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Impaired Planning in Parkinson’s Disease is Reflected by Reduced Brain Activation and Connectivity

James P. Trujillo,1,2 Niels J.H.M. Gerrits,2,3 Chris Vriend,2,3,4 Henk W. Berendse,3,5 Odile A. van den Heuvel,2,3,4 and Ysbrand D. van der Werf1,2,3,*

1Department of Emotion & Cognition, Netherlands Institute for Neuroscience, An Institute of the Royal Netherlands Academy of Arts and Sciences, Amsterdam, The Netherlands
2Department of Anatomy and Neurosciences, VU University Medical Center (VUMC), Amsterdam, The Netherlands
3Neuroscience Campus Amsterdam (NCA), Amsterdam, The Netherlands
4Department of Psychiatry, VUMC, Amsterdam, The Netherlands
5Department of Neurology, VUMC, Amsterdam, The Netherlands

Abstract: Objective: Parkinson’s disease (PD) often entails impairments of executive functions, such as planning. Although widely held that these impairments arise from dopaminergic denervation of the striatum, not all executive functions are affected early on, and the underlying neural dynamics are not fully understood. In a combined longitudinal and cross-sectional study, we investigated how planning deficits progress over time in the early stages of PD compared to matched healthy controls. We used functional magnetic resonance imaging (fMRI) to identify accompanying neural dynamics. Methods: Seventeen PD patients and 20 healthy controls performed a parametric Tower of London task at two time points separated by ~3 years (baseline and follow-up). We assessed task performance longitudinally in both groups; at follow-up, a subset of participants (14 patients, 19 controls) performed a parallel version of the task during fMRI. We performed meta-analyses to localize regions-of-interest (ROIs), that is, the bilateral dorsolateral prefrontal cortex (DLPFC), inferior parietal cortex, and caudate nucleus, and performed group-by-task analyses and within-group regression analyses of planning-related neural activation. We studied task-related functional connectivity of seeds in the DLPFC and caudate nucleus. Results: PD patients, compared with controls, showed impaired task performance at both time-points, while both groups showed similar performance reductions from baseline to follow-up. Compared to controls, patients showed lower planning-related brain activation together with decreased functional connectivity. Conclusion: These
INTRODUCTION

Although classically considered a motor disorder, Parkinson’s disease (PD) patients also often suffer from cognitive deficits [Aarsland et al., 1999; Kaasinen and Rinne, 2002; Kudlicka et al., 2011; Owen, 2004]. In particular, PD is associated with impairment of the executive functions, including working memory [Kehagia et al., 2010], set shifting [Cools et al., 2001], and planning of goal-directed behaviour [Owen et al., 1995]. Executive functioning refers to higher cognitive functions that strongly rely on the prefrontal cortex and the associated functional circuits that involve the striatum, as well as more posterior regions, such as the inferior parietal cortex [Elliott, 2003]. The striatum is highly connected to the cortex via multiple cortico-striato-thalamo-cortical (CSTC) loops, which are not only involved in motor functions [Voorn et al., 2004], but also in cognitive performance [Chudasama and Robbins, 2006; Elliott, 2003]. Dopaminergic afferents to the striatum critically influence activity within the prefrontal cortical areas involved in executive functions through the CSTC loops [Cools, 2011]; the prefrontal cortex in turn is imbedded in additional cortico-cortical loops, such as the fronto-parietal system, which is especially important in visuospatial working memory [Diwadkar et al., 2000] and spatial planning [Newman et al., 2003].

Executive dysfunction in PD is assumed to be caused, at least partly, by dopaminergic denervation of the striatum [Cools, 2006] and concomitant dysfunction of the CSTC loops. Imaging studies demonstrate that these CSTC loops and fronto-parietal networks are impaired even in early stages of PD before executive impairment at the behavioural level is evident [Monchi et al., 2007; Trujillo et al., 2015]. In early-stage PD, functional connectivity is impaired in both task-related and resting-state networks [Olde Dubbelink et al., 2013, 2014; Stoffers et al., 2008]. As progressive dopamine depletion is associated with further decline in the signal-to-noise ratio between dopaminergic neuronal assemblies [Cools and D’Esposito, 2011; Kroener et al., 2009], decreased functional connectivity in PD is believed to be the result of impaired communication between brain areas [Mattay et al., 2002]. Whether task-related network activation in early stage PD is increased or decreased seems to depend on the characteristics and complexity of the employed cognitive task. In tasks of visuospatial working memory [Trujillo et al., 2015] and set-shifting [Gerrits et al., 2015], for example, PD patients perform the tasks with similar accuracy to that of healthy controls, but show hyperactivation of task-related regions relative to controls. Conversely, behavioural deficits are apparent during verbal working memory tasks that were accompanied by decreased task-related activity [Ekman et al., 2012; Lewis et al., 2003]. These results suggest that in early PD, impaired performance is accompanied by decreased task-related activity, whereas intact behavioural performance is associated with compensatory increased activity.

