Longitudinal functional connectivity changes related to dopaminergic decline in Parkinson’s disease

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\textbf{ABSTRACT}

\textbf{Background:} Resting-state functional magnetic resonance imaging (fMRI) studies have demonstrated that basal ganglia functional connectivity is altered in Parkinson’s disease (PD) as compared to healthy controls. However, such functional connectivity alterations have not been related to the dopaminergic deficits that occurs in PD over time.

\textbf{Objectives:} To examine whether functional connectivity impairments are correlated with dopaminergic deficits across basal ganglia subdivisions in patients with PD both cross-sectionally and longitudinally.

\textbf{Methods:} We assessed resting-state functional connectivity of basal ganglia subdivisions and dopamine transporter density using \textsuperscript{11}C-PE2I PET in thirty-four PD patients at baseline. Of these, twenty PD patients were rescanned after 19.9 ± 3.8 months. A seed-based approach was used to analyze resting-state fMRI data. \textsuperscript{11}C-PE2I binding potential (BPND) was calculated for each participant. PD patients were assessed for disease severity.

\textbf{Results:} At baseline, PD patients with greater dopaminergic deficits, as measured with \textsuperscript{11}C-PE2I PET, showed larger decreases in posterior putamen functional connectivity with the midbrain and pallidum. Reduced functional connectivity of the posterior putamen with the thalamus, midbrain, supplementary motor area and sensorimotor cortex over time were significantly associated with changes in DAT density over the same period. Furthermore, increased motor disability was associated with lower intraregional functional connectivity of the posterior putamen.

\textbf{Conclusions:} Our findings suggest that basal ganglia functional connectivity is related to integrity of dopaminergic system in patients with PD. Application of resting-state fMRI in a large cohort and longitudinal scanning may be a powerful tool for assessing underlying PD pathology and its progression.

1. Introduction

Parkinson’s disease (PD) is characterised by progressive loss of dopaminergic neurons in the substantia nigra and subsequent dysfunction of dopamine neurotransmission in the striatum (Nurmi et al., 2001; Eshuis et al., 2006). The presynaptic dopamine transporter (DAT) is a plasma membrane transporter that is localised exclusively in dopamine neurones (Piccini, 2003). Hence, DAT availability measured with DAT-specific radioligands and Positron Emission Tomography (PET) is one measure of nigrostriatal dopaminergic integrity (Shih et al., 2006; Pavese and Brooks, 2009). Previous longitudinal studies have demonstrated an annualized rate of reduction in striatal DAT binding of approximately 6–13% in patients with PD compared with 0–2.5% in healthy controls (Nurmi et al., 2000; Marek et al., 2001; Parkinson Study, 2002). The reduction in striatal DAT binding correlates with increasing disability in patients with PD (Seibyl et al., 1995; Li et al., ...
Resting-state functional magnetic resonance imaging (fMRI) offers a means of assessing the status of functional networks within the brain without the confounding influence of task performance (Fox and Greicius, 2010). It has been demonstrated that functional changes within the basal ganglia network can differentiate patients with PD from healthy controls (Szewczyk-Krolkowskiet al., 2014), and are responsible for the cardinal motor features of PD (Wu et al., 2012). Hacker et al. (2012) reported lower striatal functional connectivity within the thalamus, midbrain, pons and cerebellum in patients with PD. Another study using a more spatially refined analysis found reduced functional connectivity between the posterior putamen and the sensorimotor cortex while functional connectivity was increased between the anterior putamen and the inferior parietal cortex (Helmich et al., 2010). It has been shown that functional connectivity in PD improves across striatal subdivisions under regular dopaminergic replacement therapy (Bell et al., 2015), suggesting that striatal functional connectivity is dependent on the integrity of dopaminergic system (Surmeier et al., 2010). For the substantia nigra, it was found that functional connectivity is decreased to the putamen in PD patients compared to healthy controls (Ellmore et al., 2013).

