Propagation of Beta/Gamma Rhythms in the Cortico-Basal Ganglia Circuits of the Parkinsonian Rat

Timothy O. West¹²*, Luc Berthouze³⁴, David M. Halliday⁵, Vladimir Litvak², Andrew Sharott⁶, Peter J. Magill⁶⁷, Simon F. Farmer⁸⁹

¹CoMPLEx, Centre for Mathematics and Physics in the Life Sciences and Experimental Biology, UCL
Department of Physics and Astronomy, Gower Street, London, WC1E 6BT

²Wellcome Trust Centre for Neuroimaging, UCL Institute of Neurology, Queen Square, London, WC1N 3BG

³Centre for Computational Neuroscience and Robotics, University of Sussex, Falmer, UK

⁴Institute of Child Health, UCL, Guildford St., London, WC1N 1EH

⁵Department of Electronic Engineering, University of York, YO10 5DD, UK.

⁶Medical Research Council Brain Network Dynamics Unit, University of Oxford, Oxford, OX1 3TH, United Kingdom.

⁷Oxford Parkinson’s Disease Centre, University of Oxford, Oxford OX1 3QX, United Kingdom.

⁸Department of Neurology, National Hospital for Neurology & Neurosurgery, Queen Square, London WC1N 3BG

⁹Sobell Department of Motor Neuroscience and Movement Disorders, Institute of Neurology, UCL, London WC1N 3BG.

*Corresponding Author

Abstract

Some motor impairments associated with Parkinson’s disease are thought to arise from pathological activity in the neuronal networks formed by the basal ganglia (BG) and motor cortex. To evaluate several hypotheses proposed to explain the emergence of pathological oscillations in Parkinsonism, we investigated changes to the directed connectivity in these networks following dopamine depletion.
We recorded local field potentials in the cortex and basal ganglia of anesthetized rats rendered Parkinsonian by injection of 6-hydroxydopamine (6-OHDA), with dopamine-intact rats as controls. We performed systematic analyses of the networks using a novel tool for estimation of directed neural interactions, as well as a conditioned variant which permits the analysis of the dependence of a connection upon a third reference signal. We find evidence of the dopamine dependency of both low beta (14-20 Hz) and high beta/low gamma (20-40 Hz) directed interactions within the BG and cortico-BG networks. Notably, 6-OHDA lesions were associated with enhancement of the cortical “hyper-direct” connection to the subthalamic nucleus (STN), as well the STN’s feedback to the cortex and striatum. We find beta synchronization to be robust to conditioning using signals from any one structure. Conversely, we find that high beta/gamma drive from the cortex to subcortical regions is weakened by 6-OHDA lesions and is susceptible to conditioning. Furthermore, we provide evidence that gamma is routed from striatum in a pathway that is independent of STN. These results further inform our understanding of the substrates for pathological rhythms in salient brain networks in Parkinsonism.

Keywords
Parkinson’s disease; basal ganglia; local field potential; synchronization; connectivity;

New & Noteworthy
We present a novel analysis of electrophysiological recordings in the cortico-basal ganglia network with the aims of evaluating several hypotheses concerning the origins of abnormal brain rhythms associated with Parkinson’s disease. We present evidence for changes in the directed connections within the network following chronic dopamine depletion in rodents. These findings speak to the plausibility of a “short-circuiting” of the network that gives rise to the conditions from which pathological synchronization may arise.

Introduction
The basal ganglia (BG) are host to a small but important cluster of dopaminergic neurons that act to modulate the activity of a large re-entrant network that comprises the cortico-basal ganglia-thalamo-cortical circuit (DeLong and Wichmann 2010; Lanciego et al. 2012). Investigation of the structure of this network (Smith et al. 1998; Bolam et al. 2000) has led to a canonical view of the circuit (depicted in figure 1) from which multiple process theories of BG function have arisen (for a review, see Schroll and Hamker (2013)).

Recent theory concerning the organisation of brain networks and communication within them via synchronized oscillations (Varela et al. 2001; Fries 2005, 2015; Bressler and Menon 2010; Thut et al. 2012) has emphasised the importance of understanding the dynamics of these networks beyond that afforded by studying structure alone (Deco et al. 2008, 2012). Neural oscillations and their
synchronization can be detected across multiple scales of brain activity, from single neuronal
discharges up to the level of mesoscale neural ensembles such as those measured in the local field
potential (LFP) or electrocorticogram (ECoG). Pathological oscillations and their synchronization
across disparate regions of the brain have been detected in brain disorders such as Parkinson’s disease
(PD), schizophrenia, and epilepsy, leading to the hypothesis that the oscillations themselves bear a
causal role in their associated behavioral impairments (Schnitzler and Gross 2005; Uhlhaas and Singer
2006; Hammond et al. 2007).

Excessive beta oscillations (14-30 Hz) in the BG associated with dopamine depletion have been
observed reliably in untreated patients with PD (Levy et al. 2000; Brown et al. 2001; Weinberger et
al. 2006; Hammond et al. 2007). Beta rhythms are attenuated by treatments such as dopamine
replacement therapy (Kühn et al. 2006; Weinberger et al. 2006; West et al. 2016; Beudel et al. 2017;
Levy 2002) and deep brain stimulation (DBS) (Ray et al. 2008; Eusebio et al. 2011; Whitmer et al.
2012) in a way that correlates with the degree of the improvement of motor symptoms. This has
strengthened the argument that the pathological beta rhythms are directly related to the functional
impairment seen in patients (Hanslmayr et al. 2012; Brittain and Brown 2014). Furthermore, gamma
activity in the motor system has been hypothesized to be prokinetic (Schoffelen et al. 2005). In PD, the
spectral power of multiunit recordings from STN at 40-90 Hz have been demonstrated to be
negatively correlated with bradykinetic symptoms in patients (Sharott et al. 2014).

The pathological oscillations observed in mesoscale electrophysiological signals are a direct
consequence of changes to the underlying networks of neuronal ensembles that generate them. This
understanding has led to the re-classification of multiple neurological diseases such as PD or
Tourette’s as ‘circuit disorders’ (DeLong and Wichmann 2010). Knowledge of how dopamine
depletion results in changes to the network, and the subsequent emergence of pathological synchrony
is likely to lead to a better understanding of the causes of impairment and its treatments (Shen et al.
2008; Schroll et al. 2014). Thus improving insight into how changes to connections in the network
lead to the emergence of pathological dynamics remains an important line of enquiry (Wichmann and
DeLong 1999; Dostrovsky and Bergman 2004; Holgado et al. 2010).

Previous work aiming to understand the origins of the pathological beta rhythm has involved
systematic lesioning of the BG network (Ni et al. 2000; Tachibana et al. 2011), computational
modelling (Holgado et al. 2010; Moran et al. 2011; Marreiros et al. 2013; Nevado-Holgado et al.
2014; Pavlides et al. 2015), and techniques from signal analysis (Sharott et al. 2005a; Mallet et al.
2008a, 2008b; Litvak et al. 2011). In this paper, we take the latter approach and, through analysis of
neural recordings, aim to infer the changes in neural transmission that occur in cortico-BG circuits
following chronic dopamine depletion.
Connectivity between parts of the brain can be inferred from the statistical dependencies that arise due to neural transmission: we refer to this as functional connectivity (Friston 2011). Previous studies have aimed to describe ‘effective’ connectivity (i.e. causal interactions) within this network and have employed the dynamic causal modeling (DCM) framework in order to do so. To date, two such studies have utilised the inversion of biophysical models upon cross spectral densities from recordings in either anaesthetised 6-OHDA lesioned rats (Moran et al. 2011), or awake DBS patients (Marreiros et al. 2013). Both found evidence for the strengthening of the cortico-subthalamic connection (termed the ‘hyper-direct’ pathway (Nambu et al. 2002)) in the dopamine-depleted state.

From this work amongst others, several hypotheses have arisen concerning the emergence of pathological beta rhythms as a result of the dopamine depletion associated with PD (for a review see Holgado et al. 2010). These include the dopamine-dependent modulation of recurrent loops within the network, either between the reciprocally-coupled network of neurons of the subthalamic nucleus (STN) and the external globus pallidus (GPe) (Plenz and Kital 1999; Bevan et al. 2002; Terman et al. 2002; Holgado et al. 2010); or of a longer loop involving feedback from BG output nuclei to the cortex via thalamo-cortical tracts (Leblois et al. 2006; Pavlides et al. 2012, 2015). Alternatively, it has been proposed that dopamine depletion disrupts mechanisms which regulate the gain of cortical afferents to the BG and somehow disrupt striatal outflow (Brown 2007; Hammond et al. 2007).

Here, using a recently described non-parametric (model-free) signal analysis technique (Halliday et al., 2015), we study the effects of dopamine depletion upon neural connectivity in the network formed by elements of the BG and motor cortex in 6-OHDA-lesioned and dopamine-intact control rats. We employ this method as a measure of directed functional connectivity (hereon shortened to directed connectivity). It is a model-free estimate that makes no assumptions as to the causes of the data (Bastos and Schoffelen 2016), only that temporal precedence implies a driving neuronal influence (please see later sections for discussion). Furthermore, we use a a multivariate extension of the framework (Halliday et al. 2016) in order to determine whether the interaction between two areas shares correlation with activity recorded at a third structure in the network. This approach provides insight into frequency-specific directional connectivity and the dependence of coupled signals upon activity at a third region. By recording LFPs and ECoG in 6-OHDA–lesioned animals and dopamine-intact controls we aim to identify changes to connectivity that occur as a result of the loss of dopamine from these circuits. Our findings are interpreted within the context of the canonical circuit (figure 1), other existing models of basal ganglia connectivity, and several hypotheses concerning the generation and propagation of pathological beta rhythms in the network.
Methods

Experimental Data

Electrophysiological Recordings

Experimental procedures were carried out on adult male Sprague-Dawley rats (Charles River, Margate, UK) and were conducted in accordance with the Animals (Scientific Procedures) Act, 1986 (UK). Recordings were made in eight dopamine-intact control rats (288–412 g) and nine 6-OHDA-lesioned rats (285–428 g at the time of recording), as described previously (Magill et al. 2006; Mallet et al. 2008a, 2008b; Moran et al. 2011). Briefly, anaesthesia was induced with 4% v/v isoflurane (Isoflo, Schering-Plough Ltd., Welwyn Garden City, UK) in O2, and maintained with urethane (1.3 g/kg, i.p.; ethyl carbamate, Sigma, Poole, UK), and supplemental doses of ketamine (30 mg/kg; Ketaset, Willows Francis, Crawley, UK) and xylazine (3 mg/kg; Rompun, Bayer, Germany).

The ECoG was recorded via a 1 mm diameter steel screw juxtaposed to the dura mater above the right frontal (somatic sensory-motor) cortex (4.5 mm anterior and 2.0 mm lateral of Bregma and approximately centred over M2, (Paxinos and Watson 2007)) and was referenced against another screw implanted in the skull above the ipsilateral cerebellar hemisphere. Raw ECoG was band-pass filtered (0.3–1500 Hz, -3 dB limits) and amplified (2000x; DPA-2FS filter/amplifier: Scientifica Ltd., Harpenden, UK) before acquisition. Extracellular recordings of LFPs in the dorsal striatum (STR), GPe and STN were simultaneously made in each animal using ‘silicon probes’ (NeuroNexus Technologies, Ann Arbor, MI). Each probe had one or two vertical arrays of recording contacts (impedance of 0.9–1.3 MΩ measured at 1000 Hz; area of ~400 μm²).