Proper functioning of these executive circuits is thus necessary for executive tasks such as planning, which involves thinking ahead in a goal-directed manner before taking any action [Jurado and Rosselli, 2007]. An often-used measure of goal-directed planning is performance on the Tower of London (ToL) task, which requires participants to mentally manipulate a configuration of beads stacked on posts of varying lengths to reach an end-goal configuration [Shallice, 1982]. Neuroimaging studies have consistently shown that this task causes robust activation of the bilateral dorsolateral prefrontal cortex, inferior parietal cortex, and the caudate nucleus in healthy populations [Boghi et al., 2006; Dagher et al., 1999; van den Heuvel et al., 2003]. Therefore, the ToL task is a well-suited paradigm to probe CSTC and fronto-parietal functioning in psychiatric and neurological disorders characterized by fronto-striatal and fronto-parietal failure, such as obsessive-compulsive disorder [van den Heuvel et al., 2005] and related anxiety disorders [van den Heuvel et al., 2011], schizophrenia [Eisenberg and Berman, 2010], and PD [Owen, 2004; Williams-Gray et al., 2007]. With our study we sought to expand upon previous findings by using cross-sectional neuroimaging, combining task-related activation, network connectivity, and performance over time.

We therefore used a parametric self-paced visuospatial ToL task [van den Heuvel et al., 2003]. Participants performed this task at two separate time points, that is, at baseline and after ~3 years, to compare performance longitudinally in a group of PD patients as well as a group of matched healthy controls. We additionally administered the test during a functional magnetic resonance imaging (fMRI) session at the second time point, to study task-related neural activity and network connectivity. In this way, we aimed to gain insight into (1) group differences in planning-related neural activation and network connectivity during task performance, and (2) how behavioural performance declines with aging across the groups, contingent upon variable cognitive load. Based on an a priori interest in the fronto-striatal and fronto-parietal systems, we chose a meta-analytic approach for defining and
localizing regions-of-interest (ROIs) specific to our task and contrasts, without being biased by our own results. Based on our meta-analyses of studies using the ToL, we specifically investigated the bilateral dorsolateral prefrontal cortex, bilateral inferior parietal cortex, and bilateral caudate nucleus. Additionally, we assessed task-related functional connectivity of the dorsolateral prefrontal cortex using a generalized form of task-dependent psychophysiological interactions (gPPI) [O’Reilly et al., 2012] analysis.

Muslimović demonstrated that PD patients are already impaired on the ToL task at an early stage [Muslimović et al., 2005] and exhibit a stronger decline in planning accuracy over time when compared to controls [Muslimović et al., 2009]; we therefore hypothesize that our patient group will likewise show decreased task performance with a more pronounced decline in performance at follow-up and, based on the complexity of the task, decreased activity in task-related areas compared to controls. We also expect, based on recent connectivity analyses from our group [Trujillo et al., 2015], decreased task-related network connectivity in the PD group compared to controls.

**MATERIALS AND METHODS**

**Participants**

Participants were recruited as part of a follow-up study from a previously established cohort from our group [Gerrits et al., 2015; Trujillo et al., 2015; Vriend et al., 2015]. Twenty non-demented patients with early stage, mostly de novo (i.e., naive to dopamine replacement therapy) PD and 21 healthy controls participated in this study. Participants performed two parallel versions of a Tower of London task, twice outside the scanner (ToL-out), at two time points separated by an average of 3.16 years, and once in the MR scanner (ToL-in), at the second time point. The study design and exclusion process are graphically depicted in Figure 1a. Outliers were defined as having an overall average task accuracy of more than 1.5 times the interquartile interval below the group median, or below chance level (50% for ToL-in, 20% for ToL-out). After exclusion, our total sample size consisted of 17 PD patients (mean age: 60.2 ± 8.9 years) and 20 healthy controls (mean age: 58.4 ± 9.7 years) in the ToL-out version, and 14 PD patients and 19 healthy controls in the ToL-in version (note that the ToL-in and ToL-out groups, although slightly different, did not differ significantly in their demographic data, and all subjects included in the MRI analysis were also included in the behavioural analyses).

The patients were diagnosed by a movement disorders specialist according to the UK PD Brain Bank criteria [Daniel and Lees, 1992] for idiopathic PD, supported by abnormal dopamine transporter single-photon emission computed tomography (DaT-SPECT) scans in 10 patients. The Unified PD Rating Scale Part III (UPDRS-III) (Fahn et al. 1987) and Hoehn and Yahr stage [Hoehn and Yahr, 1967] were administered to assess disease severity and stage, respectively. We determined disease subtype (i.e., tremor dominant/akinetic) and disease laterisation using the method described by Eggers et al. (2011). At baseline, two patients were already using dopaminergic medication; at follow-up, all patients had begun dopamine replacement therapy. At the time of assessment at follow-up, all patients were on their regular dopaminergic medication, and considered optimal ON. Total levodopa equivalent daily dosage (LEDD) scores were calculated at follow-up [Olde Dubbelink et al., 2013]. All participants were screened for general cognitive status using the Mini-Mental State Examination (MMSE) [Cockrell and Folstein, 1988], depressive symptoms using the Beck Depression Inventory (BDI) [Beck et al., 1996], and anxiety using the Beck Anxiety Inventory (BAI) [Beck et al., 1988]. None of the participants had a score of <24 on the MMSE, of >15 on the BDI [Beck and Beck, 1972], or >25 on the BAI [Beck et al., 1988]. We screened for the presence of psychiatric disorders using the short screening version of the Structured Clinical Interview for DSM-IV Axis-I Disorders (SCID-I) [Spitzer et al., 1992]. The UPDRS-III, Hoehn and Yahr stage, BDI, BAI, SCID-I, and MMSE were administered at both time points. All participants provided informed consent, obtained according to the Declaration of Helsinki, and the study protocol was reviewed and approved by the Medical Ethical Committee of the VU University medical center.

