Subthalamic nucleus gamma activity increases not only during movement but also during movement inhibition

Petra Fischer¹,²*, Alek Pogosyan¹,², Damian M Herz¹,², Binith Cheeran², Alexander L Green², James Fitzgerald², Tipu Z Aziz², Jonathan Hyam³, Simon Little³, Thomas Foltynie³, Patricia Limousin³, Ludvic Zrinzo³, Peter Brown¹,², Huiling Tan¹,²

¹Medical Research Council Brain Network Dynamics Unit at the University of Oxford, Oxford, United Kingdom; ²Nuffield Department of Clinical Neurosciences, John Radcliffe Hospital, University of Oxford, Oxford, United Kingdom; ³Unit of Functional Neurosurgery, Sobell Department of Motor Neuroscience and Movement Disorders, University College London Institute of Neurology, London, United Kingdom

Abstract Gamma activity in the subthalamic nucleus (STN) is widely viewed as a pro-kinetic rhythm. Here we test the hypothesis that rather than being specifically linked to movement execution, gamma activity reflects dynamic processing in this nucleus. We investigated the role of gamma during fast stopping and recorded scalp electroencephalogram and local field potentials from deep brain stimulation electrodes in 9 Parkinson’s disease patients. Patients interrupted finger tapping (paced by a metronome) in response to a stop-signal sound, which was timed such that successful stopping would occur only in ~50% of all trials. STN gamma (60–90 Hz) increased most strongly when the tap was successfully stopped, whereas phase-based connectivity between the contralateral STN and motor cortex decreased. Beta or theta power seemed less directly related to stopping. In summary, STN gamma activity may support flexible motor control as it did not only increase during movement execution but also during rapid action-stopping.

DOI: 10.7554/eLife.23947.001

Introduction

Previous studies have described a neuronal stopping network involving prefrontal and supplementary motor cortical regions, as well as the subthalamic nucleus (STN) (Aron et al., 2014; Jahanshahi et al., 2015; Rae et al., 2015). The STN is well-positioned to cancel actions as it receives cortical input via the hyperdirect pathway and can inhibit the thalamus and brainstem via the basal ganglia output nuclei as well as the striatum via the globus pallidus externus (GPe) (Mink, 1996; Wei and Wang, 2016). In spite of recent advances in understanding functional and effective connectivity within the stopping network using fMRI (Rae et al., 2015, 2016; Xu et al., 2016), the fast temporal dynamics of population activity accompanying the stopping process are not entirely clear.

When rats attempted to cancel an action, increased STN firing activity was found irrespective of whether cancellation was successful or not (Schmidt et al., 2013), but more recently, micro-electrode recordings in the human STN revealed two distinct subpopulations that selectively increased firing rate either during successful response inhibition or during motor execution (Bastin et al., 2014; Benis et al., 2016). Also in the GPe a subpopulation termed arky pallidal cells, which seem to receive input not only from the striatum but also from the STN (Nevado-Holgado et al., 2014), has specifically been linked to action cancellation (Mallet et al., 2016). It is unclear, though, how...
different populations within the basal ganglia are activated in a selective and flexible way. Oscilla-
tions, particularly in the gamma band (>30 Hz), have been proposed to be a key mechanism for
coordinating spatially separate but functionally related assemblies (Bosman et al., 2012; Fries, 2015;
Nikolić et al., 2013; Schoffelen et al., 2005). We hypothesized that gamma activity may thus also
facilitate coordinated activation of task-relevant subpopulations for efficient movement cancelation.

A local field potential study in Parkinson’s disease patients, however, has shown increased 55–75 Hz
gamma activity when patients failed to stop (Alegre et al., 2013), which is in line with the prevailing
view that gamma activity is pro-kinetic (Cassidy et al., 2002; Fogelson et al., 2005; Litvak et al.,
2012) or related to response vigour (Jenkinson et al., 2013). Beta activity, instead, is widely viewed
as a marker of broad motor suppression within the STN (Wessel et al., 2016a) as well as cortex
(Swann et al., 2012). High STN beta activity for example was linked to elongated response times
during incongruent trials in a Stroop task (Brittain et al., 2012) and to stronger suppression of cor-
tico-spinal excitability during speech inhibition (Wessel et al., 2016a). However, as movements are
known to coincide with decreasing beta and increasing gamma activity (Joudi et al., 2012; Lalo et al.,
2008), comparisons between executed and withheld movements might reflect the lack
of movement rather than the stopping process per se.

Ideally, stopping would be recorded as a continuous variable that measures how fast an ongoing
movement is terminated instead of whether an action has been started at all. Motor inhibition has
traditionally been investigated with stop signal or Go/NoGo tasks, in which movements are trig-
gered by cues (Huster et al., 2013; Swick et al., 2011). In the stop signal paradigm, subjects press
a button in response to a go cue and in some trials a stop signal instructs them to withhold the
movement. Go/NoGo tasks instead rely on a large fraction of go trials to catch participants out on
rare trials, in which a NoGo cue signals them to withhold the pre-potent motor response. Both tasks
require participants to decide whether to stay or move but not to interrupt an ongoing action. Suc-
cessful stopping is achieved in these tasks by successfully delaying or canceling action initiation
rather than terminating an action that is already ongoing. Our aim was to extend existing studies by
investigating rhythmic movements that can be interrupted halfway and are not directly preceded by go-cues but are self-initiated. Patients were asked to tap rhythmically to a metronome. Under these circumstances, subjects anticipate the metronome instead of reacting only after each sound, and so movements can be considered self-initiated. They were instructed to stop upon hearing a different cue that was timed such that they were able to stop only in approximately half of their attempts (Figure 1). The neural response to the stop signal was not intermixed with a foregoing response to a go cue as the last metronome sound was delivered about 700 ms prior to the stop signal.

The delay of the stop signal was set by the experimenter after a training period at the start and then kept constant for the rest of the experiment. It was delivered relative to the tap instead of the metronome sound to keep movement variability to a minimum and to prevent the strategy of delaying the tap relative to the metronome sound. This, in combination with the instruction to synchronize accurately to the metronome, provided trains of self-initiated actions that were well-matched across trials. The task was also well-suited to investigate endogenous fluctuations in readiness to stop. We analyzed STN local field potentials (LFP) and scalp electroencephalography (EEG) activity recorded in this task from nine Parkinson’s disease patients, who underwent deep brain stimulation surgery. To differentiate volitional motor inhibition from salience detection, six of them were recorded in an additional control condition with identical auditory cues but different instructions. Their task in this condition was to finish the tapping sequence with two more taps upon hearing the stop signal instead of attempting to stop (Figure 1).

**Results**

**Behavioural results**

The mean stop signal delay time with a 55% ± (SD) 10% successful stopping rate was 707 ± 49 ms (range = 620–760 ms). The mean interval between the preceding tap and the unsuccessfully inhibited tap in trials where stopping failed was 864 ± 36 ms, and was significantly shorter than the 900 ms interval dictated by the metronome (Wilcoxon signed-rank test, p=0.004). In these trials, patients would have still had on average 156 ms to stop.

Movement trajectories preceding successfully or unsuccessfully inhibited stops were overlapping (Figure 1, trajectories measured by a pressure sensor and goniometer). Thus any electrophysiological differences in this window are unlikely related to movement differences per se.

Stopping performance was quantified as movement extent, which was the extent of downward movement after the stop signal relative to the amplitude of the preceding upward movement. 0% movement extent thus refers to a full stop. 50% describes a movement that was interrupted halfway and 100% would correspond to a full tap, i.e. failed stopping. Correlations between movement extent and various properties of the last regular tap were computed for each patient and then subjected to t-tests to assess if the Fisher’s z-transformed correlation coefficients significantly differed from zero on the group-level. In 7 of 9 subjects, movement extent correlated with the tap-to-sound offset, which indicates that stopping performance was worse when the foregoing tap was relatively late in a trial corresponding to previous results (Fischer et al., 2016). However, none of the tested variables were associated with successful stopping after FDR-correction of the resulting p-values (see Table 1).

Previous research suggests that a surprising sound alone already elicits motor slowing of verbal reports (Wessel and Aron, 2013). We thus checked if the tap performed after the salient ‘stop signal’ (which served as ‘continue signal’ in the control condition) was delayed or slowed down in the control condition when stopping was not even required. The median intertap interval directly preceding the stop signal (median ITI = 893 ms) did not differ significantly from the one directly after the stop signal (median ITI = 889 ms, Wilcoxon signed-rank test p=1.0).

**LFP and EEG power differences following the stop signal**

We tested for rapid LFP and EEG power changes between the stop cue and the average timing of the tap when inhibition failed, which was on average 156 ms after the cue and puts a limit on the window within which successful movement inhibition had to occur.