The same probe was used throughout these experiments but it was cleaned after each experiment in a proteolytic enzyme solution to ensure that contact impedances and recording performance were not altered by probe use and re-use. Monopolar probe signals were recorded using high-impedance unity-gain operational amplifiers (Advanced LinCMOS: Texas Instruments, Dallas, TX) and were referenced against a screw implanted above the contralateral cerebellar hemisphere. After initial amplification, extracellular signals were further amplified (1000x) and low-pass filtered at 6000 Hz using programmable differential amplifiers (Lynx-8: Neuralynx, Tucson, AZ). The ECoG and probe signals were each sampled at 17.9 kHz using a Power1401 Analog-Digital converter and a PC running Spike2 acquisition and analysis software (Cambridge Electronic Design Ltd., Cambridge, UK). Recording sites in the BG were verified by post hoc histology, as described previously (Magill et al. 2006; Mallet et al. 2008a, 2008b).

Neuronal activity was recorded during episodes of spontaneous ‘cortical activation’, which contain patterns of activity that are similar to those observed during the awake, behaving state (Steriade 2000). Cortical activation was defined according to ECoG activity. Neuronal activity patterns present under this anaesthetic regime may only be qualitatively similar to those present in the un-anesthetized...
brain. However, the urethane-anesthetized animal still serves as a useful model for assessing ensemble
dynamics within the basal ganglia. Indeed, in 6-OHDA-lesioned animals, exaggerated beta
oscillations emerge in cortico-basal ganglia circuits during activated brain states thus accurately
mimicking the oscillatory activity recorded in awake, un-medicated PD patients. Examples of the raw
and pre-processed electrophysiological signals as well as the corresponding power spectra for control
and lesioned animals are shown in figure 2.

6-Hydroxydopamine Lesions of Dopamine Neurons
Unilateral 6-OHDA lesions were carried out on 200–250 g rats, as described previously (Mallet et al.
2008a, 2008b). Twenty-five minutes before the injection of 6-OHDA, all animals received a bolus of
desipramine (25 mg/kg, i.p.; Sigma) to minimize the uptake of 6-OHDA by noradrenergic neurons
(Schwarting and Huston 1996a). Anaesthesia was induced and maintained with 4% v/v isoflurane (see
above). The neurotoxin 6-OHDA (hydrochloride salt; Sigma) was dissolved immediately before use
in ice-cold 0.9% w/v NaCl solution containing 0.02% w/v ascorbate to a final concentration of 4
mg/ml. Then 3 ml of 6-OHDA solution was injected into the region adjacent to the medial substantia
nigra (4.5 mm posterior and 1.2 mm lateral of bregma, and 7.9 mm ventral to the dura. The extent of
the dopamine lesion was assessed 14–16 days after 6-OHDA injection by challenge with apomorphine
(0.05 mg/kg, s.c.; Sigma) (Schwarting and Huston 1996b). The lesion was considered successful in
those animals that made >80 net contraversive rotations in 20 min. Electrophysiological recordings
were carried out ipsilateral to 6-OHDA lesions in anesthetized rats 21–42 days after surgery, when
pathophysiological changes in the basal ganglia are likely to have levelled out near their maxima
(Mallet et al. 2008a).

Data Acquisition and Analysis
Data Conversion and Pre-Processing
To isolate LFPs and ECoGs, all electrophysiological data were down-sampled from a hardware native
17.9 kHz to 250 Hz using Spike2 acquisition and analysis software (version 4; Cambridge Electronic
Design Ltd., Cambridge, UK). Data were then imported from Spike2 into MATLAB (The
Mathworks, Nantucket, MA, USA) where they were analysed using custom scripts utilizing routines
from the Fieldtrip software package (packaged in SPM 12.3) (Oostenveld et al. 2011), as well as
Neurospec (http://www.neurospec.org/). Data were pre-processed as follows: i) data were first
truncated to remove 1 second from either end of the recording, ii) mean subtracted; iii) band-passed
filtered with a finite impulse response, two-pass (zero-lag) filter designed such that the filter order is
rounded to the number of samples for 3 periods of the lowest frequency, between 4-100 Hz; iv) data
were then split into 1 second epochs; v) each epoch was subjected to a Z-score threshold criterion
such that epochs containing any high amplitude artefacts were removed. Examples of outcomes from
this pre-processing are shown in figure 2.
Signal Analysis Techniques

Power analyses were made using the averaged periodogram method across 1 second epochs and using a Hanning taper to reduce the effects of spectral leakage. Frequencies between 49-51 Hz were removed so that there was no contribution from 50 Hz line noise. We computed spectral power, imaginary coherence, directional analyses (NPD), and their conditioned variants using the Neurospec analysis toolbox.

Non-zero Phase Lag Functional Connectivity Analysis: Imaginary Coherence

The standardly used spectral coherence (Halliday et al. 1995) is sensitive to spurious correlations resulting from instantaneous volume conduction between the two signals of interest (Bastos and Schoffelen 2016). In order to circumvent this issue, several methods have been developed such as taking the imaginary part of coherence (Nolte et al. 2004), the phase lag index (PLI) (Stam et al. 2007), or the weighted phase lag index (Vinck et al. 2011). For this study, we use arguably the simplest method available (imaginary coherence- iCOH) that is derived from the complex coherency. This measure shares the property of disregarding zero lag correlations with the non-parametric directionality analysis that we later introduce for estimates of directed connectivity. We also note the concerns in Stam et al. (2007) on the validity of imaginary coherence analysis, so include additional analyses based on non-parametric directionality.

Non-Parametric Directionality

Estimates of directed connectivity were computed using non-parametric directionality (NPD) (Halliday 2015) as implemented in the Neurospec software package. This analysis combines Minimum Mean Square Error (MMSE) pre-whitening with forward and reverse Fourier transforms to decompose coherence estimates at each frequency into three components: forward, reverse and zero lag. These components are defined according to the corresponding time lags in the cross-correlation function derived from the MMSE pre-whitened cross-spectrum. The advantage of this approach is that it allows us to decompose the signal into distinct forward and reverse components of coherence separate from the zero-lag (or instantaneous) component of coherence which is assumed to reflect volume conduction. The method uses temporal precedence to determine directionality. For example, STN activity lagging M2 activity results in a significant forward component of coherence between M2 and STN (when M2 is assumed to be the reference), whereas STN activity leading M2 activity results in a significant reverse component of coherence.

In addition we used a multivariate extension of the non-parametric directionality (Halliday et al. 2016), which allows the directional components of coherence to be conditioned on a third signal. This analysis decomposes the partial coherence into the same three directional components: forward, reverse and zero-lag. This analysis can indicate if the signals reflected in the correlation are common to other parts of the network. For example, the partial correlation between A and B with C as
predictor can be used to determine if the flow of information from A → B is independent of area C, or
whether the flow of information is A → C → B, in which case the partial coherence between A and B
with C as predictor should be zero. The partial coherence can also be used to investigate if the flow of
information is C → A and C → B, or if it is A → B → C or C → A → B, which for the latter case the
partial coherence, and any directional components should be zero.

This assumes that the conditioning signal, C, is representative of the activity in the relevant brain area.
If the signal, C, only captures part of the activity in the brain area then the partial coherence estimate
may still have residual features. The most robust interpretation of the partial coherence and
multivariate non-parametric directionality is where the partial coherence (and any directional
components) are not significant compared to the directional components for the ordinary coherence.

Statistics
In order to make statistical comparisons of power, connectivity and directionality spectra between
lesioned and control animals we used cluster based permutation testing (Maris and Oostenveld 2007)
which avoids introducing bias through the prior specification of frequency bands. It requires no
assumption of normality, and affords a correction for the multiple comparison problem by controlling
the family-wise error rate via an approximation of the reference distribution for the test statistic. For
more details see Maris (2012). The cluster-forming threshold was p<0.05 and the permutation test
threshold was set at p<0.025 (as it is a two-sided test). The number of permutations was set to 5000
which tenders a lowest possible P-value equal to 0.0004. Cluster statistics were computed using the
‘ft_freqstatistics’ routine in Fieldtrip.

Results
Spectral Power
Examples of spectra computed from LFP and ECoG signals recorded in individual animals can be
seen in figure 2 (B and D). All the 6-OHDA-lesioned rats demonstrated a clear peak in the spectra in
the range 18-22 Hz (encompassing low beta/lower end of high beta frequencies) for LFP recordings
across all subcortical recording sites as well as for the sensorimotor ECoGs. In some animals, cortical
beta was weaker than that observed subcortically. None of the LFP data from control animals
contained a similar beta peak in the spectra although some (4 of 8) showed a wide, low amplitude
peak around 20-40 Hz most prominent in the recordings at M2 (an example of which is seen in figure
2B). Analysis of the group averaged spectra (figure 3) shows that the beta peak is significantly
increased in the dopamine-depleted animals. Cluster-based permutation testing demonstrated
significant differences in group level spectra between control and lesion conditions with clusters
showing increases in power in the STN (16-21 Hz, P=0.023), GPe (17-22 Hz, P=0.009) and STR (18-
21 Hz, P=0.019) for the dopamine-depleted animals. Cluster-based permutation did not reveal
significant increases in beta power in the ECoG signals recorded in the M2 of lesioned rats when compared to the controls (although a positive cluster was found that exceeded the cluster forming threshold but was not significant under permutation testing). No differences between lesioned and control animals were found for frequencies >22 Hz in any structures.

Functional Connectivity: Imaginary Coherence (iCOH)

Initial analyses of connectivity of the recorded LFPs using magnitude squared coherence showed large magnitude (>0.9) wideband (0 - 60Hz) coherences that were indicative of a large field spread effect (data not shown). This was most apparent in subcortical-subcortical analyses but was also detected for cortical-subcortical pairings. To estimate coherence avoiding contamination by volume conduction we opted to calculate non-zero phase lag correlations using the imaginary part of coherence (iCOH) (see figure 4).

We found that activity in the low beta range (14-20 Hz) associated with 6-OHDA dopamine depletion is spread diffusely across the network with all inter-regional comparisons showing a significant beta peak in the iCOH spectrum. Notably, the strongest beta band involved STN, with STN/STR and STN/GPe pairs both showing coefficients greater than 0.2. Within region connectivity (i.e. STN contact 1 to contact 2) was found to be present in this range for only recordings within STN or GPe, where there is a clear beta peak.

Analysis of statistical differences using the cluster based permutation testing between control and lesioned animals showed significant increases of iCOH in the beta band in the lesioned animals and for 5/10 pairs tested: STN/STR (14-21 Hz, P=0.004), STN/STN (19-25 Hz, P = 0.010), GPe/STR (14-16 Hz, P = 0.009), GPe/STN (14-21 Hz, P=0.005), and GPe/GPe (19-23 Hz, P=0.004). Notably, no pairs involving M2 showed significant modulations of beta-band activity with dopaminergic state as detectable with the cluster statistics. Taken generally, these results are indicative of widespread, non-zero lag, low beta-band connectivity across the entire cortico-BG network that is increased in the dopamine-depleted rats.