There are few studies assessing longitudinal changes in resting-state functional connectivity in PD patients (Olde Dubbelink et al., 2014; Manza et al., 2016). Olde Dubbelink et al. (2014) reported a gradual loss of global resting-state functional connectivity in PD, mainly in sensorimotor cortex, over the course of three years, which was modestly related to motor deficits (Olde Dubbelink et al., 2014). However, it is unclear how changes in resting-state functional connectivity are related to dopaminergic decline in patients with PD over time.

The primary aim of this study was to examine whether functional connectivity impairments are correlated with dopaminergic deficits across basal ganglia subdivisions in patients with PD both cross-sectionally and longitudinally. Here, we used resting-state fMRI to assess functional connectivity in thirty-four PD patients and fifteen age-matched healthy controls. To estimate DAT binding we used 11C-PE2I PET, a highly specific radioligand which has been shown to be closely associated with PD motor severity and progression (Li et al., 2017). We hypothesized that longitudinal changes in resting-state functional connectivity of basal ganglia subdivisions would be associated with reductions in DAT binding and increasing motor disability in PD patients.

2. Method

2.1. Participants

Patients with PD were recruited from the Transeuro programme cohort (FP7 EC http://www.transeuro.org.uk/). All had a diagnosis of idiopathic PD in accordance with Queen Square Brain Bank criteria (Lees et al., 2009). A total of thirty-four patients with PD (PD BL) completed 11C-PE2I PET, structural MRI and resting-state fMRI scans at baseline, however four were excluded from the current analyses due to significant head motion. Of these, twenty PD patients completed follow-up measures after an interval of 19.9 ± 3.8 months, hereby referred to as the PD follow-up subgroup (PD FU). From this group, three patients were excluded due to severe signal loss, and two patients were excluded due to excessive head motion during resting-state fMRI. Therefore, a total of fifteen patients were included in the longitudinal analysis.

Motor severity was assessed by two experienced clinicians using the motor sub-score of the Unified Parkinson’s Disease Rating Scale (UPDRS-part III) (Goetz et al., 2008) and the Hoehn and Yahr staging scale (Hoehn and Yahr, 1967) whilst they were in an “off” medication state. We further subdivided the UPDRS-part III into tremor and bradykinesia-rigidity sub-scores. A total of fifteen age- and gender-matched healthy controls completed structural MRI and resting-state fMRI scans at baseline. None of the participants scored < 26 on the Mini-Mental State Examination (MMSE), had atypical or secondary Parkinsonism, or were ineligible for MRI and PET scanning (claustrophobia, metallic implants, pregnancy or breastfeeding, recent exposure to ionising radiation). All participants gave written informed consent in accordance with the Declaration of Helsinki. All aspects of the study were approved from the appropriate research ethics committees of the UK (REC 12/EE/0096 & 10/40805/73) and Sweden (EPN 2013/758 & IK 2013/685) National Research Ethics Service Committee.

2.2. Image acquisition

All scans were conducted at the Invicro Imaging Centre (Hammersmith Hospital Campus, London). During scans, participants were positioned supine; head movement was minimised using memory foam padding and video monitoring utilised to aid detection and subsequent repositioning.

MRI scans were performed on a 3T Siemens Magnetom Trio with 32-channel phased-array head coil. High resolution structural images were acquired using a 3-dimensional T1-weighted sagittal magnetization-prepared rapid gradient-echo sequence (MPRAGE: TR/TE = 2300/2.98 ms; TI = 900 ms; FA = 9°; bandwidth = 240 Hz/Px; GRAPPA acceleration factor = 2; slice thickness = 1 mm (no gap); FoV = 240 × 256 mm; matrix size = 240 × 256) in which one whole brain volume was obtained consisting of 160 slices lasting 301 s. Resting-state fMRI scans were acquired for 368 s using a T2*-weighted single-shot gradient-echo echo planar imaging (EPI) sequence (TR/TE = 2500/31.3 ms; FA = 80°; bandwidth = 2298 Hz/Px; GRAPPA acceleration factor = 2; slice thickness = 3 mm (no gap); FoV = 192 × 192 mm; matrix size = 64 × 64). 144 brain volumes were obtained per individual, each consisting of 45 interleaved axial slices acquired parallel to the AC-PC line. Participants were instructed to remain as still as possible with their eyes closed, and to avoid falling asleep during the scan.