The STN contralateral to the tapping hand responded to the stop signal with a 60–90 Hz gamma power increase when compared to activity from the tap before (Figure 2.1 shows the reference data
Figure 1. Behavioural task and representative data. (A) Schematic of the task in the STOP condition (top row) and in the control condition (2nd row). In the STOP condition participants had to tap (=ellipses) to a metronome (=rectangles) and stop after 5–9 taps. The red ellipse denotes a tap that was unsuccessfully stopped. (B) Pressure sensor, FDI muscle activity and goniometer data from one representative patient. Black lines are trials where the tapping movement after the stop signal was successfully stopped, red lines are trials where stopping failed. The markers around 0 ms represent the temporal offset between the last regular sound and the tap (o = successful stop trials, x = failed stop trials). The markers at 680 ms show the time of the stop signal, which was always triggered relative to the last regular tap that was registered by the pressure sensor at 0 ms. Note that the black and red trajectories overlap, which shows that stopping performance did not depend on the preceding movement trajectory.

DOI: 10.7554/eLife.23947.003
Table 1. Correlations between movement parameters of the last regular tap and the movement extent after the stop signal (mean ± SD). In 7 of 9 subjects, movement extent correlated with the soundOffset (=tap-to-sound offset; negative values represent taps that occurred before the sound). But none of the p-values resulting from one-sample t-tests of the Fisher’s z-transformed intra-individual correlation coefficients of the nine subjects survived FDR-correction. downTime = duration of finger contact with the pressure sensor, maxPres = peak pressure during the tap, tapNr = number of taps preceding delivery of the stop signal, peakVelDown=peak velocity of the downward movement of the previous tap, upMvmt = amount of up-movement, peakVelUp=peak velocity of the upward movement.

| Variable       | Rho ± SD   | p-value | FDR-corrected p-value |
|----------------|------------|---------|-----------------------|
| soundOffset    | 0.29 ± 0.18| 0.020   | 0.137                 |
| downTime       | −0.10 ± 0.19| 0.174   | 0.407                 |
| maxPres        | −0.04 ± 0.23| 0.460   | 0.644                 |
| tapNr          | −0.13 ± 0.20| 0.061   | 0.215                 |
| peakVelDown    | −0.05 ± 0.23| 0.377   | 0.644                 |
| upMvmt         | 0.00 ± 0.23| 0.952   | 0.952                 |
| peakVelUp      | 0.04 ± 0.29| 0.810   | 0.945                 |

DOI: 10.7554/eLife.23947.004

Figure 2. Contralateral STN power changes around the stop signal. T-scores calculated over all patients (n = 9, normalized by the average power during regular tapping) for (1) the last regular tap aligned to the timepoint when the stop signal would have occurred if it would have been delivered one tap earlier (vertical dashed line). The black line shows the tapping movement measured by the goniometer. The downward movement was accompanied by a beta decrease and gamma increase as expected. The following three columns show changes in response to the stop signal (vertical dashed line) (2) irrespective of whether stopping was successful or not, (3) during successful stops only, and (4) during failed stops only. Note that when a stop signal was present and especially when stopping was successful (column 3), gamma increased strongly. Differences between 2–1 and 3–4 are contrasted in Figure 3. The tapping trajectory of failed stops does not reach the bottom line even though the finger touched the table because trajectories were normalized to the minimum of all four trajectories, which occurred with the last regular tap, where the spring was extended more vigorously than during attempted inhibition.

DOI: 10.7554/eLife.23947.005
from the tap before aligned to where the stop signal would have occurred if it would have been presented one tap earlier; Figure 2.2 shows the response to the stop signal; Figure 3A shows the contrast between the two). Importantly, this gamma increase was significantly and consistently higher during successful movement inhibition (Figure 2.3 + 2.4, and 3B). The effect size of this difference was very large (60–90 Hz power difference between successful-failed stops: Cohen’s d mean<sub>winO</sub> = 1.2, max<sub>winOdiff</sub> = 2.6). Note that during regular tapping we observed the typical pattern of movement-related gamma power increase and beta power decrease (Figure 2.1 and Figure 3—figure supplement 1). Gamma power thus increased during both movement execution and movement inhibition. The movement-related peak was broader and weaker than the stop-related increase that peaked around 70 Hz (Figure 3—figure supplement 2).

Cortical EEGs recorded a low-frequency increase in response to the stop signal in all channels (Figure 3A), which was – in contrast to STN gamma activity – not significantly higher during successful stopping (Figure 3B). Only 8–30 Hz power over contralateral C3/C4, Cz and Fz was significantly higher when stopping was successful. However, there was no overall power increase following the stop signal in the 8–30 Hz band in these channels when compared to the tap before (Figure 3A), not even when only successful stop trials were considered (Figure 3—figure supplement 3). In previous studies, such increase was observed when an action had to be withheld before being initiated (Kühn et al., 2004; Swann et al., 2009).

To exclude that the STN gamma increase merely reflects processing of the salient stop cue, six patients additionally performed a control condition before and after the main stopping task (Figure 1). The stimulus sequence of the control condition was identical to the main condition and the instruction differed only in that patients had to finish the tapping sequence with two more taps upon hearing the stop signal instead of inhibiting the tap immediately. Importantly, no gamma increase was observed in this control condition, even though the difference between successful and unsuccessful stops was still significant despite the reduced sample size of 6 patients (Figure 4C).

It has been suggested that specifically the right STN may mediate stopping (Aron and Poldrack, 2006). To evaluate the role of the right STN alone, individual gamma differences between successful and unsuccessful stops of all right STNs are displayed in Figure 4B, showing no significant increase. Three right-handed patients performed the task with the left hand and thus in those the right STN was the contralateral one. However, in the remaining six, the right STN was the ipsilateral STN, and thus the lack of significant right STN gamma increase indicates that the gamma increase was specific to the contralateral STN.

To further corroborate the functional significance of our finding we also tested whether the average gamma increase peaked earlier during successful stops than during failed stops. Indeed, the average gamma peak of successful stops at 106 ± (SD) 59 ms occurred earlier than the average unsuccessful tap (at 156 ± 50 ms), whereas the average gamma peak of failed stops occurred later (at 179 ± 84 ms). These gamma peak timings significantly differed from each other (t(8)=-2.9, p=0.019, CI<sub>diff=[−131, −16 ms], Cohen’s d = 1.0).

We also examined within-subject correlations between movement extent (i.e. inhibition failure) and gamma within the stopping window (0–156 ms) after the stop signal (see Figure 4—figure supplement 1). This was significant in 8 of 9 patients (uncorrected tests; P3’s confidence intervals were borderline significant, Spearman’s rho p=0.049) when gamma was taken from the contralateral STN, meaning that in all but one patient we found that when gamma was higher, movement extent was less and stopping was more successful. In contralateral C3/C4 and in ipsilateral STN such a relationship was present only in three patients, and in ipsilateral motor cortex only in one patient, further indicating specificity to the correlation with contralateral STN activity. Note though that correlations might be harder to detect with EEG data due to the reduced signal-to-noise ratio in comparison with LFPs.

To see if the gamma increase was highest specifically during full stops, we classified the movement after the stop signal into full stops (<10% downward movement), intermediate stops (>10% but pressure sensor was not touched) and failed stops (all trials where the pressure sensor was touched). Only four patients made five or more full stops (mean number of full stops = 9.5), so formal statistics were not applied. Still, in full stop trials, gamma increased most strongly. It increased moderately for intermediate stops and remained flat for failed stops (Figure 4—figure supplement 2). As expected, activity recorded from the first dorsal interosseous muscle of the tapping hand...
Figure 3. Contrasts between power changes following the stop signal. (A) T-scores calculated over all patients of the contrast between power aligned to the stop signal (vertical dashed line) averaged across all trials irrespective of stopping performance (Figure 2.2) and the regular tap made before (Figure 2.1, aligned to where the stop signal would have occurred if it would have been presented one tap earlier). Red clusters denote that power significantly increased in response to the stop signal. (B) T-scores of power differences between successful and failed stops. Red clusters denote that power was significantly higher if participants successfully inhibited the upcoming tap (Figure 2.3–2.4).