In the control rats, connectivity in the beta range was reduced. Instead, there was wide-band iCOH in the high beta/low gamma bands, ranging from 20 Hz to 50 Hz in most cases but up to 70 Hz for the STN/M2 interactions. The majority of gamma interactions where iCOH was high (> 0.2) were found in connections involving the STN. Additionally, iCOH in these bands is evident between GPe/M2 and GPe/STR although this was weaker (at around ~0.1) than connections analysed with pairs involving the STN. iCOH in these bands is present in both the lesioned and control animals and does not show a strong modulation by dopamine as evidenced by the lack of significant clusters in the permutation tests for these bands.
Non-parametric Directionality (NPD)

We next investigated directed connectivity between recorded regions. The results of the analysis using the NPD measure are presented in figure 5. The iCOH and the sum of the non-instantaneous parts (forward and backward) of the NPD are similar, and both methods reveal similar patterns of connectivity (data not shown). Analysis of the instantaneous NPD in isolation demonstrated the existence of high amplitude, wide-band interactions that were similar to those found with magnitude squared coherence (data not shown), and are likely due to zero-phase field spread of activity between recordings. Analyses of directional interactions of the LFPs and ECoG hereon will use the forward and backward components of the NPD to discern directional connectivity between structures.

We observed that directional interactions of low beta-band activity are predominant in the direction leading from M2 and descending the hierarchy of the BG. This effect is most clear in the dopamine-depleted animals. Interestingly we noted a significant difference in the cortical-subthalamic beta band interaction between lesioned and control animals only in the feedback connection STN → M2 (16-18 Hz, P=0.018) which would suggest that STN feedback to M2 is strengthened in the dopamine-depleted state. In the case of the STN/GPe circuit, and unlike iCOH, the non-instantaneous components of NPD do not show 6-OHDA related increases in beta coupling in either direction for the lesioned rats. Rather, NPD suggests a directional asymmetry in activity in the high beta/gamma band with forward connections from GPe → STN connection far stronger than in the reverse direction (cluster statistics testing differences between forward and backward spectra in the 6-OHDA recordings: 4-43 Hz, P<0.001). Notably we see a feedback in the STN → STR that is only present in the lesion condition, a feature that will be relevant with respect to results later in this article.

The pattern of activity in the high beta/gamma range between cortical and subcortical regions appeared to be principally cortically leading with interactions in the 20-40 Hz range being most prominent in the dopamine-intact control rats (top row of figure 5). Cluster-based permutation analysis shows a significant increase in the high/gamma M2 → GPe NPD in the control vs the lesion condition (25-30 Hz, P=0.017). High beta/gamma connections from subcortical structures feeding back to M2, are weaker than the forward connections, but are still present for striatal and pallidal feedbacks to M2 (first column, row 2 and 4, figure 5). Again, there is a clear peak in the high beta NPD from STN to STR in the lesioned animals although a dependence on dopamine was not seen to be significant when cluster statistics were applied. Finding of NPD from STN to STR does not accord the canonical circuit (Figure 1) but may instead imply feedback to striatum via subcortical thalamo-striatal loops that will be discussed later.

Inferring Routing of Brain Rhythms: Partialized Non-Parametric Directionality

We repeated the NPD analysis as before but this time by systematically partialising out each structure by conditioning the analysis on the LFPs or ECoG recorded at other regions in the network. We again
employed cluster statistics to determine significant differences between the non-conditioned NPD spectra (presented in the previous section) and the conditioned variant shown in this section of the results.

Conditioning the NPD using Local Field Potentials Recorded from the STN

We first conducted a partialization of the NPD estimate using LFPs recorded from within the STN (figure 6). Conditioning with signals from the STN does not remove beta connectivity between the remaining structures in the network although it does weaken the majority of comparisons in the control (6 of 6 comparisons) but not the lesion (2 of 6 comparisons) experiments (see figure 6, red and blue bars respectively). Cluster statistics indicate that the following NPDs for the control experiments were significantly reduced by conditioning with the STN signal: M2 → STR (14-33 Hz, P<0.001), M2 → GPe (14-33 Hz, P<0.001; 37-49 Hz, P=0.010), STR → GPe (10-49 Hz, P<0.001), GPe → STR (18-49 Hz, P<0.001) as well as feedback connections (returning to cortex): STR → M2 (14-27 Hz, P<0.001), GPe → M2 (18-49 Hz, P<0.001). Furthermore, conditioning the NPD with the signal from STN does not disrupt the 6-OHDA associated increases of M2 input to either the STR (14-21 Hz, P<0.001) or GPe (14-21 Hz, P<0.001). We also found in the dopamine-depleted state that there was increased (relative to the controls) feedback to M2 from both GPe (16-20 Hz, P=0.013) and STR (16-20 Hz, P=0.006).

Notably we observed some separation in the effects of the conditioning between the control and lesion experiments. In the control animals conditioning the NPD on LFPs recorded at STN acted to reduce activity in a wide band (~12-40 Hz) for the forward connections (flowing down the indirect pathway; see figure 1), whilst the return connections (STR → M2, and GPe → M2) were only affected by conditioning a tighter band corresponding to low beta. Lesioned animals only showed reductions at higher frequencies (~24-45 Hz, high beta/low gamma) and only between GPe and STR. We observed that conditioning of the NPD with the STN signal acted to significantly reduce interactions between STR and GPe in both the forward (23-49 Hz, P<0.001) and reverse (27-49 Hz, P=0.001) directions.

Conditioning the NPD using Local Field Potentials Recorded from the GPe

Next, we performed the NPD analysis of recorded signals but this time conditioning the interactions with LFPs recorded from within the GPe (figure 7). We found that the conditioning had the effect of reducing NPD estimates in 6 out of 6 possible connections in the controls and 3 out of 6 in the 6-OHDA-lesioned rats. Most notably we found that the conditioning significantly attenuated (when compared to the unconditioned NPD) the low beta band interaction in the M2 → STR connection for both recordings made in control (14-39 Hz, P<0.001) and lesioned (14-21 Hz, P<0.001) animals. Secondly, we found a reduction of interactions between STR → STN across a wide range of frequencies, again for both control (6-49 Hz, P<0.001) and lesioned (4-49 Hz, P<0.001) recordings suggesting signal routing is strongly mediated by GPe in accordance with the canonical indirect
pathway. Interestingly we found that although beta NPD in the M2 → STN connection was attenuated by conditioning in the control recordings, for the 6-OHDA recordings the prominent low beta peak in the NPD remained and no significant effect of conditioning was observed. Similarly, the STN → M2 feedback also retained a sharp beta peak that remained significantly increased in recordings corresponding to the 6-OHDA lesion experiments (14-20 Hz, P=0.002). Additionally, we found that when conditioning the NPD with the GPe signal, we found that the STR→M2 demonstrated a significant increase in strength in the 6-OHDA animals (16-21 Hz, P<0.001).

In the high beta/gamma band we found that conditioning with GPe has a large effect in attenuating the NPD in the forward connections (from M2 descending the indirect pathway) in the control animals: M2 → STR (14-39 Hz, P<0.001), M2 → STN (16-49 Hz, P<0.001), and STR → STN (6-49 Hz, P<0.001). In the lesion animals only, 2 of the 6 comparisons made with NPD were significantly attenuated in the 20-50 Hz range: STR → STN (4-49 Hz, P<0.001) and STN → STR (21-27 Hz, 31-45 Hz, P=0.003). This would imply that interactions in both directions between STN and STR are mediated via GPe.

Conditioning the NPD using Local Field Potentials Recorded from the STR

The third set of analyses used the local field potentials recorded at the STR to condition the NPD estimates (figure 8). We found that this had the effect of destroying large parts of the descending interactions (connections from M2 descending the hierarchy of the indirect pathway) in the control animals, namely for M2 → GPe (16-37 Hz, P<0.001) and M2 → STN (16-37 Hz, P<0.001). In the lesion recordings, the effect of conditioning split into two ways: 1) The interactions between the STN/GPe were significantly reduced across a very wide band ranging low-beta to gamma frequencies in both the STN → GPe (8-49 Hz, P<0.001) and GPe → STN (6-49 Hz, P<0.001) condition. 2) Interaction in the “hyper-direct” M2 → STN connection was not attenuated, although the M2 → GPe (likely routed at least in part via the indirect pathway) was suppressed by conditioning with the striatal signal (18-24 Hz, P=0.001). This peak is also seen in the feedback connection from STN → M2 where the significant 6-OHDA associated increase in beta feedback reported in previously analysis was found to remain (18-20 Hz, P=0.012).

Similar to the NPD estimates conditioned with signals recorded at GPe, we found that the conditioning with LFPs recorded at STR acts to largely remove the high beta/gamma interactions. In the M2 → GPe connection we found that high beta gamma activity was attenuated by conditioning in the control condition (16-37 Hz, P<0.001) as well as a significant condition dependent 6-OHDA associated suppression of activity in this band was observed (27-37 Hz, P=0.001; 41-45 Hz, P=0.005). Additionally, we also found that feedback in this frequency range (for control recordings) from GPe → M2 was significantly attenuated by conditioning with striatal signals (14-41 Hz, P<0.001; 45-49 Hz, P<0.001).

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Furthermore, we found that activity in this feedback connection was significantly increased in the controls when compared to the 6-OHDA lesioned animals (35-41 Hz, P=0.002).

Conditioning NPD Using Field Potentials Recorded from M2

The final analyses utilized ECoG signals recorded from the M2 to condition the NPD estimates (results in figure 9). We found that the NPD estimates conditioned on M2 are greatly flattened and devoid of distinct peaks at either low beta or high beta/gamma frequencies that were seen typically in the other analyses. Altogether 5 of 6 NPD spectra had no distinct spectral peaks. When testing for significant attenuation of NPD following conditioning we found that only control recordings were found to be significantly attenuated. However, the loss of features found in the unconditioned NPD (such as beta or gamma peaks) were equivalent for both the control and 6-OHDA recordings.

When testing for the effects of 6-OHDA, we found that the STN → STR connection was significantly altered. We observed a broad peak from 20-40 Hz in the lesion recordings that was not attenuated by M2 conditioning and demonstrated a significant increase in strength associated with dopamine depletion (21-27 Hz, P=0.010).

Summary of Connectivity Analyses

Using recordings made in control and lesioned rats, we identified functional connectivity between cortical and BG sites that involved either low beta or high beta/gamma oscillations. Broadly speaking, we found that gamma connectivity is sensitive to the conditioning of structures upstream of the STN, particularly GPe and STR, which removes any detectable gamma from the spectra. In contrast, beta connectivity is robust to partializing using LFPs of any single structure. Cortico-subthalamic connectivity in the beta range is unaffected by partialising of GPe or STR, suggesting M2/STN low beta connectivity is not routed via the indirect pathway. In the next section, we will outline several putative models of oscillatory dynamics and present evidence from our analyses that either support or weaken the plausibility of each model.