11C-PE2I (N-(3-iodopro-2E-enyl)-2β-carbomethoxy-3β-(4’-methylpheny1)nortropane) PET imaging was conducted on a Siemens Biograph TruePoint HI-REZ 6 PET/CT system. Radioligand volumes (11C-PE2I = 350 MBq) were prepared to 10 ml using saline solution and administered intravenously as single bolus injections followed immediately by 10 ml saline flush. Administration was at 1 ml/sec. Dynamic emission data were acquired continuously for 90 min post-injection then reconstructed into 26 temporal frames using a filtered back-projection algorithm (direct inversion Fourier transform; matrix size = 128x128, zoom = 2.6, 5 mm transaxial Gaussian filter, pixel size = 2 mm isotropic). A low-dose CT transmission scan (0.36 mSv) was acquired for attenuation correction prior to the injection of 11C-PE2I. PD patients were instructed to withdraw their dopaminergic medication at least 24 h prior to scanning for standard release preparations and 48 h for prolonged release medications (Li et al., 2017).

2.3. Image preprocessing

Standard image preprocessing of resting-state fMRI was performed using Statistical Parametric Mapping (SPM12, Wellcome Trust Centre for Neuroimaging, London, UK, http://www.fil.ion.ucl.ac.uk/spm/). Each participant’s structural image was segmented for cerebrospinal fluid (CSF) and white matter (WM). Resting-state fMRI data were slice-time corrected, realigned to the first image using rigid-body motion transforms and co-registered to the structural image. For every time point t, the framewise displacement (FD) was calculated, and every time point that exceeded a pre-defined head motion limit (FD (t) > 0.5 mm) was removed (Power et al., 2012). The FD values were later used as nuisance regressors and confounding covariates in the individual and group-level analyses, respectively. Then, a ‘scrubbing’ procedure was performed using the ‘fsl_motion_outliers’ tool in FSL to remove the time points affected by head motion (motion spike) (Power et al., 2012). A nuisance regression was conducted using a Volterra...
2.4. Functional connectivity analysis

Resting-state functional connectivity was assessed using a seed-based approach with bilateral caudate, anterior putamen, posterior putamen and substantia nigra as seeds. Functional connectivity maps were obtained by computing the Pearson’s correlation coefficient between the averaged time series of each seed region and that of all other voxels in the brain for each individual participant. These correlation coefficients were converted to normally distributed values using Fisher’s r-to-Z transform for use at the group level. The individual z-maps were then entered into a random effect one-sample t-test to identify brain regions showing significant positive correlations with the seed ROIs. Then, the individual z-maps were entered into a random effect two-sample t-test to identify differences in functional connectivity between groups (PD BL vs. healthy controls at baseline; PD BL vs. PD FU). Statistical maps were thresholded at voxel-level $p_{FWE} < 0.05$ and cluster-level $p_{FDR} < 0.05$.

2.5. Multiple regression analysis

For patients with PD at baseline (n = 30), to investigate the associations between regional $^{11}$C-PE2I BPND and functional connectivity of the seed regions, voxel-based multiple regression analysis was conducted using functional connectivity maps as the dependent variable, the regional $^{11}$C-PE2I BPND as the independent variable, and age/ gender as nuisance covariates. The FD, averaged across the entire time series, was also entered as a covariate to minimize the potential effect of motion-related variance on group inference. The correlation analysis was assessed using r-statistics. To explore correlates of functional connectivity with disease severity, multiple regression analysis was performed with UPDRS-III, bradykinesia-rigidity and tremor sub-scores entered as covariates of interest. All statistical maps were thresholded at voxel-level $p < 0.001$ uncorrected and cluster-level false discovery rate corrected $p_{FDR} < 0.05$.