**Task Paradigm**

A pseudo-randomized self-paced version of the ToL was used, discussed in detail elsewhere [van den Heuvel et al., 2003]. The version of the paradigm used outside the scanner consisted of five planning conditions, ranging in solution length from 1 to 5 moves (indicated hereafter as S1–S5), while the MRI version also included a counting condition (indicated hereafter as S0) which was used in the fMRI analysis as control condition. In the planning conditions, subjects saw a starting configuration together with a target configuration, with the instruction to “count the number of steps.” In both conditions, three coloured beads were placed on three vertical posts, which could accommodate up to one, two, or three beads each. One bead could be moved at a time and only when there was no other bead on top. Subjects were requested to determine the minimum number of moves necessary to reach the target configuration and to (a) press on the keyboard the number representing the correct number of steps (ToL-out), or (b) press the button corresponding to the correct answer printed on the screen (ToL-in). In the counting condition, subjects simply had to count the total number of yellow and blue beads on the screen. See Figure 1b for a visual representation of the task layout. The trials were presented in a pseudo-randomized order to prevent...
overflow effects (i.e., perseverance of task-related cognitive processes after a difficult trial), implicating that each trial of three or more moves was followed by a baseline trial. In the scanner, participants started with a short practice session to familiarize them with the task, during which feedback was given on their performance, but no scans were acquired. During the test run, no further feedback was provided. A maximum reaction time (RT) of 45 s for each trial was applied, after which the next stimulus was presented. The ToL-out paradigm consisted of a total of...
100 trials, while the ToL-in paradigm consisted of 90 planning trials and 36 counting trials; both versions lasted on average approximately 20–25 min.

**Image Acquisition**

Functional MRI data were acquired on a Discovery MR750 3.0T MRI scanner (General Electric, Milwaukee) at the VU University medical center using a gradient echo-planar imaging (EPI) sequence (TR = 2100 ms; TE = 50 ms; 64 × 64 matrix; field of view = 24 cm; flip angle = 30°) with 40 descending slices per volume (3.75 × 3.75 mm in-plane resolution; slice thickness = 2.8 mm; inter-slice gap = 0.2 mm), that provided whole-brain coverage. A sagittal three-dimensional gradient-echo T1-weighted sequence (256 × 256 matrix; voxel size = 1 × 0.977 × 0.977 mm; 172 sections) was acquired for co-registering the functional images.

**DATA ANALYSIS**

**Behavioural Data**

We assessed planning performance by calculating the overall percentage of correct responses with each condition per participant. For ToL-out, we compared the two groups longitudinally using a mixed ANOVA with task-load (levels: S1/S2/S3/S4/S5) and time (levels: baseline/follow-up) as within-subject factors and group (levels: PD patients/healthy controls) as between-subject factor. For the ToL-in, we similarly used an ANOVA with task-load (levels: S1/S2/S3/S4/S5) as the within-subject factor and group (levels: PD patients/healthy controls) as between-subject factor. In both analyses, we employed the Greenhouse-Geisser correction when the assumption of sphericity was violated. ToL-out scores at follow-up were compared, per participant. For ToL-in, we compared the two groups employing only a ROI approach were excluded), (3) studies employing only a ROI approach were excluded), (3) they used a similar contrast (planning: all planning-related conditions compared against a baseline; and/or task-load: more difficult conditions compared against easier conditions), (4) they used mental calculation, as opposed to physically transforming the starting configuration into the

**Image Processing and Analysis**

Pre-processing and statistical analyses were performed in SPM8 (Wellcome Department of Imaging Neuroscience, London, UK) running in a Matlab environment (version 7.5, The MathWorks, Natick, MA, 2000). The echo-planar imaging (EPI) images were reoriented and first slice-time corrected, then realigned to the first image and unwarped using a least squares approach and a six parameter (rigid body) spatial transformation to correct for motion. They were subsequently normalised to the Montreal Neurological Institute (MNI) T1-template, employing the co-registered individual T1-weighted image for estimation. Lastly, the images were smoothed with an 8 mm Gaussian kernel.

A design matrix was created to examine within-subject effects in a first level GLM. We employed an event-related design modelling all trials of all six conditions, using total reaction time as the event-length. The six first regressors modelled the correctly answered trials in each of the six conditions of increasing difficulty: “S0,” “S1,” “S2,” “S3,” “S4,” and “S5.” Incorrect trials per condition (when present) as well as the six movement parameters that were calculated during the realignment were added to the model as nuisance covariates. The contrasts of interest were “planning,” defined as all planning conditions against control (S1 S1 S1 S1 S1), and “task-load”, defined across the five planning conditions only, with the vector (0 –2.5 –1.5 –0.5 1 3.5). The weighted nature of the task-load contrast is based on the uneven number of conditions and on the significant increase in reaction time during S4 compared to S3, and between S5 compared to S4 as opposed to the more gradual increase between the lower conditions. These contrasts were used for the whole brain GLM, region of interest, and gPPI analysis to examine the effect of planning cognitive load, corrected for baseline features of the task such as visuospatial processing and motor responses.

Contrast images derived from the first level analyses were used at the second (group) level, employing whole-brain voxel-wise independent t-tests. Brain regions were identified using the WFU-Pick Atlas (Wake Forest University, Winston-Salem, NC, USA) [Maldjian et al., 2003].