Discussion

Hypotheses and evaluation of evidence for signal propagation in the network

We have undertaken a systematic analysis of a dataset involving multisite LFP recordings of the cortico-basal ganglia circuit that contains data from a set of dopamine-intact control rats and another set of rats with chronic dopamine depletion induced by a unilateral injection of 6-OHDA. We will next discuss evidence for competing theories of the propagation of oscillatory activity across the Parkinsonian cortico-basal ganglia circuit.
Mechanisms of the Flow of Beta Rhythms in the Basal Ganglia Circuit

Here we will evaluate the evidence provided by the analyses reported here in light of a number of proposed theories concerning the generation and propagation of beta-band activity in the network and the changes that occur during dopamine depletion that lead to its amplification. This work is summarised in table 1.

Hypothesis 1: Dopamine depletion in the basal ganglia induces increased beta resonance in the cortical/STN “long-loop”.

Previous authors have suggested that pathological beta rhythms are generated from the strengthening of a long cortical feedback loop that returns from basal ganglia output nuclei via the thalamus. Strengthened coupling is proposed to facilitate pathological resonance at beta frequencies (Brown 2007; van Albada and Robinson 2009; Dovzhenok and Rubchinsky 2012; Pavlides et al. 2015). The first step towards verifying the plausibility of this hypothesis involves determining whether there is indeed functional connectivity between STN and M2 in the beta band, and whether this occurs independently of the cortico-striatal inputs to the indirect pathway.

Analysis of the iCOH for the M2/STN pairing suggests that functional connectivity in the beta band is significantly strengthened in the lesioned animals compared to controls (figure 4). Analysis with NPD has demonstrated that there is a beta peak in the directed coherence in the low beta range in the forward M2 → STN connection for both the control and 6-OHDA animals. However, in the lesioned animals, the feedback connection (STN → M2) is significantly strengthened over that measured in the controls. Neither the hyper-direct M2 → STN connection, nor its feedback (STN → M2) are diminished by either conditioning with the signals from the GPe or STR in the lesioned animals (figure 7 and figure 8). This suggests that outflow from STN is routed independently of STR or GPe, likely directly via the BG output nuclei. In contrast, in control rats, NPD of the feedback connections at beta frequencies are significantly decreased by conditioning with the STR signal in the forward (M2 → STN), and backward (STN → M2) directions, suggesting that in the dopamine-intact anaesthetised state, any residual beta band activity is routed via STR, whilst the hyper-direct pathway is relatively quiescent. These findings support the idea that the dopamine-depleted state is associated with a strengthening of the hyper-direct pathway and its subsequent feedback from STN to cortex.

We also provide evidence of multiple coexisting pathways for propagation of beta oscillations. Notably, it was found that conditioning of the NPD with LFPs recorded from the STN (figure 6) does not act to remove the 6-OHDA lesion associated beta NPD in the structures ‘upstream’ of the STN (i.e. the STR and GPe). NPD in the low beta range is significant in both the forward and backward directions along parts of the network involving either M2, STR or GPe. Most notably, we find that striatal-subthalamic interactions are strongly modulated by the GPe signal, a finding in line with the canonical indirect pathway. Future work to validate the long-loop hypothesis would involve the
conditioning of the STN \(\rightarrow\) M2 NPD using signals recorded from BG output nuclei (either internal globus pallidus (GPi /EPN in rat) and/or SNr) or their major targets in the thalamus. If these signals were available then it would be possible to determine if there is enhanced cortical return of BG beta rhythms in the dopamine-depleted state.

Hypothesis 2: Pathological beta is generated from strengthening of the reciprocally coupled STN/GPe circuit.

A separate hypothesis concerning the generation of pathological beta rhythms in the basal ganglia considers the reciprocally coupled STN/GPe circuit from which increased coupling associated with the loss of dopamine induces a pathological beta resonance that spreads across the rest of the network (Plenz and Kitai 1999; Bevan et al. 2002; Holgado et al. 2010; Tachibana et al. 2011).

We note that conditioning the NPD with the M2 signal does not remove the strong STN \(\rightarrow\) GPe directed connectivity, but it does attenuate the GPe \(\rightarrow\) STN (figure 9). This would suggest that activity feeding back onto GPe from STN has sufficiently unique information to not be partialized out by the cortical ECoG, perhaps generated by some resonance phenomena. However, a number of the analyses presented here suggest that pathological beta does not originate from STN/GPe circuit resonance. This is summarised as follows: i) Comparison of forward and backward NPD for STN/GPe interactions shows strong asymmetry, with the GPe \(\rightarrow\) STN connection predominating; ii) conditioning of the NPD using the LFPs recorded at the STR significantly reduces the strength of both GPe \(\rightarrow\) STN and STN \(\rightarrow\) GPe NPDs in both the control and lesion animals (figure 8), suggesting that beta in these structures results from beta propagating through striatum; iii) conditioning the NPD with LFPs recorded at the STN (figure 6) does not act to remove the upstream 6-OHDA associated beta NPD between STR or GPe (although it does significantly weaken beta NPD in the control animals); iv) GPe conditioned NPD analysis does not show impairment of the pathological M2/STN beta interactions (figure 7) suggesting that the beta found at STN can be, at least in part, generated independently of a signal found at GPe. The evidence summarised in point (i), may speak to the hypothesis that the STN/GPe circuit acts as a pacemaker that is in turn tuned by abnormal striatal drive, but by some nonlinear transformation that is not discernible using the conditioned NPD. Taken together we argue these findings give evidence against pathological beta rhythms arising from the pacemaker-like activity of the STN/GPe sub-circuit.

Hypothesis 3: Beta arises through aberrant striatal activity and facilitation of downstream hypersynchrony.

It has been proposed that aberrant cortico-striatal activity is involved in the emergence of pathological beta rhythms (Kumar et al. 2011; McCarthy et al. 2011; Damodaran et al. 2015). We provide evidence that elements of the BG downstream of STR require a signal that is also shared with the cerebral cortex (M2). In the lesioned animals, there is significant beta band NPD in the M2 \(\rightarrow\) STR and STR
GPe connections (figure 5) suggesting a propagation of beta activity down the indirect pathway. With conditioning, we find that the beta peak in the STR $\rightarrow$ GPe connection is removed in its entirety when conditioning the NPD with the cortical signal (figure 9) in both the control and lesion recordings (interestingly however, the STR $\rightarrow$ STN beta NPD does remain) suggesting that beta band correlations are at least in part explained by cortical activity. Importantly, we found that conditioning the STR $\rightarrow$ STN connection with the LFP recorded from GPe (figure 7) acts to significantly reduce the NPD in both control and lesion experiments. This would suggest that beta flow is mediated via GPe in agreement with the canonical view of the indirect pathway demonstrating the ability of partial NPD to discern hierarchical connectivity. Further to this, the finding that upstream GPe/STR beta NPD is susceptible to conditioning with LFPs from STN (figure 6) in dopamine-intact animals may reinforce the idea that in the healthy network, routing of beta occurs predominantly through the indirect pathway, whilst in the pathological case we see beta that is persistent and with activity that is at least in part independent of that recorded at the STN.

Hypotheses of the Origins/Routing of High Beta/Gamma Oscillations

The presence of high beta/gamma oscillations in the subcortical network has been noted by a number of authors (Brown et al. 2002; Berke 2009; Sharott et al. 2009; van der Meer et al. 2010; Nicolás et al. 2011) but our understanding of the functional propagation of high beta/gamma oscillations through the network is limited. An evaluation of the evidence is summarised in table 2. We report gamma activity in the LFPs as well as connectivity in the range 30-60 Hz which is in good agreement with that previously reported in anaesthetised rats (Magill et al. 2004; Sharott et al. 2005b, 2009). Activity in the awake rats, during movement has also been reported, albeit at slightly higher frequencies (Brown et al. 2002; Brazhnik et al. 2012; Delaville et al. 2014).

Hypothesis 4: High beta/gamma enters the subcortical network via the hyper-direct M2 $\rightarrow$ STN connection.

Results from analyses which used iCOH to investigate non-zero lag correlations between BG structures and the cortex suggested that gamma interactions are routed in a way that bypasses STR as a gamma peak is absent in the M2 (figure 4). The hyper-direct pathway is the other principal source of cortical input to the BG, therefore the marked weakness of gamma interaction in the M2/STR when compared to the M2/STN iCOH spectra may imply that the hyper-direct pathway is responsible for gamma input to the network.

However, whilst there is a large peak in the high-beta/low-gamma band NPD for the M2 $\rightarrow$ STN interaction (figure 5), if we examine the same connection but conditioned on LFPs either recorded at STR (figure 7) or GPe (figure 8) we see that the conditioning significantly reduces NPD in the control animals (M2 $\rightarrow$ STN conditioned on STR 16-37 Hz, P<0.001; M2 $\rightarrow$ STN conditioned on GPe 16-47 Hz, P<0.001) suggesting any directed coherence between M2 and STN in these animals is routed...
via striatal-pallidal connections. Furthermore, if we condition the NPD with LFPs recorded at the STN (figure 6), we see that gamma interactions remain in the upstream components (M2 → STR, M2 → GPe) again suggesting striatal-pallidal connectivity is vital in the propagation of gamma rhythms. When taken together, these data do not supply strong evidence that the source of high beta/gamma input in the network is supplied by a hyper-direct cortico-subthalamic route.

Hypothesis 5: Gamma enters the network via cortico-striatal inputs and reaches STN via the indirect pathway in a dopamine dependent manner.

An alternative to high beta/gamma oscillations entering via hyper-direct STN input is that they are channelled via the cortico-striatal indirect pathway. The clearest results of the NPD analysis in the high beta/gamma band can be seen to be for the forward NPDs originating from M2 and passing on to the subcortical regions (figure 5). Connections M2 → STR, M2 → GPe, and M2 → STN all show high values of NPD in this frequency band (> 0.15) suggesting that most of the gamma is directed from the cortex. Furthermore, conditioning the NPD with either LFPs recorded at the STR (figure 8) or GPe (figure 7) acts to remove gamma interactions both upstream and downstream of the STR (with respect to the indirect pathway). Subsequently, conditioning of the NPD with STN (figure 6) was less effective at attenuating gamma band interactions than when using signals higher in the indirect pathway, suggesting that the gamma descends the hierarchy, from either a cortical or striatal source. Notably, we observed that STN conditioned NPD did not act to attenuate feedback connections from GPe or STR back to the M2. This would suggest routing of gamma to the M2 in a way that occurs independently of STN.

In attempt to elucidate the source of the gamma activity we conditioned the NPD on the cortical ECoG (figure 9). We find that gamma connectivity in the control recordings, dopamine depletions acts to significantly reduce NPD coefficients for the GPe → STN and STR → STN connections, yet the feedback connection STN → STR is unaffected. This connection in the control animals shows a peak from 18-42 Hz which is significantly larger than in the lesioned animals. This is in agreement with the idea of prokinetic gamma and in accordance with findings in patients (Sharott et al. 2014). Furthermore, these findings suggest that gamma activity is directed to upstream components of the BG in a way independent of M2, perhaps via a subcortical feedback loop.