We also assessed how changes in functional connectivity over time correlated with changes in $^{11}$C-PE2I BPND over time. Difference functional connectivity maps were created by voxel-based subtraction ($\Delta$: follow-up – baseline) and entered into a multiple regression analysis as the dependent variable with $^{11}$C-PE2I BPND as the independent variable, and age/ gender as nuisance covariates. In addition, to examine how changes in functional connectivity correlated with changes in motor disability over time, multiple regression analysis was also performed with $\Delta$UPDRS-III, $\Delta$bradykinesia-rigidity and $\Delta$tremor sub-scores entered as covariates of interest. All statistical maps...
were thresholded at voxel-level \( p < 0.001 \) uncorrected and cluster-level false discovery rate corrected \( \text{pFDR} < 0.05 \).

### 3. Results

Demographics and clinical characteristics of the study participants are summarised in Table 1.

#### 3.1. Resting-state functional connectivity in patients with PD and healthy controls

Resting-state functional connectivity of basal ganglia subdivisions at baseline in patients with PD was generally consistent with previous studies (Fig. 2). The caudate was connected to the anterior/posterior cingulate cortex and thalamus. The anterior putamen and posterior putamen exhibited similar patterns of functional connectivity with the supplementary motor area (SMA), sensorimotor cortex, thalamus and a contiguous region of grey matter extending from the thalamus through the midbrain, pons and cerebellum. The substantia nigra showed functional connectivity with the caudate, putamen, globus pallidus, thalamus and cerebellum. The functional connectivity patterns of the healthy controls were roughly similar to that of the PD patients (Supplementary Material Fig. 1). To ensure that group-level results were not influenced by spurious correlations as a result of in-scanner head movement, we compared the mean FD between the groups (healthy controls vs. PD at baseline, \( p = 0.577 \); PD at baseline vs. PD at follow up, \( p = 0.493 \)), suggesting that head movement did not differentially alter resting-state functional connectivity across groups (Supplementary Material Table 2).

At baseline, voxel-based comparisons of functional connectivity maps for each of the four seed regions between PD patients and healthy controls were conducted. Both the caudate and the anterior putamen showed decreased intraregional functional connectivity for the PD group as compared to HC (Fig. 3a and 3b), while the posterior putamen showed the opposite pattern. In the PD group, the posterior putamen exhibited an enhanced intraregional functional connectivity intermediate between anterior and posterior putamen (Fig. 3c). PD patients exhibited reduced functional connectivity between the substantia nigra and the thalamus and pallidum as well as with smaller clusters located within the caudate and putamen (Fig. 3d).

Functional connectivity maps of each of the four seed regions were also compared between baseline and follow-up for the PD patients. We found a time-related decrease of functional connectivity between the anterior putamen and sensorimotor cortex (Fig. 4a). In addition, the posterior putamen showed decreased functional connectivity with the anterior putamen and the posterior putamen (intraregional), as well as smaller clusters in the caudate, thalamus and midbrain (Fig. 4b). There were no significant differences in caudate and substantia nigra functional connectivity between the two time points.

#### 3.2. Correlation between basal ganglia functional connectivity and \(^{11}\text{C}-\text{PE2I BPND}\) and clinical measures: multiple regression analyses

At baseline, voxel-based multiple regression analysis of posterior putamen functional connectivity maps revealed clusters in the midbrain and pallidum showing a significant positive correlation with posterior putamenal \(^{11}\text{C}-\text{PE2I BPND}\) (Fig. 5a, Table 2). No significant correlations between functional connectivity maps of other seed regions (caudate, anterior putamen and substantia nigra) and \(^{11}\text{C}-\text{PE2I BPND}\) values were observed. Furthermore, we did not find any significant correlations between the functional connectivity of basal ganglia subdivisions and clinical scores.

