparent in the dlPFC. We found a slight precession of the SFC phase in all conditions, with similar error across conditions, yet no significant differences among phases for each condition. This means that the mean phase does not differ among conflict conditions, although the amplitude of SFC increases (Supplementary Fig. 9c). Thus, we observed a similar general theme of uncommon rate-coding but robust temporal coding in the dlPFC, but the specific temporal-coding schemes differed between dACC and dlPFC neurons. The overall thematic similarities demonstrate that the temporal-coding archetype is not unique to the dACC and suggest that temporal coding may be a conserved strategy across the PFC. This finding also underscores the value of asking not only whether certain types of information (for example, conflict³) are encoded in different PFC regions, but also how they are encoded.

Trial-to-trial variation in temporal coding and its relationship to behavior. Neural information processing should be stable against
noise yet flexible enough to meet unforeseen changes in cognitive demands, whether driven by external conditions or by internal fluctuations in arousal, attention, or goals. Because these demands vary on a moment-to-moment basis, their trial-to-trial encoding should be detectable. We reasoned that a temporal-coding scheme, distributed over a population of neurons, may be able to effectively support such an on-demand, within-trial implementation of controlled responses.

Using sessions with more than 100 simultaneously recorded neurons, we considered whether a distributed population of neurons can encode a particular instance of conflict as a coherent population. We examined SFC among all simultaneously recorded dlPFC neurons during each trial and again found significant spike-theta coherence that increased in trials with higher levels of conflict (Fig. 6a). Measuring SFC at the single-trial level also afforded the opportunity to examine whether trial-to-trial variation in SFC can account for trial-to-trial variation in behavior. This analysis revealed that the duration of theta coherence predicted reaction time in each trial (linear mixed model (LMM), \( t_{\text{res}} = 2.9, P = 0.002 \)). Furthermore, higher mean theta coherence across neurons during a given trial strongly predicted shorter reaction times in that trial, even after controlling for conflict level in the current trial, in the previous trial, and their interaction (Fig. 6b, theta coherence: \( t_{\text{res}} = -4.22, P = 2 \times 10^{-5} \)). To control for any differences between LFP recorded from surface ECoG, and intraparenchymally recorded LFP, we examined within-trial SFC using down-sampled and filtered LFP across the UMA between 1 and 50 Hz. Similar within-trial SFC effects were observed using intraparenchymally recorded LFP (Fig. 6c, LMM, \( t_{\text{res}} = -3.9, P = 9.5 \times 10^{-5} \)). Despite their well-established relationship to reaction time, all other effects in the model (current and previous trial conflict level and their interaction) were substantially weaker than the theta-coherence effect on reaction time (\( t_{\text{res}} \leq 1.99 \), \( 0.065 > P > 0.045 \)). Higher mean coherence was unrelated to conflict level in the previous trial \( (P = 0.24) \), also supporting the idea that the relationship between mean theta coherence and reaction times was not driven by behavioral adaptations due to previous-trial effects. These results show that a temporal code distributed across a population of dlPFC neurons closely tracks moment-to-moment fluctuations in performance, including and beyond those imposed by the task, potentially supporting a mechanism that permits flexible adjustments to cognitive demands.

Discussion

We examined the responses of single neurons and LFP in two brain regions, the dACC and dlPFC, in human neurosurgical patients performing a conflict task. We found a small but significant population of neurons in the dACC (but not the dlPFC) with changes in firing rate that encode task conflict. Our previous report demonstrated conflict encoding at the level of a population of human dACC neurons, but our current observation of explicit conflict...
encoding at the level of individual neurons has not yet been reported in humans. Our new finding is important because it is inconsistent with the idea that ostensible conflict signals in mass action measures (such as BOLD) are an epiphenomenon. The firing rate encoding of conflict in these neurons was most apparent before the decision occurred, suggesting that it may reflect conflict monitoring in the service of on-line control, rather than learning or trial-to-trial adjustment. More prominently, we found more widespread population encoding of conflict in the form of modulation of spike timing relative to ongoing LFPs. That is, a robust temporal code for decision conflict complements the more limited rate code. This temporal code seems to be independent of the spike code. Specifically, we observed this temporal code in neurons that did not exhibit a significant firing rate code, and the magnitudes of the two codes were independent across neurons.