Hypothesis 6: High beta/gamma is generated in the GPe or STR and downstream to STN

The finding that conditioning the NPD with cortical ECoG does not entirely abolish gamma connectivity within the BG may suggest that at least some high beta/gamma propagates locally within the BG. When conditioning was performed with LFPs recorded from the GPe (figure 7) or STR (figure 8), we found that interactions in the high beta/gamma frequency ranges were abolished for the majority of subcortical interactions. This would imply that these GPe and STR structures are necessary for the propagation of high beta/gamma interactions in the both the control and 6-OHDA...
lesion animals. This in combination with the evidence provided for hypothesis 5 suggests that high
beta/gamma originates at either STR or GPe and then propagates to downstream structures. Backward
gamma interactions from GPe to STR are apparent in the NPD conditioned on M2 or STN, suggesting
the STR signal is the result of local propagation of a gamma signal from GPe. From the canonical
view of the circuit it is not clear how gamma passes upstream from GPe. However, a substantial
proportion of GPe neurons that innervate the striatum have been shown to exist, with one GPe cell
type (arkypallidal neurons) projecting exclusively to striatum (Mallet et al., 2012; Abdi et al., 2015;
Hegeman et al. 2016).

Summary of Findings

In this paper, we have presented analyses that investigate propagation of oscillatory activity through
connected regions of the cortico-basal ganglia network. We have applied a novel method of
partialized directed coherence to achieve a systematic deconstruction of the propagation of rhythmic
activity between regions of the network inferred from the LFPs and ECoGs recorded at multiple sites
within that network. Using the 6-OHDA-lesioned rat model of Parkinsonism, we demonstrate marked
differences in the functional connectivity that result as a consequence of dopamine depletion.

We find diffuse beta synchronization of LFPs across the network that is strongly associated with
chronic dopamine depletion. With regards to functional beta connectivity in the network we find
evidence for:

1. An increased cortical drive to the basal ganglia following dopamine depletion.
2. Significant beta-band connectivity between structures interacting with the STN that is
   independent of activity upstream in the indirect pathway (at STR and GPe). This likely
   originates from the ‘hyperdirect’ cortico-subthalamic input.
3. Increase in feedback of BG structures to M2 after dopamine depletion, proffering evidence in
   favour of a hypothesis of dopamine-dependent modulation of the long re-entrant cortico-BG-
   thalamo-cortical loop.
4. Activity dynamics of the STN/GPe subcircuit being partly dependent upon drive from
   striatum.
5. A feedback from STN to STR that is independent of M2 and significantly strengthened after
dopamine depletion, suggesting a strengthening of recurrent subcortical circuits.

Furthermore, we provide evidence for the existence of high beta/gamma synchrony within the
network, with evidence that dopamine depletion acts to weaken these rhythms. We summarise our
findings with respect to high beta/gamma band interactions in the following:

1. Gamma propagates down the indirect pathway from STR to GPe to STN. This activity is
   likely generated at cortex.
2. At least some of the gamma activity found at STN is independent of M2 and provides evidence for a subcortical return of subthalamic outputs back to striatum.

3. At least some of the gamma activity returning to cortex is independent of STN, perhaps indicating propagation through the direct pathway.

Propagation of Low Beta via two Coexisting but Distinct Streams

In the case of low beta oscillations, we find our data most strongly support a hypothesis that in the dopamine-depleted condition, beta propagation in the network is biased to favour low beta synchrony via induction of a long cortico-subthalamic loop that inputs to the BG via the hyper-direct pathway. Furthermore, we see strong evidence that the return connection from STN to M2 is significantly stronger in the lesioned animals than in the dopamine-intact controls. This provides supporting evidence for the notion that pathological beta amplification arises from entrainment of the re-entrant cortical/STN loop (Brittain and Brown 2014). We speculate that strengthening of the hyper-direct input acts to “short-circuit” the network, such that transmission of information along the indirect pathway is compromised. In the “hold your horses” model of the STN’s role in decision making (Frank 2006; Frank et al. 2007), the hyper-direct pathway is proposed to provide a cortical veto signal which may act to suppress premature action. In the case of PD, over activity of this circuit via increased resonance may act to lock the network into a state that ultimately supresses action. These findings are in agreement with previous research which have found good evidence for bidirectional connectivity between STN and cortex (Lalo et al. 2008; Jávor-Duray et al. 2015).

This hypothesis requires further testing through analysis of the role of the BG output nuclei at GPi or SNr (or their targets in the thalamus) in the propagation of activity. This could be achieved using a functional ‘lesion’ approach like that described in this paper. Furthermore, biophysical modelling of the cortico-subthalamic loop may yield insight as to whether this is plausible given the known conduction delays for the connections in the network. Long feedback loops involving cortex have been demonstrated to be capable of generating oscillatory activity (Leblois et al. 2006; Pavlides et al. 2015). However, work by Shouno et al. (2017) suggests that the required delay for the return of the beta oscillation from STN to cortex may be too large to support resonance in the low beta band. Perhaps the engagement of shorter subcortical loops either subcortical-thalamic loops (McHaffie et al. 2005); or activity of recurrent subthalamo-striatal projections (Sato et al. 2000; Koshimizu et al. 2013) may be more suitable candidates.

The analysis presented here also suggests that a cortico-subthalamic pathway is not the exclusive pathway for beta rhythms within the network, yet may be necessary for enhancement of the STN feedback to cortex that may induce pathological resonance. We would suggest that both the hyperdirect and indirect routes for beta propagation coexist. These two pathways could originate form (be driven by) distinct populations of cortical projections neurons (namely those of the pyramidal tract...
and intra-telencephalic projections, respectively) and so are likely to show a degree of independence from one another. The data presented here also suggest a second pathway upstream of STN involving the STR that is most evident in the recordings from control rats. We suggest that both pathways contain signals shared by activity measured in the cortical ECoG as conditioning of the NPD acts to remove beta peaks from the majority of connections that were analysed, leaving just beta coherence at the STR → STN connection. These findings support the hypothesis that dopamine cell loss acts to increase the sensitivity of the STR to cortical inflow acting to gate beta to the remainder of the circuit (Magill et al. 2001; Tseng et al. 2001).

Notably, our data do not support the hypothesis of beta generation via an autonomous STN/GPe pacemaker network, as directional coherence between the two is heavily attenuated by conditioning with LFPs recorded upstream in the STR and there is significant asymmetry in the NPD with the pallidal drive predominating. In agreement, Moran et al. (2011) found evidence for a weakening of the STN to GPe feedback connection in the dopamine depleted state, conflicting with the STN/GPe resonance hypothesis. Estimates of effective connectivity from DCM studies have also suggested that input from cortex to STN is strengthened in the Parkinsonian state (Moran et al. 2011), a finding in line with the idea that dopamine enforces cortical influence upon the STN/GPe network (Magill et al. 2001; Leblois 2006; Leblois et al. 2006; Holgado et al. 2010). It may be the case that during PD cortical activity subsumes the STR as the primary driver of the STN/GPe subcircuit. It has been demonstrated that movement is associated with a decreased cortico-pallidal coherence during movement in humans (van Wijk et al. 2017) suggesting disengagement of this circuit is pro-kinetic. Thus pathological resonance may arise as a result of alterations to striatal output (Kumar et al. 2011; Damodaran et al. 2015) that in the healthy system act to decorrelate spiking activity between the two structures (Wilson 2013). The action of dopamine upon these inputs is likely to modulate the existence of beta amplifying phase alignments between STN and GPe such as that observed by Cagnan, Duff, & Brown (2015).

Dopamine Depletion is Associated with an Increased Subthalamo-Striatal Feedback

Taken together, the analyses presented here speak to the existence of a high beta/low gamma rhythm that is in general reduced by dopamine depletion. Specifically, our results indicate a band at 27-34 Hz is attenuated in the lesioned rats. Experiments investigating LFPs in the motor cortex of moving rats have demonstrated an increase in activity in this band during movement suggesting that activity at these frequencies in the M2 and SNr is prokinetic (Brazhnik et al. 2012). Our data would suggest that high beta/gamma activity in the normal network is predominantly driven by the cortex as evidenced by: i) the unconditioned NPD indicates that gamma is prominently in the forward direction leading from cortex to subcortical sources; and ii) conditioning the NPD on ECoG recorded at M2 acts to diminish the subcortical directional coherence across a wide band for all connections not involving
STN. However, studies by Zold and colleagues has demonstrated that oscillatory activity >20 Hz in corticostriatal afferents is not effectively transmitted (Zold et al. 2012).

Furthermore, following partialization some interactions involving STN do remain. In particular we provide evidence for a significant strengthening of feedback from STN to STR in the lesioned animals in the high beta/gamma band. We speculate that this signal is facilitated through the strengthening of subcortical loops such as that of the thalamo-striatal pathways (McHaffie et al. 2005). Thalamic afferents make up to at least 25% of input onto spiny projection neurons in the STR (Doig et al. 2010; Smith et al. 2014) but have been far less studied than cortical inputs. Work investigating synaptic remodelling following 6-OHDA depletion in mice has suggested that thalamo-striatal inputs to medium spiny neurons are shifted in favour of the indirect pathway (Parker et al. 2016) perhaps enhancing striatal return of subthalamic activity in a mechanism independent of cortex.

Study Limitations
Incomplete Signals for Conditioning
The use of partial coherence for inferring neural connectivity is not in itself a novel approach (Rosenberg et al. 1998; Eichler et al. 2003; Salvador et al. 2005; Medkour et al. 2009), and the application of the conditioned NPD to LFPs recorded in the rat hippocampus has been previously reported (Halliday et al. 2016). However these analyses assume that the signals used for conditioning completely capture the activity going through the proposed pathway. This however is unlikely to be completely the case due to the finite sampling of the structures afforded from the use of electrodes. However, the large number of channels for recording ensure that multiple samples are obtained from each structure. In the data presented here, subcortical structures were recorded from between 2-8 different channels which were all used to conditioning the estimate of directed coherence. It must also be noted that this limitation is likely to apply most to the larger structures that were analysed, namely the motor cortex and striatum, whereas recordings from the smaller sized STN is more likely to capture the majority of activity. This must be considered when interpreting conditioning of the NPD with respect to STR signals. It could be the case that M1 → STN connectivity remains as a result of incomplete sampling of striatal signals.

Inference of Connectivity from Non-Spiking Brain Activity
This study focuses upon conclusions that are drawn from mesoscale recordings of brain activity as measured either in the ECoG or LFPs. Transmission of information in the brain is facilitated by axonal propagation of action potentials that are not explicitly captured in these signals. LFPs and ECoG comprise a conglomerate of sub- and supra- threshold events that may or may not be tied to spike activity and so direct inference of neurophysiological connectivity per se is limited by this. Nonetheless, it has been previously demonstrated by Mallet and colleagues that beta activity in the LFPs recorded at STN and GPe are associated with increased synchronization of action potential
firing by neurons in these structures (Mallet et al. 2008a, 2008b). Furthermore, we provide evidence for the existence of temporally lagged correlations between rhythmic local field potentials recorded between distinct regions of the cortico-BG network that imply causation from one signal to another, a phenomenon that would itself not be possible without the transmission of action potentials. Future work will require the investigation to determine whether directional interactions are ascertainable from multiunit activity and how this relates to lagged synchronization of LFPs.