Longitudinally, voxel-based analysis of the change in posterior putamen functional connectivity (\( \Delta \): follow-up – baseline) revealed clusters showing significant positive correlations with \( \Delta^{11}\text{C}-\text{PE2I BPND} \) in the posterior putamen. These clusters were found in the left thalamus and a contiguous region of grey matter extending from the thalamus, through the midbrain, as well as in the SMA, precentral gyrus and postcentral gyrus (Fig. 5b, Table 2). Furthermore, clusters showing significant positive correlations between \( \Delta \)posterior putamen functional connectivity and \( \Delta \text{bradykinesia-rigidity} \) sub-scores were found in the right posterior putamen (Fig. 5c, Table 2). No other significant results

### Table 1

Demographic and clinical information for all study participants at baseline (PD: \( n = 30 \); HC: \( n = 15 \)) and for the PD subgroup at baseline and 19.9 ± 3.8 months follow-up (PD\textsubscript{FU}: \( n = 15 \)). Independent-samples t-test was used to assess the difference between PD and healthy controls at baseline. For the PD follow-up group, Wilcoxon signed-rank and paired-samples t-test were used to assess the difference between baseline and follow-up.

<table>
<thead>
<tr>
<th>Demographics and clinical characteristics</th>
<th>PD ( n = 30 )</th>
<th>HC ( n = 15 )</th>
<th>Statistic</th>
<th>( p )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender (M:F)</td>
<td>24:6</td>
<td>8:7</td>
<td>( t(19.674) = 0.696 )</td>
<td>0.494</td>
</tr>
<tr>
<td>Age (years) *</td>
<td>55.4 ± 7.1</td>
<td>53.15 ± 11.4</td>
<td>( t(14) = 26.445 )</td>
<td>&lt; 0.001 *</td>
</tr>
<tr>
<td>Disease duration (years) *</td>
<td>5.8 ± 2.2</td>
<td>N/A</td>
<td>( t(14) = 26.234 )</td>
<td>&lt; 0.001 *</td>
</tr>
<tr>
<td>UPDRS-III *</td>
<td>31.3 ± 10.6</td>
<td>N/A</td>
<td>( t(14) = 2.003 )</td>
<td>0.065</td>
</tr>
<tr>
<td>UPDRS-III Bradykinesia-Rigidity *</td>
<td>22.4 ± 8.1</td>
<td>N/A</td>
<td>( t(14) = 1.925 )</td>
<td>0.075</td>
</tr>
<tr>
<td>UPDRS-III Tremor *</td>
<td>6.2 ± 5.3</td>
<td>N/A</td>
<td>( t(14) = 1.514 )</td>
<td>0.152</td>
</tr>
<tr>
<td>Hoehn &amp; Yahr Scale *</td>
<td>2.0 (0.0)</td>
<td>N/A</td>
<td>( Z = 1.7 )</td>
<td>0.083</td>
</tr>
<tr>
<td>LED *</td>
<td>678.3 ± 361.8</td>
<td>N/A</td>
<td>( t(14) = 2.156 )</td>
<td>0.046 *</td>
</tr>
<tr>
<td>Follow-Up: ( \text{PD}_\text{FU} ) ( n = 15 )</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gender (M:F)</td>
<td>12:3</td>
<td>12:3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age (years) *</td>
<td>53.0 ± 7.2</td>
<td>54.7 ± 7.1</td>
<td>( t(14) = 26.445 )</td>
<td>&lt; 0.001 *</td>
</tr>
<tr>
<td>Disease duration (years) *</td>
<td>6.0 ± 2.2</td>
<td>7.7 ± 2.0</td>
<td>( t(14) = 26.234 )</td>
<td>&lt; 0.001 *</td>
</tr>
<tr>
<td>UPDRS-III *</td>
<td>31.4 ± 12.0</td>
<td>35.9 ± 10.9</td>
<td>( t(14) = 2.003 )</td>
<td>0.065</td>
</tr>
<tr>
<td>UPDRS-III Bradykinesia-Rigidity *</td>
<td>22.4 ± 9.6</td>
<td>25.2 ± 8.9</td>
<td>( t(14) = 1.925 )</td>
<td>0.075</td>
</tr>
<tr>
<td>UPDRS-III Tremor *</td>
<td>6.9 ± 4.5</td>
<td>8.3 ± 4.2</td>
<td>( t(14) = 1.514 )</td>
<td>0.152</td>
</tr>
<tr>
<td>Hoehn &amp; Yahr Scale *</td>
<td>2.0 (0.0)</td>
<td>2.0 (0.0)</td>
<td>( Z = 1.7 )</td>
<td>0.083</td>
</tr>
<tr>
<td>LED *</td>
<td>771.7 ± 386.5</td>
<td>906.3 ± 235.7</td>
<td>( t(14) = 2.156 )</td>
<td>0.046 *</td>
</tr>
</tbody>
</table>