An emerging concept bridging these perspectives is that the relationship between neuronal spiking and ensemble oscillatory activity is a critical feature that the brain uses to encode complex functional representations. Perhaps the most celebrated example of spike-oscillation synchrony is the theta phase precession observed in rat hippocampal and entorhinal place cells. Similar temporal codes underlie the encoding of complex visual stimuli in the primary visual cortex, acoustic stimuli in the auditory cortex, and coordination of gaze and reaching movements in the parietal cortex. It seems that there are similar neural representations of cognitive variables and spatial maps in the human ACC and mesial temporal lobe, respectively, and that both are linked to theta rhythmicity and phase coding. Recent theoretical work has proposed a dynamical control mechanism for phase coding, thus providing explanatory power and opportunities for testing specific hypotheses. These factors, together with the wide-ranging interconnected nature of the medial PFC, suggest that temporal coding in the medial PFC could both be a prominent coding scheme, and support a dynamical, distributed mechanism to influence a diverse array of brain areas.

Some previous work suggests that the dACC may integrate control-related variables into a general control signal and that the dlPFC may implement their effects; that is, the dACC may be a monitor and the dlPFC may be a controller. Our results are broadly consistent with that proposed division of labor. First, in contrast to the dACC, we found a minimal effect of conflict on firing rate changes in dlPFC neurons, as if the dlPFC does not need to signal conflict per se. We did find a strong oscillatory code for conflict in the dlPFC, although it differed qualitatively from the one observed in the dACC: it was an increase in SFC without a concurrent alignment to phase such as that observed in the dACC. Oscillatory dynamic processes expand the coding space to allow optimized information formatting and consequently facilitate the sparsening of downstream representations, while simultaneously conferring stability of representations in the presence of noise. From this perspective, a natural interpretation is that the dACC plays an upstream function, closer to the input (that is, the source of conflict), such as a detection or signaling role. Indeed, theoretical work suggests that impulses occurring at earlier phases of the cycle exert more control on the system. That this phase relationship was not apparent in the dlPFC is consistent with the idea that it is functionally downstream of the dACC. In the same vein, theta-SFC coding in the dlPFC suggests its involvement in the downstream implementation of controlled behavior, particularly given that theta coherence is strongly correlated with faster responses even after controlling for external, task-derived demands, such as the amount of conflict in the current or previous trial cue. This finding accords with the idea that spike-theta coherence in the dlPFC may be a ‘lingua franca’ to support the implementation of controlled decision-making strategies, adjusting to the presence of sources of interference regardless of whether they are external (for example, the task or environment) or internal (for example, spontaneous attentional fluctuations).

A substantial amount of research examining the neural bases of conflict detection and resolution has been conducted in non-human animals. Classically, much of this work reported no such correlates and suggested that ostensible conflict coding may be an epiphenomenon. By contrast, at least two recent studies suggest that frank encoding of conflict may be observed in single neurons in non-human animals. How can these results be reconciled? Our work suggests that two factors may be at play. First, the conflict coding observed in the dACC is greatest in the temporal, phase-locked domain and is much weaker in the rate domain, meaning that it may have been difficult to detect in some studies. Indeed, the hemodynamic correlates of conflict may be more strongly linked to the oscillatory patterns we observed than to the more modest rate code. Second, the weakness of the signal in animals may be compounded by the need for overtraining, which makes controlled behavior more automatized and may reduce the size of any neural signals. This effect would explain why firing rate effects seem to be larger in humans than in animals. If so, this example illustrates one of the
major benefits of human intracranial recordings; that they allow for studies of rapidly and flexibly learned behaviors\(^9\).

Predictions from this proposed functional specialization model can be tested with future work, including simultaneous recordings of larger neuronal populations in the dACC to measure the temporal structure of population representations, or closed-loop, single-pulse electrical stimulation at particular LFP phases to dynamically modulate cognitive control. Finally, this idea of functional specialization is not meant to imply a strict functional segregation between the dACC and dIPFC. The overall use of sparse rate-coding and widespread temporal-coding schemes highlights the need for an integrated understanding of the role of various neural coding strategies across prefrontal networks\(^10\).