**ROI Selection**

For the purpose of ROI selection we employed a selective meta-analysis to find coordinates of task-related activation specifically representative of planning and task-load, separately. On the 6th of November, 2014, the electronic database PubMed was searched using the keywords “Tower of London,” “fMRI” and/or “PET.” Articles were considered for inclusion when (1) they used a similar task configuration, (2) they showed whole-brain results reported in either Talairach or MNI coordinates (i.e., studies employing only a ROI approach were excluded), (3) they used a similar contrast (planning: all planning-related conditions compared against a baseline; and/or task-load: more difficult conditions compared against easier conditions), (4) they used mental calculation, as opposed to physically transforming the starting configuration into the
target configuration, and (5) the study was performed in healthy participants.

The GingerALE 2.3 BrainMap application was used to generate quantitative voxel-wise activation-likelihood estimates (ALE) maps for the contrasts of interest. Input files of study foci were manually created for coordinate-based data in both Talairach and MNI spaces, although the final ALE analysis was performed in MNI space. Any coordinates originally reported in Talairach space were therefore automatically converted to MNI space by the GingerALE software using the icbm2tal algorithm software. ALE values were computed for each voxel in the brain using the input foci. Based on the number of subjects, a Gaussian blur is then calculated for each foci group (study/paper), and the final ALE map is calculated by finding the union of these individual maps, as described by Eickhoff et al. [2011]. The resulting ALE maps then undergo cluster-level thresholding using simulated datasets with the same characteristics as the true dataset and a False Discovery Rate of 0.05. This cluster-level corrected threshold ensures that only 5% of the simulated data will exceed the calculated minimum cluster volume. The resultant clusters are representative of areas with a high likelihood of task-related activation and are assigned anatomical labels at their weighted centers by the Talairach Daemon.

To investigate the primary regions of the fronto-striatal and fronto-parietal systems, we selected coordinates associated with the bilateral inferior parietal cortex, bilateral caudate nucleus, and bilateral dorsolateral prefrontal cortex from the list of task-related clusters of the meta-analysis. These six ROIs were selected separately from the meta-analyses on the planning (vs. baseline) as well as the task-load contrasts, resulting in a total of 12 ROIs (six for planning, six for task-load). Coordinates for these regions are given in Table I.

MarsBaR (http://marsbar.sourceforge.net) was used to create spherical ROIs, centered around the meta-analysis coordinates, with a 10 mm diameter. The average parameter estimates (beta weights) of the whole ROI were then extracted from the corresponding contrast per participant and were subsequently compared, first using a repeated-measures ANOVA with region as within-subjects factor and group as between-subject factor. Significant group effects were further analyzed with post-hoc multivariate ANOVAs.

Time courses were extracted from each of the individual ROIs, using the corresponding contrast, creating volumes of interest that were used in the gPPI analyses. These were extracted at a threshold of $P = 0.5$ to ensure robust time-series.

### Functional Connectivity

We assessed the task-related functional connectivity of the bilateral DLPFC and bilateral caudate nucleus using a generalized form of context-dependent psychophysiological interaction (gPPI) [O’Reilly et al., 2012]. A PPI analysis statistically tests in a whole-brain voxel-wise manner whether areas were functionally connected with the seed-region during task performance [O’Reilly et al., 2012]. We chose gPPI, instead of the traditional PPI [Friston et al., 1997], as it allowed us to model all psychological task conditions into one first-level design, thus improving the model-fit [McLaren et al., 2012]. For the purpose of the gPPI analyses, the intersection between the ROI mask and the 1st level mask was individually calculated per subject by applying SPM imcalc with trilinear interpolation to these masks.

Our first-level model included the correct trials for the six task conditions, incorrect trials per condition, the six convoluted PPI terms, the time-series of the seed-region, and the six movement parameters. We defined the planning and task-load contrasts in the same way as for the GLM analysis, this time using the convoluted PPI terms and leaving the psychological variable (task conditions) and movement parameters as nuisance variables. For the eight separate analyses (left and right DLPFC—planning, task-load; left and right caudate nucleus—planning, task-load), all subjects in the GLM analysis could be included and the psychological variable (task conditions) and movement parameters as nuisance variables. For the eight separate analyses (left and right DLPFC—planning, task-load; left and right caudate nucleus—planning, task-load), all subjects in the GLM analysis could be included in the gPPI analysis, since all subjects exhibited significant BOLD effects in the ROI and had voxels that were functionally connected to the seed regions.

At second level, we compared the contrasts between groups using an independent samples t-test and an uncorrected statistical threshold of $P < 0.001$ with a spatial extent threshold of $k > 5$. Group differences (i.e., voxels more active in one group compared to the other) were masked inclusively with the main effects of the group of interest, so that only voxels that were also found in the main effects are reported in the group interactions. The same analysis procedures were employed for all regions and contrasts.