\* Data are presented as Mean ± SD.

\( \Delta \) indicates \( p < 0.05 \).

LED = L-dopa equivalent dose (mg).

UPDRS = Unified Parkinson’s disease rating scale.

The UPDRS and Hoehn scale were assessed in the practically defined off-medication state.
were found for longitudinal analyses.

4. Discussion

Our findings demonstrate that resting-state functional connectivity of the posterior putamen is associated with the integrity of the dopaminergic system in patients with PD. PD patients with more severe dopaminergic deficits showed larger decreases in posterior putamen functional connectivity with the midbrain and pallidum. In addition, longitudinal comparison between the two timepoints revealed that patients with PD showed reduced functional connectivity of the posterior putamen with the thalamus and midbrain. These functional connectivity changes were correlated with dopamine deficits of disease progression. Lastly, increased motor disability was associated with lower intraregional functional connectivity of the posterior putamen. These results suggest an essential role of dopamine in striatal function.
and demonstrate that resting-state functional connectivity of basal ganglia subdivisions is compromised by the dopaminergic pathology of PD.

A previous study using resting-state fMRI demonstrated reduced functional connectivity within the basal ganglia of patients with early PD as compared to patients with Alzheimer’s disease (AD) and healthy controls (Rolinski et al., 2015). These changes were not seen in AD patients, suggesting that aberrant functional connectivity within the
basal ganglia of PD patients is disease specific. In the current study, reduced intrarregional functional connectivity of the caudate and the anterior putamen was found in PD patients compared to healthy controls. In contrast, we found increased intraregional functional connectivity of the posterior putamen, suggesting a compensatory mechanism that develops in response to marked dopamine decline in the posterior putamen (Helmich et al., 2010). It has been demonstrated that noradrenaline, serotonin and acetylcholine are also affected in PD, although to a lesser degree than dopamine, thus it is possible that other neurotransmitter systems might play a role in mediating changes in basal ganglia functional connectivity patterns (Shine et al., 2019). Lastly, we found reduced substantia nigra functional connectivity with the thalamus extending to the caudate and putamen, which is generally consistent with results reported by Wu et al. (2012) who also showed additional reductions to the globus pallidus, insula and SMA in PD patients as compared to the healthy controls (Wu et al., 2012).

Previous studies have demonstrated significant correlations between motor symptomatology and altered functional connectivity of basal ganglia subdivisions in patients with PD (Wu et al., 2009; Hacker et al., 2012; Agosta et al., 2014). Greater severity of UPDRS-III motor scores appears to be associated with reduced functional connectivity between the striatum and midbrain (Hacker et al., 2012) and increased functional connectivity between the putamen and occipital/parietal cortex (Agosta et al., 2014), suggesting a potential role basal ganglia functional connectivity alterations in the occurrence of motor impairment (Cerasa et al., 2016). However, in the current study, baseline functional connectivity did not correlate with any clinical measures of motor severity. In line with our findings, a study examined functional connectivity within the basal ganglia network also did not find any significant correlations between the basal ganglia network connectivity and clinical measures (Szewczyk-Krolkowski et al., 2014). They suggested that functional connectivity alterations of basal ganglia network may be a trait biomarker of PD, reflecting a constitutional fault of the network and not the resulting motor symptoms (Szewczyk-Krolkowski et al., 2014).