Limits to Inference of Directed Connectivity from Neurophysiological Signals Alone

In this paper, we aim to infer how neural activity propagates across the BG network by investigating the statistical relationships between brain signals. The challenges that this approach face are well documented (Friston 2011; Bastos and Schoffelen 2016). With respect to this study, the benefits that we put forth for using a model free, non-parametric approach (namely agnosticism to the underlying generating mechanisms of the data) may in turn limit the degree of inference that can be made. Estimates of directed functional connectivity in this paper follow from the assumptions that temporal precedence is indicative of causation. It is however well documented that zero lag synchronization can emerge from neural circuits with particular (but not unusual) network motifs (Vicente et al. 2008; Viriyopase et al. 2012; Gollo et al. 2014). Additionally, “anticipatory” synchronization in which positive lags arise from a directed input have also been described in theoretical neural dynamics (Ambika and Amritkar 2009; Ghosh and Roy Chowdhury 2010). The anatomically tightly coupled STN-GPe subcircuit is a prime candidate for which these phenomena may permit vanishingly small phase lags that make the interactions blind to NPD. Answers to these problems may be given in the future by the fitting of biophysical models to the data presented in this paper. This would provide a well-defined, quantitative description of the potential mechanisms that act to generate the phenomena we have described.

Conclusion

Overall, we provide a systematic deconstruction of the propagation of pathological rhythms across the Parkinsonian cortico-basal ganglia circuit in vivo. These findings strengthen our understanding of how normal and pathological rhythms propagate across the network. Our work highlights the importance of considering non-canonical connections in the network, in particular the activity of recurrent subcortical projections that may act to amplify pathological activity within the BG. Future work will aim to understand the exact changes to the network required to generate the patterns of functional connectivity presented here, as well as to investigate the relationship with spiking activity in the network.

Acknowledgments

We thank Dr N. Mallet for acquiring some primary data sets.
**Funding**

Medical Research Council UK (awards UU138197109, MC_UU_12020/5 and MC_UU_12024/2 to P.J.M.; MC_UU_21024/1 to A.S.). Parkinson’s UK (Grant G-0806 to P.J.M.). S.F.F. receives funding from UCLH BRC. Engineering Research Council UK (awards EPSRC EP/F500351/1 to T.O.W.; EP/N007050/1 to D.H.). The Wellcome Trust Centre for Neuroimaging is funded by core funding from the Wellcome Trust (539208).

**References**

van Albada SJ, Robinson PA. Mean-field modeling of the basal ganglia-thalamocortical system. I: Firing rates in healthy and parkinsonian states. *J Theor Biol* 257: 642–663, 2009.

Ambika G, Amritkar RE. Anticipatory synchronization with variable time delay and reset. *Phys Rev E* 79: 56206, 2009.

Bastos AM, Schoffelen J-M. A Tutorial Review of Functional Connectivity Analysis Methods and Their Interpretational Pitfalls. *Front Syst Neurosci* 9: 175, 2016.

Berke JD. Fast oscillations in cortical-striatal networks switch frequency following rewarding events and stimulant drugs. *Eur J Neurosci* 30: 848–859, 2009.

Beudel M, Oswal A, Jha A, Foltynie T, Zrinzo L, Hariz M, Limousin P, Litvak V. Oscillatory Beta Power Correlates With Akinesia-Rigidity in the Parkinsonian Subthalamic Nucleus. *Mov Disord* 32: 174–175, 2017.

Bevan MD, Magill PJ, Terman D, Bolam JP, Wilson CJ. Move to the rhythm: oscillations in the subthalamic nucleus–external globus pallidus network. *Trends Neurosci* 25: 525–531, 2002.

Bolam JP, Hanley JJ, Booth PA, Bevan MD. Synaptic organisation of the basal ganglia. *J Anat* 200: 527–42, 2000.

Brazhnik E, Cruz A V., Avila I, Wahba MI, Novikov N, Ilieva NM, McCoy AJ, Gerber C, Walters JR. State-Dependent Spike and Local Field Synchronization between Motor Cortex and Substantia Nigra in Hemiparkinsonian Rats. *J Neurosci* 32, 2012.

Bressler SL, Menon V. Large-scale brain networks in cognition: emerging methods and principles. *Trends Cogn Sci* 14: 277–290, 2010.

Brittain J-S, Brown P. Oscillations and the basal ganglia: motor control and beyond. *Neuroimage* 85: 637–647, 2014.

Brown P. Abnormal oscillatory synchronisation in the motor system leads to impaired movement. *Curr Opin Neurobiol* 17: 656–664, 2007.
Brown P, Kupsch A, Magill PJ, Sharott A, Harnack D, Meissner W. Oscillatory Local Field Potentials Recorded from the Subthalamic Nucleus of the Alert Rat. 2002.

Brown P, Oliviero A, Mazzone P, Insola A, Tonali P, Di Lazzaro V. Dopamine Dependency of Oscillations between Subthalamic Nucleus and Pallidum in Parkinson’s Disease [Online]. J Neurosci 21: 1033–1038, 2001. http://www.jneurosci.org/content/21/3/1033.abstract [4 Jan. 2016].

Cagnan H, Duff EP, Brown P. The relative phases of basal ganglia activities dynamically shape effective connectivity in Parkinson’s disease. Brain 138: 1667–1678, 2015.

Damodaran S, Cressman JR, Jedrzejewski-Szmek Z, Blackwell KT. Desynchronization of Fast-Spiking Interneurons Reduces β-Band Oscillations and Imbalance in Firing in the Dopamine-Depleted Striatum J Neurosci 35, 2015.

Deco G, Jirsa VK, Robinson PA, Breakspear M, Friston K. The dynamic brain: from spiking neurons to neural masses and cortical fields. PLoS Comput Biol 4: e1000092, 2008.

Deco G, Senden M, Jirsa V. How anatomy shapes dynamics: a semi-analytical study of the brain at rest by a simple spin model. Front Comput Neurosci 6: 68, 2012.

Delaville C, Cruz A V, McCoy AJ, Brazhnik E, Avila I, Novikov N, Walters JR. Oscillatory Activity in Basal Ganglia and Motor Cortex in an Awake Behaving Rodent Model of Parkinson’s Disease. Basal Ganglia 3: 221–227, 2014.

DeLong M, Wichmann T. Changing views of basal ganglia circuits and circuit disorders. Clin EEG Neurosci 41: 61–7, 2010.

Doig NM, Moss J, Bolam JP. Cortical and Thalamic Innervation of Direct and Indirect Pathway Medium-Sized Spiny Neurons in Mouse Striatum. J Neurosci 30, 2010.

Dostrovsky J, Bergman H. Oscillatory activity in the basal ganglia--relationship to normal physiology and pathophysiology. Brain 127: 721–722, 2004.

Dovzhenok A, Rubchinsky LL. On the Origin of Tremor in Parkinson’s Disease. PLoS One 7, 2012.

Eichler M, Dahlhaus R, Sandkuhler J. Partial correlation analysis for the identification of synaptic connections. Biol Cybern 89: 289–302, 2003.

Eusebio a, Thevathasan W, Doyle Gaynor L, Pogosyan a, Bye E, Foltynie T, Zrinzo L, Ashkan K, Aziz T, Brown P. Deep brain stimulation can suppress pathological synchronisation in parkinsonian patients. J Neurol Neurosurg Psychiatry 82: 569–573, 2011.

Frank MJ. Hold your horses: A dynamic computational role for the subthalamic nucleus in decision making. Neural Networks 19: 1120–1136, 2006.
Frank MJ, Samanta J, Moustafa AA, Sherman SJ. Hold your horses: impulsivity, deep brain stimulation, and medication in parkinsonism. *Science* 318: 1309–12, 2007.

Fries P. A mechanism for cognitive dynamics: neuronal communication through neuronal coherence. *Trends Cogn Sci* 9: 474–480, 2005.

Fries P. Rhythms for Cognition: Communication through Coherence. *Neuron* 88: 220–235, 2015.

Friston KJ. Functional and Effective Connectivity: A Review. *Brain Connect* 1: 13–36, 2011.

Ghosh D, Roy Chowdhury A. Lag and anticipatory synchronization based parameter estimation scheme in modulated time-delayed systems. *Nonlinear Anal Real World Appl* 11: 3059–3065, 2010.

Gollo LL, Mirasso C, Sporns O, Breakspear M. Mechanisms of Zero-Lag Synchronization in Cortical Motifs. *PLoS Comput Biol* 10: e1003548, 2014.

Halliday DM, Senik MH, Stevenson CW, Mason R. Non-parametric directionality analysis – Extension for removal of a single common predictor and application to time series. *J Neurosci Methods* 268: 87–97, 2016.

Hammond C, Bergman H, Brown P. Pathological synchronization in Parkinson’s disease: networks, models and treatments. *Trends Neurosci* 30: 357–364, 2007.

Hanslmayr S, Staudigl T, Fellner M-C. Oscillatory power decreases and long-term memory: the information via desynchronization hypothesis. *Front Hum Neurosci* 6: 74, 2012.

Holgado AJN, Terry JR, Bogacz R. Conditions for the Generation of Beta Oscillations in the Subthalamic Nucleus–Globus Pallidus Network. *J Neurosci* 30, 2010.

Jávor-Duray BN, Vinck M, van der Roest M, Mulder AB, Stam CJ, Berendse HW, Voorn P. Early-onset cortico-cortical synchronization in the hemiparkinsonian rat model. *J Neurophysiol* 113, 2015.

Koshimizu Y, Fujiyama F, Nakamura KC, Furuta T, Kaneko T. Quantitative analysis of axon bouton distribution of subthalamic nucleus neurons in the rat by single neuron visualization with a viral vector. *J Comp Neurol* 521: 2125–2146, 2013.

Kühn AA, Kupsch A, Schneider G-H, Brown P. Reduction in subthalamic 8-35 Hz oscillatory activity correlates with clinical improvement in Parkinson’s disease. *Eur J Neurosci* 23: 1956–60,
Kumar A, Cardanobile S, Rotter S, Aertsen A. The role of inhibition in generating and controlling Parkinson’s disease oscillations in the Basal Ganglia. *Front Syst Neurosci* 5: 86, 2011.

Lalo E, Thobois S, Sharott A, Polo G, Mertens P, Pogosyan A, Brown P. Patterns of Bidirectional Communication between Cortex and Basal Ganglia during Movement in Patients with Parkinson Disease. *J Neurosci* 28, 2008.

Lanciego JL, Luquin N, Obeso JA. Functional neuroanatomy of the basal ganglia. *Cold Spring Harb Perspect Med* 2: a009621, 2012.

Leblois A. Competition between Feedback Loops Underlies Normal and Pathological Dynamics in the Basal Ganglia. *J Neurosci* 26: 3567–3583, 2006.

Leblois A, Boraud T, Meissner W, Bergman H, Hansel D. Competition between Feedback Loops Underlies Normal and Pathological Dynamics in the Basal Ganglia. *J Neurosci* 26: 3567–3583, 2006.

Levy R. Dependence of subthalamic nucleus oscillations on movement and dopamine in Parkinson’s disease. *Brain* 125: 1196–1209, 2002.

Levy R, Hutchison WD, Lozano AM, Dostrovsky JO. High-frequency Synchronization of Neuronal Activity in the Subthalamic Nucleus of Parkinsonian Patients with Limb Tremor. *J Neurosci* 20: 7766–7775, 2000.