**Conclusion: the soloists and the choir.** Perhaps the most intriguing finding in our study is that the conflict rate-coding neurons, despite their low numbers, were associated with a disproportionately large relationship with LFP throughout the prefrontal network. We propose that this rate-coding minority may serve a specialized function of being particularly sensitive in detecting and signaling changes in conflict or, possibly, in demand for control. In other words, conflict-sensitive neurons may function as specialized ‘soloists’ in the medial PFC\(^4\). These soloists may serve as an early signal for the need to establish control, and then catalyze more widespread oscillatory activity throughout the network\(^4\). The larger population of temporal-coding neurons may then act as a ‘choir’ that stabilizes and amplifies the representation of task-relevant information. If this speculation is correct, then the soloist neurons may serve a local, intra-areal purpose, and the resulting oscillatory activity may serve to coordinate cross-regional coherent responses. This second form of responding would presumably be more robust to noise and more sensitive to regulation by top-down factors. This multi-faceted approach to representing relevant information may facilitate its efficient use and communication across neuronal systems.

**Online content**
Any methods, additional references, Nature Research reporting summaries, source data, statements of code and data availability and associated accession codes are available at [https://doi.org/10.1038/s41593-019-0494-0](https://doi.org/10.1038/s41593-019-0494-0).

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Author contributions

S.A.S., C.B.M., M.J.Y., and E.H.S. designed the experiments. E.H.S., C.B.M., M.J.Y., Y.J.P., C.A.S., G.M.M., and S.A.S were involved with collecting the data. E.H.S., G.P.B., and G.H. analyzed the data. E.H.S., S.A.S., G.H., B.Y.H., M.M.B., and M.J.Y., wrote the manuscript. All authors provided edits to the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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Methods

Participants and ethics statement. Two cohorts of individuals participated in this research. One (Cohort 1, Supplementary Table 2) was a cohort of six patients (one female) with medically refractory epilepsy who were undergoing intracranial monitoring to identify seizure onset regions. These patients had been implanted with stereo-electroencephalography (sEEG) depth electrodes using standard stereotactic techniques. One or more of the sEEG electrodes in this cohort spanned the dIPFC to the dACC (Brodman’s areas 24, a, b, and c, and 32), providing LFP recordings from both regions, as well as single unit recordings in the dACC (see Data acquisition and preprocessing below).

The other cohort (Cohort 2, Supplementary Table 3) was a cohort of nine patients: eight (two female) with movement disorders (Parkinson’s disease or essential tremor) who were undergoing DBS surgery, and one male patient with epilepsy undergoing intracranial seizure monitoring. The entry point for the trajectory of the DBS electrode is typically in the inferior portion of the superior frontal gyrus, or supra-pontine portion of the middle frontal gyrus, within 2 cm of the coronal suture. This area corresponds to the dIPFC (Brodman’s areas 9 and 46).

Behavioral Task. All participants performed the MSST (Fig. 1b)\textsuperscript{\textsuperscript{19}}, in which each trial consisted of a 500 ms fixation period, followed by a cue consisting of three integers ranging from 0 to 3. One of these three numbers (the ‘target’) was different from the other two numbers (the ‘distractors’). Participants were instructed to indicate the identity of the ‘target’ on a button pad, on which each button represented the integers 1 (left-hand button), 2 (middle button), and 3 (right-hand button). This task therefore presented two types of conflict: spatial conflict if the target was in a different position in the cue than on the three-button pad (that is, ‘0 0 1’; the target is in the right-hand position, but the left-hand button is the correct choice), and distractor conflict if the distractor numbers were possible button choices (for example ‘3 2 3’, in which 3 corresponds to a possible button choice, versus ‘0 2 0’, in which 0 does not correspond to a possible button choice). After each participant registered his or her response, the cue disappeared and feedback appeared, consisting of the target number in a different color, with a variable delay of 300–800 ms. Valenced feedback (green for correct, red for incorrect) alternated with neutral feedback (blue regardless of correctness) in blocks of ten trials. The inter-trial interval varied uniformly randomly between 1 and 1.5 seconds.