Postmortem and neuroimaging studies have demonstrated that the posterior putamen suffers most from degeneration of the nigrostriatal dopaminergic pathway (Kish et al., 1988; Nurmi et al., 2001; Bruck et al., 2006; Helmich et al., 2010). The most consistent finding across resting-state fMRI studies in patients with PD appears to be a reduction of posterior putaminal functional connectivity that correlates with greater motor impairment (Herz et al., 2014). Few studies have examined how striatal functional connectivity varies with dopaminergic deficit in PD. One study found that reduced functional connectivity between the putamen and the midbrain was related to dopaminergic status in the striatum (Rieckmann et al., 2015) while another demonstrated that functional connectivity was modulated by striatal dopamine levels (Baik et al., 2014). Here, in thirty PD patients, we found that reduced functional connectivity between the posterior putamen and the midbrain was significantly associated with dopamine levels in the posterior putamen. In a preliminary analysis, we also found in a subset of fifteen PD patients that changes in this functional connectivity pattern correlated with changes in dopamine levels in the posterior putamen. Taken together, these findings suggest that resting-state functional connectivity of the posterior putamen may reflect impairment of the nigrostriatal dopaminergic projection in the PD brain.

Although PD is the most common form of parkinsonism, there are other atypical neurodegenerative conditions that mimic the clinical features of PD, such as multiple system atrophy (MSA) and progressive supranuclear palsy (PSP) (Politiis, 2014). DAT radioligands with PET have shown promising sensitivity and specificity for distinguishing patients with parkinsonian syndromes from healthy controls (Antonini et al., 2001; Jakobson Mo et al., 2018). However, dopamine-specific radioligands cannot accurately differentiate between PD, MSA and PSP. Advanced MRI techniques, such as resting-state fMRI and arterial spin labeling (ASL), may therefore help to provide complementary information for more accurate differential diagnosis. Recently developed hybrid PET/MR systems allow simultaneous acquisition of MRI and PET data, which in theory enables more precise spatial co-registration and importantly offers greater convenience for motor impaired patients (Tondo et al., 2019). Indeed, combined acquisition of ASL and 18F-FDG PET on a hybrid PET/MR system was recently shown to increase sensitivity in differentiating PD and MSA, based on changes in the caudate nucleus, pons and cerebellum (Ruan et al., 2019). Another study demonstrated a close relationship between global grey matter changes and the degeneration of striatal dopaminergic circuits in patients with PD (Choi et al., 2016). Taken together, as a non-invasive and more comprehensive approach, the combined use of advanced MRI and PET on hybrid scanners may play an increasingly important role for evaluating the underlying mechanisms of patients with PD.

This study has some limitations that need to be addressed. First, although our longitudinal results accord with well-founded predictions they were based on a relatively small number of PD participants and as such should be validated in larger samples. We believe that our findings provide promising new insights into changes in resting-state functional connectivity and their relation to DAT decline and motor dysfunction in PD patients. Second, only a single time interval of ~20 months was studied, thus future resting-state fMRI studies should be designed to evaluate patients more than twice and at longer intervals, which may provide additional evidence on the staging of PD progression and the
Results of the voxel-based multiple regression analysis for PD patients at baseline (n = 30; a) and follow-up (n = 15; b and c). (a) The posterior putamen functional connectivity revealed clusters in the midbrain and pallidum, that showed a significant positive correlation with $^{11}$C-PE2I BPND in the posterior putamen. (b) Changes in posterior putamen $^{11}$C-PE2I-BPND are positively correlated with changes in functional connectivity between the posterior putamen and thalamus, midbrain, SMA, precentral gyrus and postcentral gyrus. (c) Changes in bradykinesia-rigidity sub-score are positively correlated with changes in posterior putamen intraregional functional connectivity. All statistical maps ($p < 0.001$ uncorrected at voxel-level, and $p_{	ext{FDR}} < 0.05$ FDR-corrected at cluster-level) are overlaid on a T1-weighted MNI template. L. left; R, right.