DOI: 10.7554/eLife.23947.006

The following source data and figure supplements are available for figure 3:

Fischer et al. eLife 2017;6:e23947. DOI: 10.7554/eLife.23947

Figure 3 continued on next page
(presented to the right in Figure 4—figure supplement 2) suggests an inverse relationship to the gamma increase. Finally, we examined if the cortical 3–5 Hz power increase, which was clearly present in the stop condition (Figure 3A), was also present in the control condition when movement inhibition was not even attempted. The grey power trajectory representing the control condition shows a very similar peak in Cz (Figure 4D, n = 6). Significance testing within the crucial reaction time window (ranging from the stop signal to the average time of the failed tap, 156 ms later) resulted in no significant differences between the control condition and either the power increase during failed or successful stopping. The direct comparison between failed or successful stops was not significant either. Also a peak-extraction analysis failed to detect a difference between low-frequency peaks (Cz successful stops vs. control: t(5)=0.2, p=0.848, CI_{diff}=[-0.205, 0.205]; failed stops vs. control: t(5)=0.5, p=0.641, CI_{diff}=[-0.149, 0.220]). The 3–5 Hz increase only seemed to be reduced in the control condition in Fz and both M1 (Figure 4—figure supplement 3), however this was also not significant.

Changes in connectivity between cortex and STN following the stop signal

In a next step, we computed intersite phase clustering (ISPC) values between filtered oscillations in the EEG recordings and the LFP signal from the STN contralateral to the tapping hand. To get an estimate of the temporal development, we subdivided a –350:160 ms time window around the stop signal into equal bins in which ISPC was computed for each trial and then averaged over trials (see Materials and methods). ISPC describes whether phase differences between two sites are randomly distributed (small ISPC → low connectivity) or clustered (high ISPC → high connectivity) and was obtained by taking the length of the mean vector of all phase differences from all time points within one bin.

ISPC of 60–75 Hz gamma between the contralateral motor cortex and the contralateral STN decreased strongly and significantly in response to the stop signal relative to the average of the –350:0 ms window preceding the stop signal (Figure 5). We also observed an increase of 6–12 Hz ISPC to all cortical channels.

Power differences preceding the stop signal

Finally we assessed if gamma power was already tonically elevated prior to the stop signal, before participants knew they had to stop. We tested for significant differences within a 350 ms window before the stop signal. If the upcoming tap was inhibited more successfully, STN gamma power was already higher prior to the stop signal (Figure 6).

20–30 Hz beta power over C3/C4 ipsilateral to the tapping hand also was significantly higher preceding successful stops. If the data were re-aligned to the last regular tap instead of the stop signal, a second significant cluster at 20–30 Hz over C3/C4 contralateral to the tapping hand was found, in line with previous reports (Fischer et al., 2016) (see Figure 6—figure supplement 1).

Discussion

We found that when finger tapping had to be stopped abruptly, the stop signal elicited a fast increase in 60–90 Hz gamma activity in the contralateral STN and a pronounced theta increase in cortex. However, only the former was significantly higher when stopping was successful. The gamma increase occurred within 156 ms, which was the brief time window between the stop signal and the
average failed tap. In a control condition, in which participants were presented with the same stop signal while tapping, but stopping was not attempted, only cortical theta but not STN gamma power increased. This shows that STN gamma activity does not only reflect pro-kinetic activity as previously suggested (Litvak et al., 2012) nor does it merely reflect processing of the salient stop signal.

The alternative hypothesis that stopping of the tapping movement itself involved an active movement seems unlikely on two grounds. First, the gamma increase was less if the tap was terminated mid-flight rather than before the downward finger movement was started. Second, gamma connectivity between the STN and C3/C4 also sharply decreased directly after the stop cue, which differs
from the movement-related increase usually observed (Litvak et al., 2012) and may indicate disengagement from the obsolete motor plan.

Two previous studies have reported a different relationship between gamma and stopping success to the one that we have found (Alegre et al., 2013; Ray et al., 2012). Ray and colleagues (2012) reported a gamma increase in response to the stop signal as we do (see Ray et al., 2012: Figure 4b) but did not detect significantly higher gamma during successful stops. This discrepancy may result from extensive temporal smoothing (their sliding window was 333 ms long), and from the pre-selection of a window of interest between 200–400 ms after the stop signal, which also would have failed to detect the gamma difference in our data occurring right after the cue. If we apply the same temporal smoothing to our data, the power trajectories of successful and failed stops would look very similar (data not shown) as extensive smoothing flattens the brief gamma increase, such that part of it appears before the stop signal. The late gamma increase during failed stopping, which would be too late to affect the stopping outcome, would remain as a prominent difference about 300 ms after the stop signal. In Alegre et al.’s study, which differed methodologically in using a visual stop signal, one conclusion was that successful inhibition was associated with a bilateral gamma power decrease. The precise time-frequency decomposition parameters used in that study are unclear and so we have not re-analysed our data in the same way. However, as their window of interest was relatively long (0–0.4 s), they might have predominantly captured the pro-kinetic gamma component that is relatively reduced when a motor response is withheld. Even so, similar as in our study, gamma appeared to increase briefly when aligned to the stop signal in patients on medication during both successful and failed stopping attempts (Alegre: Figure 5) and a drop in STN-M1 coherence during successful inhibition also was found (Alegre: Figure 7). In addition, when comparing results between studies it is important to acknowledge that differences in disease phenotype, medication history, electrode models and in the precision of the targeting achieved are factors that may

Figure 5. Connectivity changes following the stop signal. Intersite phase clustering (ISPC) values are normalized by a –350.0 ms baseline preceding the stop signal. The dashed line denotes the time of the stop signal. Gamma ISPC between contralateral STN and contralateral C3/C4 decreased significantly between 60–80 Hz (encircled in red), whereas ISPC in low frequencies between STN and cortical electrodes increased.

DOI: 10.7554/eLife.23947.016

The following source data is available for figure 5:

Source data 1. MATLAB data file containing source data related to Figure 5.

DOI: 10.7554/eLife.23947.017
also contribute to variability in study findings. The normalisation of LFP measures will have only partially mitigated this variability.

Periods of high gamma activity in the STN have been reported to coincide with an overall increase in firing rate and phase-locking of spikes to the gamma cycle peak (Pogosyan et al., 2006; Trottenberg et al., 2006). From the LFP we cannot infer changes in firing rate, but it suggests that the number of neurons or inputs to these neurons synchronizing at 60–80 Hz was coupled with stopping outcome, and that increased synchronisation occurred early enough to influence such outcome. After observing that the strength of gamma synchronization in the STN or its coherence with C3/C4 did not depend on the exact movement performed, Litvak and colleagues suggested that STN gamma activity modulates rather than explicitly encodes motor commands (Litvak et al., 2012). Our results take this hypothesis further by extending the concept of modulation to include a possible role for movement cancelation. This notion is also compatible with observations that have linked STN gamma activity to effort (Jenkinson et al., 2013; Oswal et al., 2013; Tan et al., 2013) and arousal (Brücke et al., 2013; Jenkinson et al., 2013; Kempf et al., 2009). The fact that stopping was more likely successful after STN gamma was relatively high already 200 ms before the stop cue (i.e. before patients knew they had to stop) may reflect such arousal-related function and the need for proactive inhibition.

The present study is correlative in nature, so we cannot infer that gamma oscillations are causally involved in stopping. However, we would like to speculate that a strong surge in STN gamma activity may shift the excitable period of the otherwise observed pro-kinetic gamma increase such that pre-synaptic spikes arrive at a period of relative inhibition and motor output thus may be interrupted. Inter-individual variability of the peak frequency and the strength of the STN gamma increase may have been related to differences in disease progression, individual stopping speed or electrode placement and type. We did not find a significant gamma increase in cortical electrodes, which may be due to the reduced signal-to-noise ratio of the EEG. However, a broad gamma increase was
observed during stopping in electrocorticography recordings from the pre-supplementary motor area and right inferior frontal gyrus (Swann et al., 2012), raising the possibility of cortical involvement in generating the gamma increase via the hyperdirect pathway.

In comparison to tetrode recordings in rats (Schmidt et al., 2013), our study is limited in that the recording contacts may not have been directly in the STN. The SNr is located in close proximity, ventrally adjacent to the STN, and thus we cannot exclude that we picked up activity from neighbouring structures. However, gamma has been reported to be specifically localized in the dorsal part of the STN (Trottenberg et al., 2006), so that contacts selected according to the strongest gamma modulation are likely located closer to the dorsal border of this nucleus. However, this remains speculative.

Another limitation of this study is that we recorded from patients that may exhibit pathological STN hyperactivity (Hamani et al., 2004) expressed in abnormal firing rates and patterns (Magnin et al., 2000; Remple et al., 2011). Even though these pathological changes are attenuated by dopaminergic medication (Brown et al., 2001; Heimer et al., 2006; Levy et al., 2002), which was taken as usual, and patients were able to perform the task well, neuronal dynamics may still have differed from those of healthy subjects with intact basal ganglia circuits.