### Correlation with Dopamine Transporter Binding

For 10 of the included PD patients, single-photon emission computerized tomography (SPECT) scans with a $[^{123}]N$-fluoropropyl-2β-carbomethoxy-3β-(4-iodophenyl) nortropaine ($[^{123}]$FP-CIT) tracer binding to the dopamine transporter (DaT) were available with an average interval between SPECT acquisition and baseline assessment of 55

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**TABLE I. MNI coordinates for ROI analyses**

<table>
<thead>
<tr>
<th></th>
<th>Planning</th>
<th>Task-load</th>
</tr>
</thead>
<tbody>
<tr>
<td>L DLPFC</td>
<td>-40</td>
<td>-2</td>
</tr>
<tr>
<td>R DLPFC</td>
<td>32</td>
<td>34</td>
</tr>
<tr>
<td>L caudate</td>
<td>-12</td>
<td>10</td>
</tr>
<tr>
<td>R caudate</td>
<td>12</td>
<td>8</td>
</tr>
<tr>
<td>L IPC</td>
<td>-58</td>
<td>-38</td>
</tr>
<tr>
<td>R IPC</td>
<td>48</td>
<td>-42</td>
</tr>
</tbody>
</table>

L: left, R: right, DLPFC: dorsolateral prefrontal cortex, IPC: inferior parietal cortex.
TABLE II. Mean and standard deviations for demographic and clinical data at follow-up

<table>
<thead>
<tr>
<th></th>
<th>HC (n = 20)</th>
<th>PD (n = 17)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demographics</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age (years)</td>
<td>58.38 ± 9.67 (40 – 74)</td>
<td>60.18 ± 8.91 (40 – 76)</td>
<td>P = 0.56</td>
</tr>
<tr>
<td>Years between measures</td>
<td>3.29 ± 1.13 (1.5 – 5.1)</td>
<td>3.01 ± 0.9 (1.8 – 5.2)</td>
<td>P = 0.47</td>
</tr>
<tr>
<td>Gender (% male)</td>
<td>14 (70.0%)</td>
<td>11 (64.7%)</td>
<td>P = 0.73</td>
</tr>
<tr>
<td>Handedness (% right)</td>
<td>19 (95.0%)</td>
<td>14 (82.4%)</td>
<td>P = 0.39</td>
</tr>
<tr>
<td>Education</td>
<td>6.00 ± 1.00 (5 – 7)</td>
<td>6.00 ± 2 (3 – 7)</td>
<td>P = 0.57</td>
</tr>
<tr>
<td>Clinical measures</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MMSE</td>
<td>29.00 ± 1.00 (26 – 30)</td>
<td>28.00 ± 1.50 (24 – 30)</td>
<td>P = 0.02*</td>
</tr>
<tr>
<td>MMSE-Δ</td>
<td>0.002 ± 0.04 (−0.07 – 0.07)</td>
<td>−0.02 ± 0.05 (−0.07 – 0.08)</td>
<td>P = 0.24</td>
</tr>
<tr>
<td>BDI</td>
<td>2.00 ± 5.25 (0 – 7)</td>
<td>4.00 ± 8.00 (1 – 15)</td>
<td>P = 0.01*</td>
</tr>
<tr>
<td>BAI</td>
<td>0.50 ± 3.25 (0 – 7)</td>
<td>6.00 ± 8.00 (2 – 25)</td>
<td>P &lt; 0.01*</td>
</tr>
<tr>
<td>UPDRS-III</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>27.29 ± 7.82 (16 – 40)</td>
<td>1.28 ± 4.23 (−0.27 – 16.50)</td>
<td></td>
</tr>
<tr>
<td>UPDRS-III-Δ</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Subtype (% mixed/% akinetic)</td>
<td>4 (23.5%)/13 (76.5%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lateralization (% mixed/% left)</td>
<td>15 (88.2%)/2 (11.8%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hoehn and Yahr</td>
<td>2 ± 0.5 (1.5 – 3)</td>
<td></td>
<td></td>
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<tr>
<td>LEDD</td>
<td>450 ± 255 (230 – 1760)</td>
<td></td>
<td></td>
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<tr>
<td>Behavioural measures</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Task performance</td>
<td>89.5 ± 16.25 (62 – 96)</td>
<td>82 ± 18.92 (44 – 96)</td>
<td>P = 0.01*</td>
</tr>
<tr>
<td>Task reaction time (seconds)</td>
<td>9.94 ± 2.06 (6.88 – 15.29)</td>
<td>11.11 ± 4.08 (7.5 – 24.8)</td>
<td>P = 0.02*</td>
</tr>
<tr>
<td>Task performance-Δ</td>
<td>−0.01 ± 0.07 (−0.21 – 0.06)</td>
<td>−0.07 ± 0.12 (−0.30 – 0.07)</td>
<td>P = 0.43</td>
</tr>
<tr>
<td>Reaction time-Δ</td>
<td>−0.05 ± 0.10 (−0.33 – 0.14)</td>
<td>−0.05 ± 0.16 (−0.28 – 0.25)</td>
<td>P = 0.83</td>
</tr>
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(= 230) to 1760). For an overview of demographic and clinical scores at follow-up, see Supporting Information Table I.

RESULTS

Demographics

Our groups were matched for age, gender, handedness and the interval between baseline and follow-up (see Table II). Although PD patients had significantly higher anxiety (535.0, P < 0.001) and depression (835.0, P = 0.01) scores at follow-up when compared to controls, both groups’ scores are still in the lower range of the scale and are not clinically relevant. PD patients, compared to controls, also had lower MMSE scores, both at baseline and follow-up, though none fell below the typical cut-off score for dementia of 23. Patients and controls did not differ significantly in their decline in MMSE scores between baseline and follow-up. At follow-up, the mean UPDRS-III score and median Hoehn and Yahr stage were 27 and 2, respectively. Whereas none but two patients had started dopamine replacement therapy at baseline, at follow-up all patients were using dopaminergic medication with an average LEDD of 450 ± 230 (range: 230–1760). For an overview of demographic and clinical scores at follow-up, see Table II; for clinical scores at baseline, see Supporting Information Table I.