The task was presented on a computer monitor controlled by the Psychophysics Matlab Toolbox (www.psyctoolbox.org, The MathWorks; Matlab v. 2016b–2018b). This software integrates visual and data acquisition functionality (National Instruments) that allowed for synchronization of behavioral events and neural data with sub-millisecond precision. Differences in log reaction times over conflict conditions were tested using mixed-effects models (LMM): $y = X\beta + Zu + \epsilon$

where $y$ represents reaction times, $\beta$ a represents a design matrix of conflict condition factors, $\beta$ represents the fixed-effects regression coefficients, $Z$ represents the random-effects design matrix accounting for random participant-session reaction times, $u$ represents the random component of the fixed effects in $\beta$, and $\epsilon$ represents the model residuals, or, in Wilkinson notation, reaction time $-$1 conflict condition $\times$ session +1 (1 conflict condition $\times$ session $\times$ participant). Reaction times were lognormally distributed, so we modeled the log of reaction time to meet assumptions of normality. These models were fit using maximum likelihood methods using the Matlab function fitgmdist. Subsequent linear models are specified using Wilkinson notation.

Statistics. As described in the previous section, many of the hypotheses were tested with LMMs, the effects of which were evaluated with t-tests and ANOVAs. All statistical tests performed are listed in Supplementary Table 4. For tests that assumed normality, a priori Lilliefors tests were performed to confirm that data were normally distributed. Sample sizes were based on numbers of trials and neurons, both of which were randomized. The number of trials was determined by the number of datasets and the number of neurons in each dataset. The number of neurons for each trial was randomly assigned from a uniform distribution. The number of neurons was based on where microelectrodes ended up recording, approximately 4 mm away from the nearest macroelectrode, which was placed on the basis of clinical parameters. Experimenters were neither able to control the number of trials a participant performed, nor to control the precise location of microelectrodes; nevertheless, we report similar numbers to those reported in previous studies\textsuperscript{38,51–53}.

Data acquisition and preprocessing. Data were acquired at two electrophysiological scales from each participant: SUA and ECoG. SUA was recorded from microelectrodes using three different techniques. In Cohort 1, the dIPFC–dACC sEEG electrodes were Behnke-Fried macro–micro-electrodes (AdTech Medical), which consist of a standard clinical depth macroelectrode shaft with a bundle of eight shielded microwires that protrude approximately 4 mm from the tip. These eight microwires are referenced to a ninth unshielded microwire. dACC sEEG was acquired with this technique (1) and (2). Cohort 2 used a dACC–dACC CECoG using two techniques (Fig. 5a). The DBS surgeries were performed according to standard clinical procedure, using clinical microelectrode recording (FHC). Before inserting the guide tubes for the clinical recordings, we placed the microelectrodes in the cortex under direct vision to record from the dIPFC, as we have previously described in the epilepsy implied in Cohort 2 included a UMA implant from the SUA, the dIPFC, as we have previously described\textsuperscript{15–17}. Data were amplified, high-pass filtered, and digitized at 30 kilosamples per second on a neural signal processor (Blackrock Microsystems) simultaneously with the ECoG data.

LFPs were recorded from subdural CECoG electrodes or standard clinical sEEG microelectrode recordings. In Cohort 1, we recorded single-unit action potentials along the clinical CECoG electrode and referenced to either a scalp electrode or a quiescent sEEG contact in the cerebral white matter, depending on the clinical recording configuration. The medial contacts were within the dACC, and lateral contacts within the dIPFC (Fig. 1a). In the DBS patients in Cohort 2, LFP data were recorded from eight-contact (3 mm electrode diameter, 5 mm inter-contact spacing) CECoG strips (PMT). The CECoG strips were sutured to the cortical surface through the burr hole adjacent to the microelectrodes and were referenced to scalp needle electrodes adjacent to the mastoid bone. For the epilepsy cohort in Cohort 2, LFP data were recorded from the nearest CECoG electrode on the grid overlying the UMA and referenced to an epidual ECoG strip. In all cases, signals from ECoG contacts were pseudodifferentially amplified tenfold and digitized at 2 kHz on the same recording system, and therefore same time base, as the SUA and task event data.

LFP data were preprocessed by first removing clearly broken CECoG or sEEG electrodes and then removing the common mode across channels by reconstructing the data without the first principal component. The resultant time series were then epoched from 2 s before until 3 s after the time of stimulus onset. Trials containing epileptiform discharges were removed based on the range of the LFP voltage across all trials. Trials in which the range of the signal was greater than 1.5 times the interquartile range of the distribution of voltage ranges across all trials were manually reviewed and excluded. LFP spectra between 0 and 150 Hz were calculated with multi-taper methods using the Chronux toolbox for Matlab with five leading tapers and a time-bandwidth product of 3.