It may also be argued that stopping may have involved muscle contractions, which were not picked up by the FDI EMG. But the short latency of the gamma increase and the absence of a similar increase in motor cortex in combination with the decrease in connectivity between STN and C3/C4, which would be expected to increase during movements (Litvak et al., 2012), renders this possibility unlikely. Additionally, we observed that gamma increased most strongly in trials where participants were able to stop fully instead of interrupting the downward movement halfway, showing that gamma increased not only during braking in the middle of a movement but that it increased even more in the absence of any movement.

As reported previously, we confirmed a link between higher post-movement C3/C4 beta activity and subsequently improved stopping performance, which we suggest was related to fluctuations in cognitive load (Fischer et al., 2016). Beta has also been implicated in time estimation (Kononowicz and van Rijn, 2015), thus it may also reflect an intention to delay the next tap’s timing, which would allow for more time to stop. Stopping success indeed was correlated with the tap-to-sound offset of the last regular tap in seven patients, such that relatively early taps (early with respect to the sound, which should be compensated for by delaying the next tap) were followed by higher stopping success. Note that this beta difference was not present in the STN. We also observed significantly higher beta power over contralateral motor and frontal cortex when stopping was successful in comparison to when it failed. As beta oscillations are less likely to occur during movement execution (Feingold et al., 2015; Kilavik et al., 2013), this difference was expected. In the past, a number of studies have suggested that beta plays an active role in motor inhibition (Bastin et al., 2014; Brittain et al., 2012; Wessel et al., 2016a). Importantly, in the present study no beta increase was observed after the stop signal in comparison to the previous regular tap − not even when only successful stop trials were considered. Thus, it seems unlikely that bursts of beta oscillations per se implemented active braking in our task. Increased beta in other studies may have reinforced the resting position as current motor state that had to be maintained (Gilbertson et al., 2005). Such resting posture was not present in our task given that the stop signal was delivered during ongoing tapping.

How can we reconcile the above with the results reported by Benis et al. (2014), who observed a weaker STN beta decrease during ‘proactively inhibited’ go-trials (‘proactively inhibited’ as participants were aware that a stop signal may come after the cue, although it did not appear in these trials) in comparison to go-trials with a cue, which was never followed by a stop signal and thus resulted in faster reaction times? The stronger beta decrease may have been related to a more vigorous response in fast go-trials (Tan et al., 2013, 2015) or reduced response uncertainty (Tzagarakis et al., 2010) and thus does not necessarily need to reflect an inhibitory process. A stronger difference in beta decrease between the two trial types was also linked to shorter stop signal reaction times across patients. But this correlation may be mediated by symptom severity, as more severe symptoms could result in less beta reactivity (Little et al., 2012), reduced modulation of response vigour and longer stop signal reaction times.

Finally, our results may also be reconciled with those reported by Wessel et al. (2016a) if elevated beta activity reflects better connectivity across task-relevant areas (Gross et al., 2004) or
reduced cognitive load (particularly for <20 Hz beta) (Fischer et al., 2016; Rouhinen et al., 2013), it could support motor suppression without actually implementing movement inhibition.

Recently, an influential hypothesis suggesting that motor suppression is implemented by fronto-central low-frequency activity has received further support (Wessel et al., 2016b). Even though the authors also observed an STN gamma increase concurrent with response slowing, this increase was associated with the cognitive demands of the verbal working memory task rather than motor inhibition. Similar to classical stop signal tasks, our auditory stop cue also elicited a slow-wave power increase. However, this increase occurred also when stopping was not even attempted. If the slow-wave power increase over Cz would have induced slowing or braking, then the intertap interval in the control condition between the tap before and the tap immediately after the stop signal should have been increased, and this was not the case.

Our data suggest an alternative account, namely that the stop signal-evoked slow-wave response does not directly correspond to movement inhibition but instead registers salient sensory stimuli and alerts stopping-relevant areas, which in turn may trigger the STN gamma increase. The increase in cortico-subthalamic low-frequency connectivity might underpin this sequence, enabling the STN to trigger the stopping process. The event-related low-frequency response would thus be necessary for, but not equivalent to motor suppression per se. In Fz and M1 the average low-frequency response seemed to be diminished in the control condition. We would expect that registration of a salient stop signal and efficiency of the transmission process (in terms of speed or extent of neuronal recruitment) depends on endogenous fluctuations in arousal, attention and cognitive load, which would reconcile the hypothesis of low-frequency power-mediated salient stimuli processing with previous results regarding motor inhibition (Wessel and Aron, 2014).

Taken together, our results showed that gamma oscillations in the contralateral STN were linked to successful stopping. This indicates that gamma oscillations in the STN are not simply pro-kinetic, but that they can also increase during movement termination. Though we can only infer an association and not causation from observational recordings, our data suggest that the observed gamma rhythm may underpin a fast stopping mechanism involving the STN. Gamma oscillations therefore seem to support fast changes in processing demands not only in cortical but also in cortico-basal ganglia networks in line with theories of gamma synchrony establishing effective, precise and selective neuronal communication (Fries, 2015).

Materials and methods

Participants
Ten Parkinson’s disease patients (mean disease duration = 8 ± 4 years, mean age = 59 ± 8 years; one left-handed/ambidextrous; two female) were recorded after obtaining informed written consent to take part in this study, which was approved by the local ethics committee (Oxfordshire REC A, 08/H0604/58). One patient had to be excluded from the analysis as they intermittently fell asleep during the testing. All patients underwent bilateral implantation of deep brain stimulation electrodes into the STN two to six days before the recording with the aim to alleviate symptoms through chronic high-frequency deep brain stimulation. Surgeries and recordings were performed either at the University College Hospital in London or the John Radcliffe Hospital in Oxford, UK. For each patient one of the following three macroelectrode models were used: Medtronic 3389 (quadripolar, for P1-4 and 8), Boston Scientific, Vercise, DB-2201 (octopolar, for P6) and Boston Scientific, Vercise directional, DB-2202 (octopolar, directional, for P5, 7 and 9). Clinical details of the patients are given in Table 2. Patients were tested on medication to ensure task performance and motor function were as normal as possible, although acknowledging that functional impairments, albeit lessened, still persist in this state.

Task
Participants were asked to tap to an isochronous metronome (900 ms inter-trial interval, ITI, 700 Hz pitch, 40 ms duration) and to interrupt tapping in response to a high pitched auditory stop cue (2000 Hz pitch, 40 ms duration) after a random number of 5–9 taps. The metronome served as regular cue. However, the taps were not triggered in reaction to the metronome sounds but had to be initiated already before the sound to achieve synchronization that depends on how well the timing
of the sound is anticipated. If the movement would be reactive, the tap would always lag behind the sound, which was not the case as evidenced by a negative tap-to-sound offset. Thus, this task is special as the metronome cues are not equivalent to go cues.

The sound was generated in Spike2 with a 1401 data acquisition unit (Cambridge Electronic Design, Cambridge, UK), played by Creative Inspire T10 speakers and recorded by the EEG amplifier (TMSI Porti amplifier, TMS International, Netherlands). The timing of the stop signal was adjusted in a training period at the beginning such that patients would be able to stop only in 50–60% of all trials. Importantly, the stop signal was triggered relative to the tap registered by the pressure sensor and not to the sound to prevent patients from delaying their taps relative to the metronome, which would improve stopping performance if the latter were the case. A ~50% success rate was desirable to capture fluctuations in alertness or stopping readiness and to distinguish related brain processes.

The actual average stopping probability was 55% ± (SD) 10%. Six patients were additionally recorded in a control condition to assess if stopping-related activity was linked to active motor inhibition or whether it merely reflected registration of the more salient stop tone. In this control condition, patients were asked to end the tapping sequence with two more taps after hearing the high pitched sound instead of stopping immediately. The control condition thus posed much less of a challenge than the main stopping task.