Litvak V, Jha A, Eusebio A, Oostenveld R, Foltynie T, Limousin P, Zrinzo L, Hariz MI, Friston K, Brown P. Resting oscillatory cortico-subthalamic connectivity in patients with Parkinson’s disease. *Brain* 134: 359–374, 2011.

Magill P, Bolam J, Bevan M. Dopamine regulates the impact of the cerebral cortex on the subthalamic nucleus–globus pallidus network. *Neuroscience* 106: 313–330, 2001.

Magill PJ, Pogosyan A, Sharott A, Csicsvari J, Bolam JP, Brown P. Changes in functional connectivity within the rat striatopallidal axis during global brain activation in vivo. *J Neurosci* 26: 6318–6329, 2006.

Magill PJ, Sharott A, Bolam JP, Brown P. Brain State–Dependency of Coherent Oscillatory Activity in the Cerebral Cortex and Basal Ganglia of the Rat. *J Neurophysiol* 92, 2004.

Mallet N, Pogosyan A, Marton LF, Bolam JP, Brown P, Magill PJ. Parkinsonian Beta Oscillations in the External Globus Pallidus and Their Relationship with Subthalamic Nucleus Activity. *J Neurosci* 28: 14245–14258, 2008a.

Mallet N, Pogosyan A, Sharott A, Csicsvari J, Bolam JP, Brown P, Magill PJ. Disrupted
Dopamine Transmission and the Emergence of Exaggerated Beta Oscillations in Subthalamic Nucleus and Cerebral Cortex. *J Neurosci* 28: 4795–4806, 2008b.

Maris E. Statistical testing in electrophysiological studies. *Psychophysiology* 49: 549–565, 2012.

Maris E, Oostenveld R. Nonparametric statistical testing of EEG- and MEG-data. *J Neurosci Methods* 164: 177–190, 2007.

Marreiros AC, Cagnan H, Moran RJ, Friston KJ, Brown P. Basal ganglia–cortical interactions in Parkinsonian patients. *Neuroimage* 66: 301–310, 2013.

McCarthy MM, Moore-Kochlacs C, Gu X, Boyden ES, Han X, Kopell N. Striatal origin of the pathologic beta oscillations in Parkinson’s disease. *Proc Natl Acad Sci U S A* 108: 11620–5, 2011.

McHaffie JG, Stanford TR, Stein BE, Coizet V, Redgrave P. Subcortical loops through the basal ganglia. *Trends Neurosci* 28: 401–407, 2005.

Medkour T, Walden AT, Burgess A. Graphical modelling for brain connectivity via partial coherence. *J Neurosci Methods* 180: 374–383, 2009.

van der Meer M, Kalenscher T, Lansink CS, Pennartz C, Berke JD, Redish AD. Integrating early results on ventral striatal gamma oscillations in the rat. *Front Neurosci* 4: 300, 2010.

Moran RJ, Mallet N, Litvak V, Dolan RJ, Magill PJ, Friston KJ, Brown P. Alterations in brain connectivity underlying beta oscillations in parkinsonism. *PLoS Comput Biol* 7: e1002124, 2011.

Nambu A, Tokuno H, Takada M. Functional significance of the cortico–subthalamo–pallidal “hyperdirect” pathway. *Neurosci Res* 43: 111–117, 2002.

Nevado-Holgado AJ, Mallet N, Magill PJ, Bogacz R. Effective connectivity of the subthalamic nucleus - globus pallidus network during Parkinsonian oscillations. *J Physiol* 7: 1429–1455, 2014.

Ni Z, Bouali-Bennazouz R, Gao D, Benabid A-L, Bennazouz A. Changes in the firing pattern of globus pallidus neurons after the degeneration of nigrostriatal pathway are mediated by the subthalamic nucleus in the rat. *Eur J Neurosci* 12: 4338–4344, 2000.

Nicolás MJ, Lópe-Azcárate J, Valencia M, Alegre M, Pérez-Alcázar M, Iriarte J, Artieda J. Ketamine-Induced Oscillations in the Motor Circuit of the Rat Basal Ganglia. *PLoS One* 6: e21814, 2011.

Nolte G, Bai O, Wheaton L, Mari Z, Vorbach S, Hallett M. Identifying true brain interaction from EEG data using the imaginary part of coherency. *Clin Neurophysiol* 115: 2292–307, 2004.
Electrophysiological Data. *Comput Intell Neurosci* 2011: 1–9, 2011.

Parker PRL, Lalive AL, Kreitzer AC. Pathway-Specific Remodeling of Thalamostriatal Synapses in Parkinsonian Mice. *Neuron* 89: 734–740, 2016.

Pavides A, Hogan SJ, Bogacz R. Computational Models Describing Possible Mechanisms for Generation of Excessive Beta Oscillations in Parkinson’s Disease. *PLOS Comput Biol* 11: e1004609, 2015.

Pavides A, John Hogan S, Bogacz R. Improved conditions for the generation of beta oscillations in the subthalamic nucleus-globus pallidus network. *Eur J Neurosci* 36: 2229–2239, 2012.

Paxinos G, Watson C. *The rat brain in stereotaxic coordinates*. Elsevier, 2007.

Plenz D, Kital ST. A basal ganglia pacemaker formed by the subthalamic nucleus and external globus pallidus. *Nature* 400: 677–82, 1999.

Ray NJ, Jenkinson N, Wang S, Holland P, Brittain JS, Joint C, Stein JF, Aziz T. Local field potential beta activity in the subthalamic nucleus of patients with Parkinson’s disease is associated with improvements in bradykinesia after dopamine and deep brain stimulation. *Exp Neurol* 213: 108–113, 2008.

Rosenberg JR, Halliday DM, Breeze P, Conway BA. Identification of patterns of neuronal connectivity—partial spectra, partial coherence, and neuronal interactions. *J Neurosci Methods* 83: 57–72, 1998.

Salvador R, Suckling J, Schwarzbauer C, Bullmore E. Undirected graphs of frequency-dependent functional connectivity in whole brain networks. *Philos Trans R Soc Lond B Biol Sci* 360: 937–46, 2005.

Sato F, Parent M, Levesque M, Parent A. Axonal branching pattern of neurons of the subthalamic nucleus in primates. *J Comp Neurol* 424: 142–152, 2000.

Schoffelen J-M, Oostenveld R, Fries P. Neuronal coherence as a mechanism of effective corticospinal interaction. *Science* 308: 111–113, 2005.

Schnitzler A, Gross J. Normal and pathological oscillatory communication in the brain. *Nat Rev Neurosci* 6: 285–96, 2005.

Schroll H, Hamker FH. Computational models of basal-ganglia pathway functions: focus on functional neuroanatomy. *Front Syst Neurosci* 7: 1–18, 2013.

Schroll H, Vitay J, Hamker FH. Dysfunctional and compensatory synaptic plasticity in Parkinson’s disease. *Eur J Neurosci* 39: 688–702, 2014.
Schwarting RK, Huston JP. Unilateral 6-hydroxydopamine lesions of meso-striatal dopamine neurons and their physiological sequelae. Prog Neurobiol 49: 215–66, 1996a.

Schwarting RK, Huston JP. The unilateral 6-hydroxydopamine lesion model in behavioral brain research. Analysis of functional deficits, recovery and treatments. Prog Neurobiol 50: 275–331, 1996b.

Sharott A, Gulberti A, Zittel S, Tudor Jones AA, Fickel U, Münchau A, Köppen JA, Gerloff C, Westphal M, Buhmann C, Hamel W, Engel AK, Moll CKE. Activity Parameters of Subthalamic Nucleus Neurons Selectively Predict Motor Symptom Severity in Parkinson’s Disease. J Neurosci 34, 2014.

Sharott A, Magill PJ, Bolam JP, Brown P. Directional analysis of coherent oscillatory field potentials in the cerebral cortex and basal ganglia of the rat. J Physiol 562: 951–63, 2005a.

Sharott A, Magill PJ, Harnack D, Kupsch A, Meissner W, Brown P. Dopamine depletion increases the power and coherence of 7-?-oscillations in the cerebral cortex and subthalamic nucleus of the awake rat. Eur J Neurosci 21: 1413–1422, 2005b.

Sharott A, Moll CKE, Engler G, Denker M, Grün S, Engel AK. Different Subtypes of Striatal Neurons Are Selectively Modulated by Cortical Oscillations. J Neurosci 29, 2009.

Shen W, Flajolet M, Greengard P, Surmeier DJ. Dichotomous Dopaminergic Control of Striatal Synaptic Plasticity. Science (80- ) 321, 2008.

Shouno O, Tachibana Y, Nambu A, Doya K. Computational Model of Recurrent Subthalamo-Pallidal Circuit for Generation of Parkinsonian Oscillations. Front Neuroanat 11: 21, 2017.

Smith Y, Bevan MD, Shink E, Bolam JP. Microcircuitry of the direct and indirect pathways of the basal ganglia. Neuroscience 86: 353–387, 1998.

Smith Y, Galvan A, Ellender TJ, Doig N, Villalba RM, Huerta-Ocampo I, Wichmann T, Bolam JP. The thalamostratial system in normal and diseased states. Front Syst Neurosci 8: 5, 2014.

Stam CJ, Nolte G, Daffertshofer A. Phase lag index: Assessment of functional connectivity from multi channel EEG and MEG with diminished bias from common sources. Hum Brain Mapp 28: 1178–1193, 2007.

Steriade M. Corticothalamic resonance, states of vigilance and mentation. Neuroscience 101: 243–276, 2000.

Tachibana Y, Iwamuro H, Kita H, Takada M, Nambu A. Subthalamo-pallidal interactions underlying parkinsonian neuronal oscillations in the primate basal ganglia. Eur J Neurosci 34: 1470–1484, 2011.
Terman D, Rubin JE, Yew AC, Wilson CJ. Activity patterns in a model for the subthalamopallidal network of the basal ganglia. J Neurosci 22: 2963–2976, 2002.

Thut G, Miniussi C, Gross J. The functional importance of rhythmic activity in the brain. Curr Biol 22: R658-63, 2012.

Tseng KY, Kasanetz F, Kargieman L, Riquelme LA, Murer MG. Cortical Slow Oscillatory Activity Is Reflected in the Membrane Potential and Spike Trains of Striatal Neurons in Rats with Chronic Nigrostriatal Lesions. J Neurosci 21, 2001.

Uhlhaas PJ, Singer W. Neural Synchrony in Brain Disorders: Relevance for Cognitive Dysfunctions and Pathophysiology. Neuron 52: 155–168, 2006.

Varela F, Lachaux J-PP, Rodriguez E, Martinerie J. The brainweb: phase synchronization and large-scale integration. Nat Rev Neurosci 2: 229–39, 2001.

Vicente R, Gollo LL, Mirasso CR, Fischer I, Pipa G. Dynamical relaying can yield zero time lag neuronal synchrony despite long conduction delays. Proc Natl Acad Sci U S A 105: 17157–62, 2008.

Vinck M, Oostenveld R, van Wingerden M, Battaglia F, Pennartz CMA. An improved index of phase-synchronization for electrophysiological data in the presence of volume-conduction, noise and sample-size bias. Neuroimage 55: 1548–1565, 2011.