Action potential sorting. SUA data were re-thresholded offline at $-4 \times$ the root mean square of the 250 Hz high-pass filtered signal. Well-isolated action potential waveforms were then segregated in a semi-supervised manner using the T-distribution expectation-maximization method on a feature space comprised of the first three principal components using Offline Sorter software (Plexon)\textsuperscript{19}. The times of threshold crossing for identified single units were retained for further analysis.

Classification of single unit selectivity. To estimate neuronal selectivity, we fit a sliding GLM (using similar methods to those used in ref.\textsuperscript{15}) to the normalized firing rate of each neuron averaged over a time window of 250 ms each, and repeated this process iteratively, shifting in steps of 20 ms for the duration of the whole trial. This GLM, implemented with custom scripts in Matlab, consisted of a three-way analysis of variance (ANOVA) with factors: decision conflict (four levels: neither conflict, spatial conflict, flanker conflict, and both conflicts), response identity (three levels: button 1, 2, or 3), and feedback valence (two levels: neutral versus valenced feedback). This ANOVA model also accounted for reaction time as a nuisance variable and included interaction terms for all four factors. Using this sliding GLM, we classified individual neurons on the basis of their firing rate selectivity for these task-relevant features.

We focused our analyses on target windows of 500 ms during a post-stimulus period centered 500 ms after stimulus onset rather than on longer time windows so as to limit the number of statistical tests. To establish statistical significance while controlling for multiple tests within each target window (that is, one test for each of 25 20-ms steps), we performed a permutation test in which neurons of neuronal spike data corresponding to a trial were randomly reshuffled across trials, for each of the neurons separately, for a total of 10,000 iterations. Using the main three-way ANOVA model with test windows of 250 ms and steps of 20 ms, a combined threshold of $P \leq 0.05$ for at least four consecutive steps yielded significant effects for the first factor (in a 500 ms target window) in less than 5% of the surrogate neurons. The $\alpha$ value of 0.03 was chosen to correct for the number of consecutive bins required to reach significance, and corresponds to a corrected $\alpha$ value of 0.05. This combined significance threshold, which had the higher yet significant $P$ value combined with longer duration, was chosen over a combined threshold of lower $P$ value with shorter duration because meaningful electrophysiological effects (neural responses) only become apparent over longer time windows that contrast with noise effects (false positives). We therefore adopted this combined height-duration threshold in all of our analyses of individual neurons to control for
multiple comparisons. McNemar's test with a Yates' corrected $p$ were used to test for differences in proportions of neurons. All tests were two-sided.

SFC. SFC was calculated using multi-taper methods with the Chronux toolbox for Matlab\(^6\). SFC coherograms were generated using a 1 s window size, a 10 ms step size, a time-bandwidth product of five, and nine leading tapers. Significant changes in coherence were determined with permutation tests, for which random trials of SUA and LFP data were paired. Such a randomized trial pairing was performed 10,000 times, and SFC was recalculated between random pairs of SUA and LFP data in order to generate SFC null distributions. Coherograms for each neuron and condition (randomly subsampling numbers of trials such that they were balanced across conditions) were then tested against these null distributions to determine statistically significant frequency bands and time ranges using cluster-corrected permutation tests with significance levels of 0.05 (ref. 30). Only significant clusters following the onset of the stimulus and before the maximum reaction times were considered. We refer to this procedure as the SFC permutation test. The phase and amplitude of coherence among conditions were then examined for neurons exhibiting significant SFC clusters. To enable comparison with the firing rate models, these phase or reliability coding effects were assessed using a sliding ANOVA over the same time period as that for which firing rate effects were assessed, specifically 250–750 ms following stimulus onset\(^30\).

Single-trial coherence for the UMA data was calculated as described above, except that coherence was calculated across simultaneously recorded neurons for each trial. Ordinary least squares linear regression was used to predict reaction times in each condition from single-trial theta coherence calculated among simultaneously recorded dlPFC neurons. To confirm that the amount of decision conflict on previous trials did not affect our ability to predict reaction time from single trial coherence across a population of dlPFC neurons, we implemented a multiple regression model of reaction time using theta coherence, current trial conflict level, previous trial conflict level, and the interaction between current and previous trial conflict levels as predictors (reaction time ~ 1 + theta coherence + current trial conflict level × previous trial conflict level).