A nearly identical task has previously been studied in young healthy subjects (Fischer et al., 2016). It differed from the present patient study only in the metronome interval duration, which was shorter (700 ms instead of 900 ms in the patients) and the number of taps (6–10 taps until the stop signal may appear instead of 5–9 taps in the patients). Intervals were chosen to be longer because stopping proved to be more feasible for patients with longer intervals, and the number of taps was reduced to increase the number of trials obtained in the time-limited recording sessions. We planned to record 100 trials in the stopping condition and 20 trials before and after the main block in the control condition. Due to fatigue and time constraints in some cases less trials were recorded. As three patients (P4, P7, P9) had severe motor symptoms on the right side, they performed the task with their left index finger. The remaining six patients used the right index finger. As we would expect the contralateral hemisphere to be more involved in the tapping, we analysed the data not

Table 2. Clinical details. Age and disease duration are given in years. UPDRS-III: Unified Parkinson’s disease rating scale part III. Levodopa equivalent dose was calculated according to Tomlinson et al. (2010).

| ID | Age/Sex/dom. Hand | UPDRS-III OFF/ON levodopa | Disease duration | Main symptom | Levodopa equivalent dose (mg / day) | DBS lead | Surgical centre |
|----|-------------------|---------------------------|-----------------|--------------|------------------------------------|----------|----------------|
| 1  | 65/f/r            | 33/11                     | 5               | Tremor/Dyskinesia | 807 mg | Medtronic 3389™ | Oxford |
| 2  | 55/m/r            | 49/25                     | 10              | Leg dragging + tremor (left side) | 2022 mg | Medtronic 3389™ | Oxford |
| 3  | 66/f/r            | 25/14                     | 17              | Freezing of gait, balance | 1089 mg | Medtronic 3389™ | London |
| 4  | 50/m/r            | 37/17                     | 5               | Tremor, Dyskinesia, especially in right foot | 958 mg | Medtronic 3389™ | London |
| 5  | 48/m/leftambi     | 46/18                     | 6               | Frequent OFFs | 800 mg | Boston Scientific DB-2202™ | Oxford |
| 6  | 54/m/r            | 61/32                     | 8               | Motor fluctuations | 455 mg | Boston Scientific DB-2201™ | Oxford |
| 7  | 60/m/r            | 37/6                      | 6               | Rigidity left side, bradykinesia, dyskinesia | 2084 mg | Boston Scientific DB-2202™ | Oxford |
| 8  | 67/m/r            | 31/13                     | 3.5             | Bradykinesia, Rigidity | 2173 mg | Medtronic 3389™ | London |
| 9  | 68/m/r            | 33/15                     | 10              | Motor fluctuations | 1765 mg | Boston Scientific DB-2202™ | Oxford |

DOI: 10.7554/eLife.23947.021
separated between left and right motor cortex and STN, but between contra- and ipsilateral C3/C4 and STN.

**Behavioural analysis**

Behavioural outliers (such as spurious goniometer deflexions) prior to the stop signal were removed following visual inspection. After further exclusion of arrhythmic taps as defined by taps that deviated more than 300 ms from the metronome sound, an average number of 65 ± (SD) 24 trials remained for further analyses. Goniometer traces and the distribution of tap onsets were strongly overlapping prior to successfully vs. unsuccessfully stopped taps (Figure 1). To get a graded measure of stopping performance for correlations, the amount of downward movement measured by the goniometer was quantified as movement extent. It was defined as the extent of the downward movement normalized by the amplitude of the upward movement done before. The time between the stop signal and subsequently failed stops was quantified as median across trials for each patient and then averaged over subjects.

**Electrophysiological recordings**

Bilateral STN local field potentials and EEG was recorded at a sampling frequency of 2048 Hz. EEG electrodes were placed over (or close to if sutures had to be avoided) Fz, Cz, Pz, Oz, C3 and C4 according to the international 10–20 system. Electrooculogram (EOG) was recorded to remove eye blink artefacts in a subsequent procedure. For one patient, EEG channels could not be recorded because large DC drifts caused amplifier saturation. Tap onsets were registered by a force-sensitive resistor measuring the pressure of the finger on its surface. Finger flexion, i.e. the tapping trajectory, was recorded with a goniometer (TMSi Goniometer F35) attached to the index finger over the metacarpophalangeal joint. To capture muscle activity, electromyogram (EMG) was recorded from the first dorsal interosseous muscle (FDI).

**Data pre-processing**

Events for tap and sound onsets were created in Spike 2 (RRID:SCR_000903, Cambridge Electronic Design). After DC component removal (2 s time constant), data were processed further with custom routines in MATLAB (RRID:SCR_001622, v. 2014b, The MathWorks Inc., Natick, Massachusetts). EEG channels were re-referenced to linked earlobes if the latter were recorded (n = 5) or to the average of all EEG channels if not (n = 3). LFP bipolar were computed by subtracting two channels of the same recording electrode (bipolar combinations varied depending on the number of available contacts). Data were down-sampled to 1000 Hz and eye artefacts were removed from the EEG signals by subtracting the filtered EOG (40 Hz low-pass Butterworth filter with a filter order of 6, passed forwards and backwards) after amplitude matching via least-squares optimization (MATLAB function fminocn). Power between 3–40 Hz was obtained by filtering the data into 3 Hz wide frequency bands shifted by 1 Hz (Butterworth, filter order = 6, two-pass, using fieldtrip functions_ft_preproc_lowpassfilter and ft_preproc_highpassfilter [RRID:SCR_004849, Oostenveld et al., 2011]) and calculating the power of the Hilbert transform. Power between 50 and 120 Hz was calculated within 10 Hz wide frequency bands in 2 Hz steps. To reduce noise, power subsequently was temporally smoothed with a 100 ms sliding window. Before exporting the data, it was further down-sampled to 200 Hz. MATLAB analyses scripts for this procedure and subsequent steps to reproduce the figures are provided as source code files (Source code 1). The data can be downloaded from the Oxford University Research Archive: https://ora.ox.ac.uk/objects/uuid:54c00c3d-1809-4a52-bba8-b491b6075f35.

**LFP bipolar selection**

As we recorded from three different electrode models, with multiple contacts of which some may not have been located in the STN, we decided to pre-select the bipolar configuration that recorded the strongest gamma reactivity during regular tapping. We chose to select the contacts based on gamma activity because gamma has been found to be highly focal to the STN (Trottenberg et al., 2006). For the quadripolar (Medtronic 3389) and the unsegmented octopolar model (Boston Scientific DB-2201), bipolars were computed between neighbouring contacts or if channels saturated and thus could not be recorded, the surrounding contacts were instead used for the bipolar subtraction. For the directional contacts (Boston Scientific DB-2202), bipolar combinations were computed.
between the small segmented ones (C2–C7), plus C1 and C8 if more than two of these channels were saturated to increase the likelihood of including activity from the presumably focal gamma source. As power was converted into relative power changes with respect to a baseline, normalized power estimates were relatively comparable despite differently sized contact surfaces or distances between contacts, as was the case for the directional electrode model.

For the selection process, we first computed the 60–90 Hz median power over all taps for each of the multiple bipolar pairs on each electrode in a time window spanning twice the tapping interval around each tap. Then the range between the maximum and minimum of the resulting power time course was divided by the average power within this window, providing the amount of movement-related gamma modulation captured by each bipolar configuration. For each recording electrode only the bipolar configuration with the highest modulation was analysed further. Note that these contacts also recorded significant movement-related beta modulation as shown in Figure 3—figure supplement 1.

**Intersite phase clustering**

Phase-based connectivity between the contralateral STN and the five EEG channels of interest (Fz, C3, C4, Cz, Pz) was computed based on the phase of the Hilbert-transformed filtered signal (bandwidth and frequency shifts as described in Data pre-processing). Intersite phase clustering (ISPC) can be defined over trials or over time. As we did not expect high-frequency oscillations to be phase-locked across trials, we calculated ISPC for each trial over multiple fixed-width windows to get an estimate of changes in ISPC over time. The fixed width was 200 ms for 50–120 Hz and 250 ms for 6–40 Hz. The frequency cut-off was 6 Hz as 250 ms would have included only one and a quarter cycle of a 5 Hz oscillation or even less for lower frequencies.

The window width was chosen to be longer for lower frequencies such that more cycles contributed to the estimate. 250 ms would for example encompass four cycles of a 16 Hz oscillation. ISPC was computed within each of these windows, which were shifted by 10 ms such that the overlapping bins resulted in a smooth image. ISPC was obtained by calculating the length of the average vector of phase differences represented as vectors with length one on a unit circle (Lachaux et al., 2000) based on the following equation (n=number of samples, MATLAB code provided):

\[
\left| \frac{1}{n} \sum_{t=1}^{n} e^{i\phi(t)_{STN}} - e^{i\phi(t)_{EEG}} \right|
\]

The amplitude of the signal thus did not contribute to the ISPC estimate. To assess whether ISPC changed in response to the stop signal, we compared whether it differed significantly from zero after normalizing it by the pre-stop signal period ranging from –350 to 0 ms before the stop cue.

**Statistical testing**

It should be noted that we analysed LFPs from electrode contact pairs of different surface areas (according to electrode type) and EEGs that had different references between subjects. Accordingly, we only considered normalised changes in power to mitigate this variability. All statistical analyses were performed in MATLAB. Correlations between stopping performance (quantified as movement extent after the stop signal) and movement parameters or features in the EEG/LFP were calculated as Spearman’s rank correlation coefficients with bootstrapped confidence intervals (using the Spearman function from the Robust correlation toolbox [Pernet et al., 2012]). To test if correlations with movement parameters differed significantly from zero on a group-level, correlation coefficients were Fisher’s z transformed for each patient and then subjected to a one-sample t-test (n = 9). The maximum correlation with EEG/LFP gamma power (Figure 4—figure supplement 1) was determined for each patient by finding the maximum correlation within 60–90 Hz and 0:156 ms after the stop signal.