### Table 2

Clusters showing positive correlations between posterior putamen functional connectivity and $^{11}$C-PE2I BPND and disease severity (Bradykinesia-Rigidity sub-score), as revealed by voxel-wise multiple regression analysis for all PD patients. Longitudinal multiple regression using difference maps ($\Delta$: follow-up – baseline) from the PD$_{FU}$ group are also shown.

<table>
<thead>
<tr>
<th>Seed</th>
<th>Covariate of Interest</th>
<th>Cluster</th>
<th>Peak T</th>
<th>MNI Coordinates</th>
<th>Brain Region</th>
</tr>
</thead>
<tbody>
<tr>
<td>Posterior putamen Baseline Analysis (n = 30)</td>
<td>$^{11}$C-PE2I BPND</td>
<td>0.027</td>
<td>22</td>
<td>4.65</td>
<td>0</td>
</tr>
<tr>
<td>Posterior putamen Follow-up Analysis (n = 15)</td>
<td>Δ $^{11}$C-PE2I BPND</td>
<td>0.007</td>
<td>580</td>
<td>5.37</td>
<td>−4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.001</td>
<td>845</td>
<td>4.66</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.031</td>
<td>376</td>
<td>4.39</td>
<td>−36</td>
</tr>
<tr>
<td></td>
<td>Δ Bradykinesia-Rigidity sub-score</td>
<td>0.032</td>
<td>48</td>
<td>6.37</td>
<td>24</td>
</tr>
</tbody>
</table>

Threshold of $p < 0.001$ uncorrected at voxel-level and $p < 0.05$ corrected for false discovery rate at cluster-level.

Coordinates of peak voxels (x, y, z) are given in Montreal Neurologic Institute (MNI) space.

*Indicates correction for false discovery rate.
relevance to dopaminergic deficit. Third, the current study did not control for the potential impact of PD-related atrophy on dopaminergic deficits and resting state functional connectivity. Previous studies in PD have shown a close relationship between atrophy and the degeneration of striatal dopaminergic circuits (Choi et al., 2016), but have failed to find an association with disruptions of functional connectivity (Helmich et al., 2010; Luo et al., 2014). Lastly, patients with PD in this study underwent scanning while in the practically-defined “off” medicated state, but the long-term impact of chronic dopaminergic treatment on resting-state functional connectivity still remains unclear and as such warrants further investigation in future studies.

5. Conclusion

In conclusion, our study demonstrates that functional connectivity of basal ganglia subdivisions in PD is closely related to both the integrity of the dopaminergic system and motor severity. Our findings suggest that basal ganglia functional connectivity in the resting-state may be a powerful imaging biomarker for assessing underlying PD pathology and its progression, and may offer a meaningful way to investigate the mechanisms whereby dopamine deficiency leads to motor symptoms.

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CRediT authorship contribution statement

Weihua Li: Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Software, Validation, Visualization, Writing - original draft, Writing - review & editing. Nick P. Lao-Kaim: Conceptualization, Data curation, Software, Writing - review & editing. Andreas-Antonios Roussakis: Conceptualization, Data curation. Antonio Martin-Bastida: Conceptualization, Data curation. Natalie Valle-Guzman: Conceptualization, Data curation. Gesine Paul: Conceptualization, Data curation, Writing - review & editing. Eyal Soreq: Software. Richard E. Daws: Software. Tom Foltynie: Conceptualization, Data curation, Writing - review & editing. Roger A. Barker: Conceptualization, Data curation, Funding acquisition, Writing - review & editing. Adam Hampshire: Methodology, Supervision, Writing - review & editing. Paola Piccini: Conceptualization, Data curation, Project administration, Resources, Supervision, Funding acquisition, Writing - review & editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Ethical approval

All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Informed consent

Informed consent was obtained from all individual participants included in the study.

Submission declaration and verification

This work has not been published previously, and it is not under consideration for publication elsewhere. Its publication is approved by all authors and that, if accepted, it will not be published elsewhere in the same form, in English or in any other language, including electronically without the written consent of the copyright-holder.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.nicl.2020.102409.

References