**Discriminability.** To roughly compare the time courses of conflict effects across coding schemes, we fit a linear discriminant model to each timepoint of the firing rate or coherence phase data across populations of each category of neuron\(^30\). This model was evaluated with fourfold cross-validation, and a permutation threshold based on reshuffling conflict labels among trial data was used to determine significance (values greater than 95% of the area under the permutation distribution).

**stLFPs.** To understand the average influence of a dACC neuron's action potential on population synaptic dynamics in the dACC and dlPFC, we aligned 3 s of LFP recorded in the dACC and dlPFC before and after the time of each dACC spike occurring between 250 ms and 750 ms following the stimulus. These LFPs were decorrelated by their covariance matrix and then $z$-scored. Averaging these LFPs generated an stLFP. stLFPs have been shown to be closely related to the cross-correlation of the intracellularly recorded membrane potential and the surrounding LFP\(^59\). To generate null distributions to test for any stLFP effect, while maintaining both the temporal structure in the population firing rate and each neuron's contribution to the mean population firing rate overall, spikes were also randomly shifted among adjacent 1 ms time bins and across adjacent trials using the raster margins model\(^30\). Null distributions for stLFP were generated by calculating stLFPs from these shifted spike times and the original LFP as described in the raster margins model. stLFP amplitude was operationally defined as the minimum stLFP during the 200 ms following the mean dACC spike time. Again an LMM was used to assess significance among stLFPs that were associated with particular classes of neurons. The model specification in this case was: stLFP ~ 1 + conflict-selective neurons × session + theta-coherent neurons × session + beta-coherent neurons × session + non-coding neurons × session + (1 + conflict-selective neurons × session + theta-coherent neurons × session + beta-coherent neurons × session + non-coding neurons × session | participant).

**Reporting Summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

Data are available from the authors upon reasonable request and with permission of the Columbia University Medical Center Institutional Review Board.

**Code availability**

All analysis code is available online at http://www.github.com/elliotsmith/MSIT-analysis

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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection
- Neural data collection utilized Blackrock Microsystems Central Software (version ). Behavioral data collection utilized Psychophysics Toolbox (version 3)

Data analysis
- Data analyses were carried out using the chronux toolbox (version 2), Plexon Offline Sorter (version 3.4), and custom Matlab scripts uploaded to github.com/elliothsmith/MSIT-analysis

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Sample size
Sample sizes were based on numbers of trials and neurons, both of which were randomized. The number of trials was determined by the patients’ willingness and motivation to perform the task. The conflict type for each trial was randomly assigned from a uniform distribution. The number of neurons was based on where microelectrodes ended up recording, approximately 4 mm away from the nearest macroelectrode, which was placed based on clinical parameters. Experimenters were neither able to control the number of trials a patient performed, nor the precise location of microelectrodes.

Data exclusions
Trials containing epileptiform discharges were removed based on the range of the LFP voltage across all trials. Those trials in which the range of the signal was greater than 1.5 times the interquartile range of the distribution of voltage ranges across all trials were manually reviewed and excluded.

Replication
Cross validation and statistical controls were used when possible. Replication for this study mainly involved determining that neuron-level effects were present across subjects.

Randomization
Randomization was carried out by the task, which was determined by a randomly seeded random number generator in Matlab.

Blinding
Subject-level blinding was not possible at the time of data acquisition, though researchers were blinded to trial-level, and neuron-level information at the time of acquisition.

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Human research participants

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Population characteristics
Subjects were neurosurgical patients undergoing monitoring for medically refractory epilepsy or surgical implantation of deep brain stimulation electrodes. Subjects ranged in age from N to N years. N out of N subjects were female.

Recruitment
All patients who were undergoing the aforementioned surgeries were asked if they would like to participate in the research. No patients declined to participate, suggesting a lack of self-selection bias. The only potential selection bias results form these patients neurological disorders.

Ethics oversight
the Columbia University Medical Center Institutional Review Board approved these experiments.

Note that full information on the approval of the study protocol must also be provided in the manuscript.