Each time-frequency matrix was normalized for each subject and frequency by the average power across all regular taps (excluding tap one and those directly followed by a stop signal) to obtain a relative power percentage change before testing for differences.

Multiple-comparison correction for power or ISPC comparisons in time windows of interest was performed by using a cluster-based permutation procedure (Maris and Oostenveld, 2007, MATLAB code provided): The original paired samples were randomly permuted 2000 times such that each pair was maintained but its order of subtraction may have changed to
create a null-hypothesis distribution. For each permutation, the sum of the z-scores within supra-threshold-clusters (pre-cluster threshold: p<0.05) was computed to obtain a distribution of the 2000 largest suprathreshold-cluster values. If the sum of the z-scores within a suprathreshold-cluster of the original difference exceeded the 95th percentile of the permutation distribution, it was considered statistically significant.

Pairwise comparisons for behavioural data or peak timings were performed using t-tests or Wilcoxon signed-rank tests if the normality assumption (assessed by Lilliefors tests) was violated and if multiple comparisons were made, p-values were subjected to false discovery rate (FDR)-correction.

Acknowledgements
This work was supported by the Medical Research Council (MC_UU_12024/1), and PF was funded by a Clarendon Scholarship and St. John’s College Award. DMH received funding from European Union’s Horizon 2020 research and innovation programme under the Marie Sklodowska-Curie grant agreement number 655605. SL was funded by clinical research training grants from the Wellcome Trust (105804/Z/14/Z). The Unit of Functional Neurosurgery is supported by the Parkinson Appeal UK, and the Monument Trust. The funding bodies played no role in the study design, in the collection, analysis and interpretation of data, in the writing of the report, and in the decision to submit the article for publication.

Additional information
Competing interests
BC: B.C. has received travel support and unrestricted educational grants for organising CPD events from Medtronic, St. Jude Medical and Boston Scientific (manufacturers of DBS electrodes). TZA: T.A. has performed consultancy for and received speaking fees from Medtronic. SL: S.L. has been a participant in a DBS teaching course funded by Medtronic, the manufacturer of the electrodes used in this study. TF: T.F. has received speaking fees and travel support from Medtronic and St. Jude Medical. PL: P.L. has received speaking fees and travel support from Medtronic and St. Jude Medical. LZ: L.Z. has received speaking fees and travel support from Medtronic and St. Jude Medical. PB: P.B. has received fees and non-financial support from Medtronic and personal fees from Boston Scientific. The other authors declare that no competing interests exist.

Funding

| Funder                                           | Grant reference number | Author          |
|--------------------------------------------------|------------------------|-----------------|
| Medical Research Council                         | MC_UU_12024/1          | Peter Brown     |
| Horizon 2020 Framework Programme                 | MSCA 655605            | Damian M Herz   |
| Wellcome                                         | 105804/Z/14/Z          | Simon Little    |
| Parkinson Appeal UK                              |                         | Petra Fischer   |
| Clarendon Scholarship and St. John’s College Award|                         |                 |
| Monument Trust UK                                |                         |                 |

The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.
Author contributions
PF, Conceptualization, Data curation, Software, Formal analysis, Investigation, Visualization, Methodology, Writing—original draft, Project administration, Writing—review and editing; AP, Conceptualization, Software, Methodology, Writing—review and editing; DMH, Methodology, Writing—review and editing, Helped with data recordings; BC, Validation, Writing—review and editing, Helped with patient selection and recruitment. B.C. also assessed anatomical electrode locations; ALG, TZA, Writing—review and editing, Performed DBS surgeries; JF, JH, TF, PL, LZ, Writing—review and editing, Performed DBS surgeries and helped with patient recruitment; SL, Writing—review and editing, Helped with patient recruitment; PB, Conceptualization, Supervision, Funding acquisition, Investigation, Methodology, Writing—review and editing; HT, Conceptualization, Supervision, Investigation, Methodology, Writing—review and editing

Author ORCIDs
Petra Fischer, http://orcid.org/0000-0001-5585-8977
Alexander L Green, http://orcid.org/0000-0002-7262-7297
Peter Brown, http://orcid.org/0000-0002-5201-3044
Huiling Tan, http://orcid.org/0000-0001-8038-3029

Ethics
Human subjects: Patients were recorded after obtaining informed written consent to take part in this study, which was approved by the local ethics committee (Oxfordshire REC A, 08/H0604/58).

Additional files

Supplementary files
• Source code 1. MATLAB analyses scripts to reproduce the figures.
DOI: 10.7554/eLife.23947.022

Major datasets
The following dataset was generated:

| Author(s)       | Year | Dataset title                                                                 | Dataset URL                                                                 | Database, license, and accessibility information |
|-----------------|------|-------------------------------------------------------------------------------|-----------------------------------------------------------------------------|--------------------------------------------------|
| Petra Fischer   | 2017 | Subthalamic nucleus activity during stopping of rhythmic finger tapping        | https://ora.ox.ac.uk/objects/uuid:54c00c3d-1804-4a52-bba8-b491b6075935        | Publicly available at the Oxford University Research Archive (https://ora.ox.ac.uk/) |

References

Alegre M, Lopez-Azcarate J, Obeso J, Wilkinson L, Rodriguez-Oroz MC, Valencia M, Garcia-Garcia D, Guridi J, Artieda J, Jahanshahi M, Obeso JA. 2013. The subthalamic nucleus is involved in successful inhibition in the stop-signal task: a local field potential study in Parkinson's disease. *Experimental Neurology* **239**:1–12. doi: 10.1016/j.expneurol.2012.08.027, PMID: 22975422

Androulidakis AG, Kühn AA, Chen CC, Blomstedt P, Kempf F, Kupsch A, Schneider GH, Doyle L, Dowsey-Limousin P, Hariz MI, Brown P. 2007. Dopaminergic therapy promotes lateralized motor activity in the subthalamic area in Parkinson's disease. *Brain* **130**:457–468. doi: 10.1093/brain/awl358, PMID: 17213215

Aron AR, Poldrack RA. 2006. Cortical and subcortical contributions to stop signal response inhibition: role of the subthalamic nucleus. *Journal of Neuroscience* **26**:2424–2433. doi: 10.1523/JNEUROSCI.4682-05.2006, PMID: 16510720

Aron AR, Robbins TW, Poldrack RA. 2014. Inhibition and the right inferior frontal cortex: one decade on. *Trends in Cognitive Sciences* **18**:177–185. doi: 10.1016/j.tics.2013.12.003, PMID: 24400116

Bastin J, Polosan M, Benis D, Goetz L, Bhattacharjee M, Piallat B, Krainik A, Bougerol T, Chabardès S, David O. 2014. Inhibitory control and error monitoring by human subthalamic neurons. *Translational Psychiatry* **4**:e439. doi: 10.1038/tp.2014.73, PMID: 25203170

Benis D, David O, Lachaux JP, Seigneur et E, Krack P, Fraix V, Chabardés S, Bastin J. 2014. Subthalamic nucleus activity dissociates proactive and reactive inhibition in patients with Parkinson's disease. *NeuroImage* **91**:273–281. doi: 10.1016/j.neuroimage.2013.10.070, PMID: 24368260
Fischer et al. eLife 2017;6:e23947. DOI: 10.7554/eLife.23947

Benis D, David O, Piallat B, Kibler A, Goetz L, Bhattacharjee M, Fraix V, Seigneuret E, Krack P, Chabardès S, Bastin J. 2016. Response inhibition rapidly increases single-neuron responses in the subthalamic nucleus of patients with Parkinson’s disease. Cortex 84:111–123. doi: 10.1016/j.cort.2016.09.006, PMID: 27745848

Bosman CA, Schoffelen JM, Brunet N, Oostenveld R, Bastos AM, Womelsdorf T, Rubehn B, Stieglitz T, De Weerd P, Fries P. 2012. Attentional stimulus selection through selective synchronization between monkey visual areas. Neuron 75:875–888. doi: 10.1016/j.neuron.2012.06.037, PMID: 22958827

Brittain JS, Watkins KE, Joudni RA, Ray NJ, Holland P, Green AL, Aziz T, Jenkinson N. 2012. A role for the subthalamic nucleus in response inhibition during conflict. Journal of Neuroscience 32:13396–13401. doi: 10.1523/JNEUROSCI.2259-12.2012, PMID: 23015430