Behavioural Results, Based on ToL-out data

Accuracy

Wilcoxon signed-rank tests showed that follow-up ToL-out accuracy scores were comparable to ToL-in scores both for the controls (Z = −0.71, P = 0.48), as well as for the PD
patients (Z = −1.03, P = 0.30). Follow-up ToL-in reaction times were significantly faster than ToL-out scores both for the controls (Z = −3.92, P < 0.01), as well as for the PD patients (Z = −2.89, P < 0.01). The omnibus test, containing performance of both groups at both time points across all difficulty levels, showed that PD patients, compared to controls, performed significantly worse on the task across both time points (F(1, 35) = 6.86, P = 0.013), that both groups performed worse with increasing task-load (F(2,51, 87.96) = 83.66, P < 0.001), and that both groups performed overall worse at follow-up compared to baseline (F(1, 35) = 10.89, P = 0.002). The overall load-effect differed between the patients and controls (F(2,51, 87.96) = 8.92, P < 0.001), with patients having more difficulty with increasing task-load. The effect of task-load was also different between the two time points, regardless of group (F(2,66, 92.96) = 4.73, P = 0.006), with a stronger decline with increasing task-load at follow-up.

For both groups, the interval between baseline and follow-up did not correlate with performance-Δ scores (PD: r² = 0.15, P = 0.58; HC: r² = −0.02 P = 0.95). Better task accuracy at follow-up was related to lower anxiety scores in the control group (r² = −0.55, P = 0.02), though not for the patients (r² = 0.25, P = 0.35). In patients task accuracy at follow-up did not correlate significantly with LEDD (r² = 0.18, P = 0.49), UPDRS-III scores (r² = 0.05, P = 0.39), or Hoehn and Yahr stage (r² = −0.24, P = 0.35).

**Reaction time**

We found that PD patients, compared with controls, performed more slowly (F(1, 35) = 5.46, P = 0.029). Across both groups we also found faster reaction times at follow-up compared to baseline (F(1, 35) = 6.01, P = 0.019), as well as an overall main effect of task-load on reaction time, with longer reaction times at higher task-loads (F(1,87, 65.27) = 308.66, P < 0.001), that was more pronounced at follow-up compared to baseline (F(1,57, 54.95) = 7.45, P = 0.003). Group did not interact with task-load (F(1,86, 65.27) = 1.09, P = 0.34), or with time point, that is, baseline versus follow-up (F(1, 35) = 0.08, P = 0.78), nor was there an interaction between group and the time-by-task-load interaction (F(1,57, 54.95) = 2.38, P = 0.11).

On average, higher accuracy on the task correlated with longer reaction time both for the controls (r² = 0.37, P = 0.005), as well as for the patients (r² = 0.37, P = 0.009). This correlation did not differ significantly between groups (Z = −0.03, P = 0.98).

For an overview of behavioural data at baseline, see Supporting Information Table I.

**Imaging Results: Regions of Interest**

We found a significant difference in the correlation between age and average activation in the left and right IPC in the planning condition, (left: Z = −2.99, P < 0.01; right: Z = −2.44, P = 0.01). We therefore included age as a covariate in all ROI analyses to control for possible confounding effects.

Across all planning-related ROIs, we found a significant main effect of group, F(1,30) = 5.73, P = 0.023, with PD patients displaying decreased activity (see Fig. 2). We found no interaction between group and region (F(3,01, 90.22) = 0.48, P = 0.69). Based on the average correlation of r = 0.36 between regions, we calculated an adjusted alpha of 0.02 for testing group differences for the individual ROIs. Under this adjusted alpha, post-hoc analyses using ANOVA revealed no significant group differences for any of the individual ROIs, although three showed trends toward being significantly higher in controls; these regions being the left caudate nucleus (F(1, 30) = 3.478, P = 0.07), the right DLPFC (F(1,30) = 4.33, P = 0.05), and the right IPC (F(1, 31) = 4.25, P = 0.05). For an overview of whole-brain results relating to the planning contrast, see Supporting Information Table II.

Across all load-related ROIs we also found a significant main effect of group (F(1, 30) = 4.85, P = 0.04), again with PD patients showing decreased task-related activity in comparison with the controls (see Fig. 3). After applying the Greenhouse-Geisser correction, we also found a significant effect of task-load on ROI activity (F(2,67, 80.16) = 3.56, P = 0.02). We found no group-by-task-load interaction (F(2,67, 80.16) = 1.74, P = 0.17), nor did we find a group-by-region interaction (F(3,15, 94.38) = 1.81, P = 0.15). For an overview of the whole-brain results relating to the task-load contrast, see Supporting Information Table III.
Imaging Results: Functional Connectivity

Main effects

In the control group, the left DLPFC showed planning-related connectivity with the right caudate nucleus, right precuneus (BA 7), right posterior cingulate cortex (BA 23 & 31) and the right middle frontal gyrus (BA 6 & 10), and load-related connectivity with the bilateral precuneus (BA 7), left superior temporal gyrus (BA 22 & 39) and left ventrolateral part of the thalamus. In the PD group, the left DLPFC showed planning-related connectivity with the left parahippocampal gyrus and the right insula; we found no load-related connectivity in the PD group for the left DLPFC.