Viriyopase A, Bojak I, Zeitler M, Gielen S. When Long-Range Zero-Lag Synchronization is Feasible in Cortical Networks. Front Comput Neurosci 6: 49, 2012.

Weinberger M, Mahant N, Hutchison WD, Lozano AM, Moro E, Hodaie M, Lang AE, Dostrovsky JO. Beta oscillatory activity in the subthalamic nucleus and its relation to dopaminergic response in Parkinson’s disease. J Neurophysiol 96: 3248–56, 2006.

West T, Farmer S, Berthouze L, Jha A, Beudel M, Foltynie T, Limousin P, Zrinzo L, Brown P, Litvak V. The Parkinsonian Basal Ganglia Network: Measures of Power, Linear and Non-Linear Synchronization and their Relationship to L-DOPA Treatment and OFF State Motor Severity. Front Hum Neurosci 10: 517, 2016.

Whittmer D, de Solages C, Hill B, Yu H, Henderson JM, Bronte-Stewart H. High frequency deep brain stimulation attenuates subthalamic and cortical rhythms in Parkinson’s disease. Front Hum Neurosci 6: 155, 2012.

Wichmann T, DeLong MR. Neurobiology: Oscillations in the basal ganglia. Nature 400: 621–622, 1999.

van Wijk BCM, Neumann W-J, Schneider G-H, Sander TH, Litvak V, Kühn AA. Low-beta cortico-pallidal coherence decreases during movement and correlates with overall reaction time.
Figure 1 – Schematic of canonical cortical-basal ganglia circuit incorporating the direct and indirect pathways, as well as the cortico-subthalamic hyperdirect pathway. The motor cortex (M2, yellow box) has major inputs to the basal ganglia (BG, orange boxes) at the striatum (STR) and subthalamic nucleus (STN). Information flow along the indirect, direct and hyperdirect pathways ultimately impinges on the output nuclei of the basal ganglia, made up of the entopeduncular nucleus (EPN) and substantia nigra pars reticulata (SNr). BG output targets thalamic relays, of which some return back to M1. Structures for which experimental data is available are shown in bold outline, otherwise in dashed.

Figure 2 - Example outcome of data pre-processing of rat subcortical monopolar LFP and cortical ECoG signals for a single animal from the control (A-B) or Parkinsonian (C-D) groups. (A) 5 second excerpt of recording from one dopamine intact, control animal. The example trace shows the time course of LFP recordings recorded using silicon electrodes implanted in the external globus pallidus (GPe), striatum (STR) and subthalamic nucleus (STN), and with multiple contacts within each structure. Additionally, there is a cortical ECoG signal recorded from a screw over motor cortex (M2). Raw data is shown in blue whilst pre-processed signals are overlaid in red. The data was demeaned and then high pass filtered at 4 Hz. (B) Spectral analysis of example control animal’s recording. Data epoched into 2-second segments contaminated by high amplitude transients were removed using Z-thresholding as described in the text. (C) Same as (A) but for an example 6-OHDA-lesioned (dopamine depleted) animal. (D) Same as (B) but for 6-OHDA-lesioned animal.

Figure 3 – Group averaged power spectra for all rats across both control and lesion conditions. Spectra are shown for signals recorded from (A) motor cortex (M2), (B) the striatum (STR), (C) the subthalamic nucleus (STN), and (D) the external globus pallidus (GPe). Individual spectra are shown in dashed lines whilst the group averages for either the 6-OHDA dopamine depleted or control animals are shown by bold lines in red or blue respectively. Results of cluster permutation tests are indicated by the bar and corresponding P-value. All recording sites presented beta peaks around 18-20 Hz. Cluster based permutation testing for significant differences between conditions showed that there was a significant increase in beta in the lesioned animals for signals recorded at STR, STN, and GPe.
Figure 4 – Functional connectivity estimates using imaginary part of coherence (iCOH). Spectra for each animal are shown by thin lines corresponding to either 6-OHDA lesioned (blue) or control (red). The group averages are shown by the bold lines. Cluster-based permutation statistics were applied to determine significant differences between conditions and significant clusters are indicated by the bold line above the spectra and corresponding P-value. The iCOH metric, robust to zero-lag interactions, presents a rich view of functional connectivity that would otherwise be missed if using standard coherence (data not shown). Beta activity is predominant across all cross-regional pairings. STN and GPe also show some intra-nuclear correlations in this range in the dopamine depleted state. Notably there is also a high beta/gamma interaction between STN/M2 and STN/STR that is visible in both control and lesion animals.

Figure 5 - Directed connectivity estimated using non-parametric directionality (NPD) between subcortically recorded LFPs (GPe, STN, and STR) and ECoG recorded from the motor cortex (M2). NPD decomposes the coherence between pairs of signals into forward and reverse components. The array of spectra in the figures reads such that each row title gives the structure with a forward coherence targeted to the structure given by the name given above the column. Spectra from individual animals are shown by the dashed lines and the group means in bold. Different spectra for the control and 6-OHDA lesioned animals are indicated by the red and blue lines respectively.

Figure 6 - Non-parametric directionality conditioned on LFPs recorded from the STN - Spectra for each animal are shown by thin lines corresponding to either 6-OHDA lesioned (blue) or control (red). The group averages are shown by the bold lines. Cluster-based permutation statistics were applied to determine significant differences between control and lesioned conditions, with significant clusters indicated by the black bold line above the spectra and corresponding P-value. Significant changes between the original NPD and the conditioned variant shown here are indicated by the coloured bold line corresponding to a difference either with the control (red) or lesioned (blue) signals.

Figure 7 - Non-parametric directionality conditioned on LFPs recorded from the GPe - Spectra for each animal are shown by thin lines corresponding to either 6-OHDA lesioned (blue) or control (red). The group averages are shown by the bold lines. Cluster-based permutation statistics were applied to determine significant differences between control and lesioned conditions, with significant clusters indicated by the black bold line above the spectra and corresponding P-value. Significant changes between the original NPD and the conditioned variant shown here are indicated by the coloured bold line corresponding to a difference either with the control (red) or lesioned (blue) signals.

Figure 8 - Non-parametric directionality conditioned on LFPs recorded from the STR - Spectra for each animal are shown by thin lines corresponding to either 6-OHDA lesioned (blue) or control
The group averages are shown by the bold lines. Cluster-based permutation statistics were applied to determine significant differences between control and lesioned conditions, with significant clusters indicated by the black bold line above the spectra and corresponding P-value. Significant changes between the original NPD and the conditioned variant shown here are indicated by the coloured bold line corresponding to a difference either with the control (red) or lesioned (blue) signals.

Figure 9 - Non-parametric directionality conditioned on the M2 ECoG - Spectra for each animal are shown by thin lines corresponding to either 6-OHDA lesioned (blue) or control (red). The group averages are shown by the bold lines. Cluster-based permutation statistics were applied to determine significant differences between control and lesioned conditions, with significant clusters indicated by the black bold line above the spectra and corresponding P-value. Significant changes between the original NPD and the conditioned variant shown here are indicated by the coloured bold line corresponding to a difference either with the control (red) or lesioned (blue) signals.

Table 1– Summary of hypotheses of the impact of dopamine depletion on the propagation of beta rhythms in the cortico-basal ganglia circuit.

Table 2 – Summary of hypotheses for gamma flow in the cortico-basal ganglia circuit.
Imaginary Coherence

Motor Cortex

Striatum

Subthalamic Nucleus

External Globus Pallidus
Cortex

Striatum

Subthalamic Nuclei

GPe

conditioned
Non-Parametric
Directionality
The diagram illustrates the spectral properties of neural connections between different brain regions, specifically focusing on the effects of lesions on NPD (Normalized Power Density) across various frequency bands for different pathways:

- **Cortex:** Connections from the cortex to subthalamic nuclei and globus pallidus.
- **Subthalamic Nuclei:** Connections from the subthalamic nuclei to M2, STN, and GPe.
- **Globus Pallidus:** Connections from the globus pallidus to M2 and STN.

Each graph shows the frequency distribution of neural activity, with different colors indicating control and lesion conditions. Significant differences (p-values) are marked on the graphs, highlighting the impact of lesions on neural directionality and non-parametric statistics.

**STR conditioned Non-Parametric Directionality**
| Mechanism: | Beta via connections involved in the “long loop” resonance | Beta generated via GPe/STN reciprocal pair | Beta via the cortico-striatal indirect pathway |
|-----------|-------------------------------------------------|-------------------------------------|--------------------------------------|
| **Pathological beta oscillations arise from inductions of a long cortico-BG-thalamic loop.** | **Pathological beta oscillations arise from resonance of reciprocally coupled GPe/STN subcircuit.** | **Pathological beta oscillations arise from changes to striatal “filtering” of cortical inputs.** |
| **Evidence for:** | • Significant beta band STN/M2 NPD in both the forward and reverse directions.  
• The low beta STN → M2 feedback coherence is significantly increased in the lesioned animals.  
• STN/M2 NPD is undiminished by conditioning with GPe or STR. | • Partialing NPD with the M2 signal does not remove beta interactions subcortically.  
• Cluster analysis on the iCOH STN/GPe connectivity shows a lesion related increase at low beta frequencies. | • Beta NPD between STR, GPe and M2 remains when partialising for STN.  
• Conditioning the NPD with signals from the STN or M2 does not remove upstream beta coherences. |
| **Evidence against:** | • Conditioning the NPD with signals from the STN or M2 does not remove beta band NPD between STR and GPe, upstream of the STN. | • Partialing NPD with the STN does not remove upstream STR/GPe beta interactions.  
• NPD conditioned on the GPe does not impair M2/STN coherences.  
• Partialling STR weakens the the STN/GPe NPD interaction (13-48 Hz, lesioned, P=0.001). | • Conditioning the NPD on M2 does not attenuate indirect pathway connections downstream of the STR and GPe. |
| Hypothesis:                                      | Cortical gamma inflow via hyper-direct pathway | Cortical-striatal indirect gamma inflow | Gamma source at GPe projected out to remainder of circuit |
|-------------------------------------------------|-----------------------------------------------|----------------------------------------|--------------------------------------------------------|
| Evidence for:                                   | • iCOH gamma present in M2/STN but not M2/STR suggests STR is bypassed.  
• Conditioning of NPD upon STR does not remove M2/STN gamma in the lesioned animals. | • Majority of gamma NPD is forwardly leading (from cortex and descending connections).  
• Partialising with STR or GPe weakens downstream gamma connectivity.  
• Partialising with STN does not weaken upstream gamma interactions. | • Partialising the NPD estimate with GPe/STR removes gamma interactions  
• Increased feedback in the STN-STR connection in the 6-OHDA state (conditioning for M2) |
| Evidence against:                               | • Partialising NPD with STN signal does not remove gamma in the upstream (M2 → STR,M2 → GPe) components.  
• Conditioning of NPD upon STR does removes M2 → STN gamma in control recordings (14-36Hz, control, P = 0.001) | • Partialising NPD with signal from M2 does not remove STR/GPe/STN coherences | • Not clear how connections pass from GPe back to M2.  
• The M2 → GPe gamma NPD shows significant decrease in response to dopamine loss (24-33Hz, P = 0.015). |