Brown P, Oliviero A, Mazzone P, Insola A, Tonali P, Di Lazzaro V. 2001. Dopamine dependency of oscillations between subthalamic nucleus and pallidum in Parkinson’s disease. Journal of Neuroscience 21:1033–1038. PMID: 11157088

Brücke C, Bock A, Huebl J, Krauss JK, Schönke T, Schneider GH, Brown P, Kühn AA. 2013. Thalamic gamma oscillations correlate with reaction time in a Go/NoGo task in patients with essential tremor. NeuroImage 75: 36–45. doi: 10.1016/j.neuroimage.2013.02.038, PMID: 23466935

Cassidy M, Mazzone P, Oliviero A, Insola A, Tonali P, Di Lazzaro V, Brown P. 2002. Movement-related changes in synchronization in the human basal ganglia. Brain 125:1235–1246. doi: 10.1093/brain/awf135, PMID: 12023312

Feingold J, Gibson DJ, DePasquale B, Graybiel AM. 2015. Bursts of beta oscillation differentiate postperformance activity in the striatum and motor cortex of monkeys performing movement tasks. PNAS 112: 13687–13692. doi: 10.1073/pnas.1517629112, PMID: 26460033

Fischer P, Tan H, Pogosyan A, Brown P. 2016. High post-movement parietal low-beta power during rhythmic tapping facilitates performance in a stop task. European Journal of Neuroscience 44:2202–2213. doi: 10.1111/ejn.13328, PMID: 27364852

Fogelson N, Pogosyan A, Kühn AA, Kupsch A, van Bruggen G, Speelman H, Tijsen M, Quartarone A, Insola A, Mazzone P, Di Lazzaro V, Limousin P, Brown P. 2005. Reciprocal interactions between oscillatory activities of different frequencies in the subthalamic region of patients with Parkinson’s disease. European Journal of Neuroscience 22:257–266. doi: 10.1111/j.1460-9568.2005.04179.x, PMID: 16029215

Fries P. 2015. Rhythms for cognition: communication through coherence. Neuron 88:220–235. doi: 10.1016/j.neuron.2015.09.034, PMID: 26447583

Gilbertson T, Lalo E, Doyle L, Di Lazzaro V, Cioni B, Brown P. 2005. Existing motor state is favored at the expense of new movement during 13–35 hz oscillatory synchrony in the human corticospinal system. Journal of Neuroscience 25:7771–7779. doi: 10.1523/JNEUROSCI.1762-05.2006, PMID: 16120778

Gross J, Schmitz F, Schnitzler I, Kessler K, Shapiro K, Hommel B, Schnitzler A. 2004. Modulation of long-range neural synchrony reflects temporal limitations of visual attention in humans. PNAS 101:13050–13055. doi: 10.1073/pnas.0404944101, PMID: 15328408

Hamani C, Zaid-Cyr JA, Fraser J, Kaplitt M, Lozano AM. 2004. The subthalamic nucleus in the context of movement disorders. Brain 127:4–20. doi: 10.1093/brain/awp029, PMID: 14607789

Heimer G, Rivlin-Etzion M, Bar-Gad I, Goldberg JA, Haber SN, Bergman H. 2006. Dopamine replacement therapy does not restore the full spectrum of normal pallidal activity in the 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine primate model of Parkinsonism. Journal of Neuroscience 26:8101–8114, doi: 10.1523/JNEUROSCI.5140-05.2012, PMID: 16885224

Huster RJ, Enriquez-Geppert S, Lavallee CF, Falkenstein M, Herrmann CS. 2013. Electroencephalography of response inhibition tasks: functional networks and cognitive contributions. International Journal of Psychophysiology 87:217–233. doi: 10.1016/j.ijpsycho.2012.08.001, PMID: 22906815

Jahanshahi M, Obeso I, Baunez C, Alegre M, Krack P. 2015. Parkinson’s disease, the subthalamic nucleus, inhibition, and impulsivity. Movement Disorders 30:128–140. doi: 10.1002/mds.26049, PMID: 25297382

Jenkinson N, Kühn AA, Brown P. 2013. γ oscillations in the human basal ganglia. Experimental Neurology 245:72–76. doi: 10.1016/j.expneurol.2012.07.005, PMID: 22841500

Joudni RA, Brittain JS, Green AL, Aziz T, Brown P, Jenkinson N. 2012. Oscillatory activity in the subthalamic nucleus during arm reaching in Parkinson’s disease. Experimental Neurology 236:319–326. doi: 10.1016/j.expneurol.2012.05.013, PMID: 22634757

Kempf F, Brücke C, Sallif F, Trottenberg T, Kupsch A, Schneider GH, Doyle Gaynor LM, Hoffmann KT, Vesper J, Währle J, Altenmüller DM, Krauss JK, Mazzone P, Di Lazzaro V, Yelnik J, Kühn AA, Brown P. 2009. Gamma activity and reactivity in human thalamic local field potentials. European Journal of Neuroscience 29:943–953. doi: 10.1111/j.1460-9568.2009.06655.x, PMID: 19291224

Kilavik BE, Zaeppfelm B, Brovelli A, MacKay WA, Riehle A. 2013. The ups and downs of β oscillations in sensorimotor cortex. Experimental Neurology 245:15–26. doi: 10.1016/j.expneurol.2012.09.014, PMID: 23022918

Kononowicz TW, van Rijn H. 2015. Single trial beta oscillations index time estimation. Neuropsychologia 75:381–389. doi: 10.1016/j.neuropsychologia.2015.06.014, PMID: 26102187

Kühn AA, Williams D, Kupsch A, Limousin P, Hariz M, Schneider GH, Yarrow K, Brown P. 2004. Event-related beta desynchronization in human subthalamic nucleus correlates with motor performance. Brain 127:735–746. doi: 10.1093/brain/awt06, PMID: 14960502

Lachaux JP, Rodriguez M, Le van Quyen M, Lutz A, Martinerie J, Varela FJ. 2000. Studying single-trials of phase synchronous activity in the brain. International Journal of Bifurcation and Chaos 10:2429–2439. doi: 10.1142/5021812740001560
Lalo E, Thobois S, Sharott A, Polo G, Mertens P, Pogosyan A, Brown P. 2008. Patterns of bidirectional communication between cortex and basal ganglia during movement in patients with Parkinson’s disease. *Journal of Neuroscience* 28:3008–3016. doi: 10.1523/JNEUROSCI.5295-07.2008, PMID: 18354004

Levy R, Ashby P, Hutchison WD, Lang AE, Lozano AM, Dostrovsky JO. 2002. Dependence of subthalamic nucleus oscillations on movement and dopamine in Parkinson’s disease. *Brain* 125:1196–1209. doi: 10.1093/brain/aww128, PMID: 12023310

Little S, Pogosyan A, Kuhn AA, Brown P. 2012. β band stability over time correlates with Parkinsonian rigidity and bradykinesia. *Experimental Neurology* 236:383–388. doi: 10.1016/j.expneurol.2012.04.024, PMID: 22572590

Litvak V, Eusebio A, Jha A, Oostenveld R, Barnes G, Foltynie T, Limousin P, Zrinzo L, Hariz MI, Friston K, Brown P. 2012. Movement-related changes in local and long-range synchronization in Parkinson’s disease revealed by simultaneous magnetoencephalography and intracranial recordings. *Journal of Neuroscience* 32:10541–10553. doi: 10.1523/JNEUROSCI.0767-12.2012, PMID: 22855804

Magnin M, Morel A, Jeanmonod D. 2000. Single-unit analysis of the pallidum, thalamus and subthalamic nucleus in parkinsonian patients. *Neuroscience* 96:549–564. doi: 10.1016/S0306-4522(99)00583-7, PMID: 10717435

Mallet N, Schmidt R, Leventhal D, Chen F, Amer N, Boraud T, Berke JD. 2016. Arkypallidal cells send a stop signal to Striatum. *Neuron* 89:308–316. doi: 10.1016/j.neuron.2015.12.017, PMID: 26777273

Maris E, Oostenveld R. 2007. Nonparametric statistical testing of EEG- and MEG-data. *Journal of Neuroscience Methods* 164:177–190. doi: 10.1016/j.jneumeth.2007.03.024, PMID: 17517438

Mink JW. 1996. The basal ganglia: focused selection and inhibition of competing motor programs. *Progress in Neurobiology* 50:381–425. doi: 10.1016/S0301-0082(96)00042-1, PMID: 9004351

Nevado-Holgado AJ, Mallet N, Magill PJ, Bogacz R. 2014. Effective connectivity of the subthalamic nucleus-globus pallidus network during Parkinsonian oscillations. *The Journal of Physiology* 592:1429–1455. doi: 10.1113/jphysiol.2013.259721, PMID: 24344162