In the control group, the right DLPFC only showed planning-related connectivity with the brainstem, but load-related connectivity with the left precuneus (BA 7), left IPC, and right posterior cingulate cortex (BA 31). Compared to the PD group, the control group showed significantly stronger right DLPFC load-related connectivity with the bilateral IPC, left precuneus (BA 7), left superior parietal cortex (BA 7), and right middle occipital gyrus. The PD group showed planning-related right DLPFC connectivity with the left inferior frontal gyrus (BA 47), and no load-related connectivity.

In the control group, we found no significant planning-related connectivity with the left caudate nucleus. In the PD group, we found planning-related connectivity between the left caudate and left IPC (BA 40), left inferior frontal gyrus (BA 46), left precentral gyrus (BA 44), and several posterior regions.

In the control group, the right caudate nucleus showed planning-related connectivity with the left inferior frontal gyrus (BA 45 & 46), right middle frontal gyrus (BA 9) and the left ventrolateral nucleus of the thalamus. In the PD group, the right caudate nucleus showed planning-related connectivity with the bilateral IPC (BA 40), left inferior frontal gyrus (BA 44 & 46), left middle frontal gyrus (BA 10), right posterior cingulate cortex (BA 31), and left precuneus (BA 7).

For an overview of the main effects by group of DLPFC connectivity see Supporting Information Tables IV and V; for the main effects by group of caudate nucleus connectivity, see Supporting Information Tables VI and VII.

Group differences

Compared to the PD group, the control group showed significantly stronger load-related right DLPFC connectivity with the left precuneus (BA 7), bilateral IPC, left superior parietal cortex (BA 7), and right middle occipital gyrus.

Dopamine-Transporter Binding

Although not directly correlated with task scores at follow-up ($r^2 = 0.48$, $P = 0.16$), higher DaT-SPECT ratios correlated significantly with a more positive relative change in accuracy scores from baseline to follow-up ($r^2 = 0.69$, $P = 0.03$, see Fig. 4).
Our finding of impaired task performance is in agreement with previous studies on planning in PD [Muslimović et al., 2005; Owen et al., 1992, 1998]. In contrast to our hypothesis with previous studies on planning in PD [Muslimović et al., 2009], however, we found similar performance decline over time in both patients and controls. This may be due to a slightly younger group of patients in our study (mean: 60 years at follow-up) compared to those showing decline in Muslimovic’s group (mean: 69 years at follow-up). Although the change in task accuracy over the course of the study was similar in both patients and controls, we additionally showed that in patients this change was predicted by the amount of dopamine-transporter binding at baseline. Our results show that patients with higher baseline DaT-binding ratios in the dorsomedial striatum had a less pronounced decline in performance from baseline to follow-up compared to those with lower DaT-binding; this indicates that baseline dopamine-transporter levels may not predict performance per se, but rather the resilience to cognitive decline; or inversely stated, lower baseline dopamine-transporter levels signify a higher rate of subsequent disease-related cognitive decline. It may also indicate that these patients, in terms of striatal dopaminergic deterioration, were simply less affected by the disease at baseline and therefore had a larger reserve serving to protect from cognitive deterioration. It is possible that this relationship is due to a selection bias, as several patients dropped out between baseline and follow-up. Erixon-Lindstroth and colleagues, however, previously demonstrated that striatal dopamine loss mediates age-related cognitive decline [Erixon-Lindstroth et al., 2005] and our results complement this finding by showing that striatal dopamine denervation—already present in early PD—may also predict the rate of cognitive decline. The parallel decline in cognitive performance with the healthy control group may also be due to the addition of dopaminergic medication, initiated after baseline measurement, which may have helped patients maintain cognitive performance over time. Such interpretations require further confirmation, for example by comparing cognitive performance in the ON and OFF states.

Studies of executive functioning in early-stage PD show seemingly incongruous results: some present increased activation while others present decreased activation in task-related areas. When paired with performance, however, increased activity appears to be associated with preserved task performance [Gerrits et al., 2015; Trujillo et al., 2015], while decreased activity is present when task performance is already significantly impaired [Ekman et al., 2012; Lewis et al., 2003]. This suggests that hyperactivation is a compensatory mechanism associated with intact behavioural performance, which was not present in our sample of PD patients, even at baseline. Our finding of decreased activation is therefore in line with previous studies in which cognitive deficits are already present. An alternative explanation for decreased task-related activity is that baseline activity is significantly increased; as increased activity would already be present at baseline, the task would not cause as large of an increase as seen in healthy subjects, resulting in the finding of decreased task-specific activity [Ko et al., 2013].

**DISCUSSION**

We initially hypothesized that patients would show impaired planning performance and a stronger decrease in performance over time compared with controls, concomitant with hypoactivation of task-related areas and decreased connectivity in task-related networks. Across both time points patients performed overall worse than healthy controls and had more difficulty with increasing task-load. Nevertheless, age-related decline was comparable between groups. In line with our hypothesis, PD patients, compared with controls, showed lower task-related network activation both related to planning as well as to increasing task-load. Lower task-related functional connectivity was seen in the fronto-parietal network of PD patients.