Ni J, Wunderle T, Lewis CM, Desimone R, Dieter I, Fries P. 2016. Gamma-Rhythmic Gain Modulation. *Neuron* 92:240–251. doi: 10.1016/j.neuron.2016.09.003, PMID: 27667008

Nikolić D, Fries P, Singer W. 2013. Gamma oscillations: precise temporal coordination without a metronome. *Trends in Cognitive Sciences* 17:54–55. doi: 10.1016/j.tics.2012.12.003, PMID: 23287106

Oostenveld R, Fries P, Maris E, Schoffelen JM. 2011. FieldTrip: open source software for advanced analysis of MEG, EEG, and invasive electrophysiological data. *Computational Intelligence and Neuroscience* 2011:1–9. doi: 10.1155/2011/156869, PMID: 21253357

Oswal A, Litvak V, Brücke C, Huebl J, Schneider GH, Kühn AA, Brown P. 2013. Cognitive factors modulate activity within the human subthalamic nucleus during voluntary movement in Parkinson’s disease. *Journal of Neuroscience* 33:15815–15826. doi: 10.1523/JNEUROSCI.1790-13.2013, PMID: 24089489

Perrett CR, Wilcox R, Rousselet GA. 2012. Robust correlation analyses: false positive and power validation using a new open source matlab toolbox. *Frontiers in Psychology* 3:606. doi: 10.3389/fpsyg.2012.00606, PMID: 23335907

Pogosyan A, Kühn AA, Trottenberg T, Schneider GH, Kupsch A, Brown P. 2006. Elevations in local gamma activity are accompanied by changes in the firing rate and information coding capacity of neurons in the region of the subthalamic nucleus in Parkinson’s disease. *Experimental Neurology* 202:271–279. doi: 10.1016/j.expneurol.2006.06.014, PMID: 16905134

Rae CL, Hughes LE, Anderson MC, Rowe JB, Rowe JB. 2015. The prefrontal cortex achieves inhibitory control by facilitating subcortical motor pathway connectivity. *Journal of Neuroscience* 35:786–794. doi: 10.1523/JNEUROSCI.3093-13.2015, PMID: 25589771

Rae CL, Nombela C, Rodríguez PV, Ye Z, Hughes LE, Jones PS, Ham T, Rittman T, Coyle-Gilchrist I, Regenthal R, Sahakian BJ, Barker RA, Robbins TW, Rowe JB. 2016. Atomoxetine restores the response inhibition network in Parkinson’s disease. *Brain* 139:2235–2248. doi: 10.1093/brain/aww138, PMID: 27343257

Ray NJ, Brittain JS, Holland P, Jounidi RA, Stein JF, Aziz TZ, Jenkinson M, Pogosyan A, Kuhn AA, Brown P. 2012. The role of the subthalamic nucleus in response inhibition: evidence from local field potential recordings in the human subthalamic nucleus. *NeuroImage* 60:271–278. doi: 10.1016/j.neuroimage.2011.12.035, PMID: 22098185

Remple MS, Bradenham CH, Kao CC, Charles PD, Neimat JS, Konrad PE. 2011. Subthalamic nucleus neuronal firing rate increases with Parkinson’s disease progression. *Movement Disorders* 26:1657–1662. doi: 10.1002/mds.23708, PMID: 21542021

Rouhinen S, Panula J, Palva JM, Palva S. 2013. Load dependence of β and γ oscillations predicts individual capacity of visual attention. *Journal of Neuroscience* 33:19023–19033. doi: 10.1523/JNEUROSCI.1666-13.2013, PMID: 24285906

Schmidt R, Leventhal DK, Mallet N, Chen F, Berke JD. 2013. Canceling actions involves a race between basal ganglia pathways. *Nature Neuroscience* 16:1118–1124. doi: 10.1038/nn.3456, PMID: 23852117

Schoffelen JM, Oostenveld R, Fries P. 2005. Neuronal coherence as a mechanism of effective corticospinal interaction. *Science* 308:111–113. doi: 10.1126/science.1107027, PMID: 15802603

Swann N, Tandon N, Canolty R, Ellmore TM, McEvoy LK, Dreyer S, DiSano M, Aron AR. 2009. Intracranial EEG reveals a time- and frequency-specific role for the right inferior frontal gyrus and primary motor cortex in stopping initiated responses. *Journal of Neuroscience* 29:12675–12685. doi: 10.1523/JNEUROSCI.3359-09.2009, PMID: 19812342

Swann NC, Cai W, Conner CR, Pieters TA, Claffey MP, George JS, Aron AR, Tandon N. 2012. Roles for the presupplementary motor area and the right inferior frontal gyrus in stopping action: electrophysiological responses
and functional and structural connectivity. *NeuroImage* **59**:2860–2870. doi: 10.1016/j.neuroimage.2011.09.049, PMID: 21979383

Swick D, Ashley V, Turken U. 2011. Are the neural correlates of stopping and not going identical? quantitative meta-analysis of two response inhibition tasks. *NeuroImage* **56**:1655–1665. doi: 10.1016/j.neuroimage.2011.02.070, PMID: 21376819

Tan H, Pogosyan A, Anzak A, Ashkan K, Bogdanovic M, Green AL, Aziz T, Fołytnie T, Limousin P, Zrinzo L, Brown P. 2013. Complementary roles of different oscillatory activities in the subthalamic nucleus in coding motor effort in Parkinsonism. *Experimental Neurology* **248**:187–195. doi: 10.1016/j.expneurol.2013.06.010, PMID: 23778147

Tan H, Pogosyan A, Ashkan K, Cheeran B, FitzGerald JJ, Green AL, Aziz T, Fołytnie T, Limousin P, Zrinzo L, Brown P. 2015. Subthalamic nucleus local field potential activity helps encode motor effort rather than force in parkinsonism. *Journal of Neuroscience* **35**:5941–5949. doi: 10.1523/JNEUROSCI.4609-14.2015, PMID: 25878267

Tomlinson CL, Stowe R, Patel S, Rick C, Gray R, Clarke CE. 2010. Systematic review of levodopa dose equivalency reporting in Parkinson’s disease. *Movement Disorders* **25**:2649–2653. doi: 10.1002/mds.23429, PMID: 21069833

Trottenberg T, Fogelson N, Kühn AA, Kupsch A, Schneider GH, Brown P. 2006. Subthalamic gamma activity in patients with Parkinson's disease. *Experimental Neurology* **200**:56–65. doi: 10.1016/j.expneurol.2006.01.012, PMID: 16499911

Tzagarakis C, Ince NF, Leuthold AC, Pellizzer G. 2010. Beta-band activity during motor planning reflects response uncertainty. *Journal of Neuroscience* **30**:11270–11277. doi: 10.1523/JNEUROSCI.6026-09.2010, PMID: 20739547

Wei W, Wang XJ. 2016. Inhibitory control in the cortico-basal ganglia-thalamocortical loop: complex regulation and interplay with memory and decision processes. *Neuron* **92**:1093–1105. doi: 10.1016/j.neuron.2016.10.031, PMID: 27866799

Wessel JR, Aron AR. 2013. Unexpected events induce motor slowing via a brain mechanism for action-stopping with global suppressive effects. *Journal of Neuroscience* **33**:18481–18491. doi: 10.1523/JNEUROSCI.3456-13.2013, PMID: 24259571

Wessel JR, Aron AR. 2014. Inhibitory motor control based on complex stopping goals relies on the same brain network as simple stopping. *NeuroImage* **103**:225–234. doi: 10.1016/j.neuroimage.2014.09.048, PMID: 25270603

Wessel JR, Ghahremani A, Udupa K, Saha U, Kalia SK, Hodaie M, Lozano AM, Aron AR, Chen R. 2016a. Stop-related subthalamic beta activity indexes global motor suppression in Parkinson’s disease. *Movement Disorders* **31**:1846–1853. doi: 10.1002/mds.26732, PMID: 27474845

Wessel JR, Jenkinson N, Brittain JS, Voets SH, Aziz TZ, Aron AR. 2016b. Surprise disrupts cognition via a fronto-basal ganglia suppressive mechanism. *Nature Communications* **7**:11195. doi: 10.1038/ncomms11195, PMID: 27088156

Xu B, Sandrini M, Wang WT, Smith JF, Sarliss JE, Awosika O, Butman JA, Horwitz B, Cohen LG. 2016. PreSMA stimulation changes task-free functional connectivity in the fronto-basal-ganglia that correlates with response inhibition efficiency. *Human Brain Mapping* **37**:3236–3249. doi: 10.1002/hbm.23236, PMID: 27144466