As expected, our meta-analyses and whole-brain analyses demonstrated a strong recruitment of the fronto-parietal and fronto-striatal systems during the ToL task, both for general planning as well as for increasing task-load, although the specific locations of the foci differed slightly between planning and task-load. This may indicate a finer functional segregation in these areas for dealing with general planning as opposed to increasing cognitive load.

Our finding of impaired task performance is in agreement with previous studies on planning in PD [Muslimović et al., 2005; Owen et al., 1992, 1998]. In contrast to our hypothesis and the findings of Muslimović et al. [2009], however, we found similar performance decline over time in both patients and controls. This may be due to a slightly younger group of patients in our study (mean: 60 years at follow-up) compared to those showing decline in Muslimovic’s group (mean: 69 years at follow-up). Although the change in task accuracy over the course of the study was similar in both patients and controls, we additionally showed that in patients this change was predicted by the amount of dopamine-transporter binding at baseline. Our results show that patients with higher baseline DaT-binding ratios in the dorsomedial striatum had a less pronounced decline in performance from baseline to follow-up compared to those with lower DaT-binding; this indicates that baseline dopamine-transporter levels may not predict performance per se, but rather the resilience to cognitive decline; or inversely stated, lower baseline dopamine-transporter levels signify a higher rate of subsequent disease-related cognitive decline. It may also indicate that these patients, in terms of striatal dopaminergic deterioration, were simply less affected by the disease at baseline and therefore had a larger reserve serving to protect from cognitive deterioration. It is possible that this relationship is due to a selection bias, as several patients dropped out between baseline and follow-up. Erixon-Lindstroth and colleagues, however, previously demonstrated that striatal dopamine loss mediates age-related cognitive decline [Erixon-Lindstroth et al., 2005] and our results complement this finding by showing that striatal dopamine denervation—already present in early PD—may also predict the rate of cognitive decline. The parallel decline in cognitive performance with the healthy control group may also be due to the addition of dopaminergic medication, initiated after baseline measurement, which may have helped patients maintain cognitive performance over time. Such interpretations require further confirmation, for example by comparing cognitive performance in the ON and OFF states.

Studies of executive functioning in early-stage PD show seemingly incongruous results: some present increased activation while others present decreased activation in task-related areas. When paired with performance, however, increased activity appears to be associated with preserved task performance [Gerrits et al., 2015; Trujillo et al., 2015], while decreased activity is present when task performance is already significantly impaired [Ekman et al., 2012; Lewis et al., 2003]. This suggests that hyperactivation is a compensatory mechanism associated with intact behavioural performance, which was not present in our sample of PD patients, even at baseline. Our finding of decreased activation is therefore in line with previous studies in which cognitive deficits are already present. An alternative explanation for decreased task-related activity is that baseline activity is significantly increased; as increased activity would already be present at baseline, the task would not cause as large of an increase as seen in healthy subjects, resulting in the finding of decreased task-specific activity [Ko et al., 2013].

![Figure 4.](image-url)
Analysis of functional connectivity revealed that while both groups generally showed functional connectivity within the task-related systems, the control group displayed more connectivity within the fronto-parietal system when compared to the PD group. This may in part explain the impaired performance seen in PD patients, as the fronto-parietal system is implicated in successful ToL problem solving [Baker et al., 1996; Newman et al., 2003]. The parietal regions found to be more strongly connected to the DLPFC in controls also overlapped with our bilateral IPC ROIs, providing further support for a disease-related impairment in the fronto-parietal system specifically activated by this task. Within-group effects of the PD patients seem to suggest that functional connectivity of the caudate nucleus is quite pronounced; we suggest caution, however, in interpreting these results as this is only apparent in the within-group effects (PD task > baseline), and not between-groups (PD > HC).

**Strengths and Limitations**

Our study provides valuable insights into the longitudinal effects of disease progression on planning capabilities in PD, and how striatal dopamine may play a role in this process. This finding is, however, based on a smaller subset of our patient group, and should thus be interpreted with caution. An advantage of our study is that all but two of the patients were naïve to dopaminergic medication at the time of the first measurement. Performing a meta-analysis provided an unbiased, objective approach to localizing our regions of interest, increasing the validity of our results. While Muslimović et al. [2009] previously demonstrated that ToL performance is impaired in early-stage PD and that cognitive decline over time is stronger in PD compared to controls [Muslimović et al., 2009], we extend these results by contributing the effect of task-related activity and connectivity, as well as by showing the predictive value of dopamine transporter binding ratios. As not all participants from the baseline study returned for the follow-up, a selection bias is possible. This may have led to, for example, only the inclusion of patients with relatively stable, or slow, disease progression. Our study also only employed fMRI at the second time-point, and DaT-SPECT at baseline, limiting what can be said about the progression of neural dynamics or dopamine neurotransmitter system integrity across time. As our study did not employ an ON versus OFF design, the effects of dopamine replacement therapy on task-related neural activity or performance cannot be fully assessed.

**CONCLUSIONS**

Goal-directed behaviour, operationalized using a planning task, is already significantly impaired in the early stages of PD. Similar to other PD-related cognitive dysfunctions, this impairment in planning is accompanied by decreased activity and functional connectivity in task-related brain networks. Possibly as a result of dopaminergic treatment, the rate of disease-related cognitive decline remains similar to the effect of normal aging in control subjects, especially for those with more intact striatal dopamine levels at baseline.

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