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Anterior Temporal Atrophy and Posterior Progression in Patients with Parkinson’s Disease

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Key Words
Parkinson’s disease · Voxel-based morphometry · Atrophy

Abstract
Background: Parkinson’s disease (PD) is characterized by specific motor and nonmotor impairments. This suggests that PD is characterized by disease-specific regional cortical atrophy. Given the change of symptoms over time, a concurrent increase in regional atrophy may further be assumed to reflect the dynamic process of disease progression. Methods: In this study we retrospectively collected T1-weighted MRI scans from previous studies performed in our center, enabling the comparison of gray matter atrophy in 77 PD patients with 87 controls using voxel-based morphometry (VBM). This large VBM analysis provided the opportunity to investigate cortical atrophy in relation with disease progression. Results: We found significant PD-related reductions of gray matter density bilaterally in the anterior temporal cortex, the left inferior frontal and left extrastriate visual cortex, independent from normal aging. The anterior temporal cortex did not show major progression, whereas particularly the posterior parts of the lateral temporal cortex and adjacent extrastriate visual cortex occurred at a later stage of disease. Conclusions: Temporal pole atrophy as an early sign of PD is consistent with the PD pathology classification of Braak. The initial anterior temporal atrophy with spread to occipitotemporal and posterior parietal regions may subserve ‘emotion-based’ sensorimotor transformations and deficits in the visual domain, respectively, which may be regarded as premotor symptoms.

Introduction
The main pathology in idiopathic Parkinson’s disease (PD) is degeneration of the substantia nigra and its dopaminergic projections to the striatum [1], which is a major cause of the motor symptoms in this disease. This is associated with misfolding of α-synuclein [2]. It is increasingly recognized, however, that PD not only involves the motor system, but inflicts deterioration in multiple domains. For instance, olfactory dysfunction is considered as a relatively early or even preclinical sign in PD [3], while in later stages pathological changes in olfaction-related brain areas have been described [4]. Cortical atrophy has been demonstrated to correlate with olfactory dysfunction in PD [5]. Especially in later stages, cognitive impairment, including visuospatial dysfunc-
tion and alterations in executive functions, occurs relatively often in PD [6]. Such cortical dysfunction suggests that PD is characterized by a disease-specific regional atrophy. Given the change of symptoms over time, an associated change in the distribution of regional atrophy may further be assumed to reflect the dynamic process of disease progression. In a network view of the brain, it is reasonable to think that initial subcortical damage in PD may inflict cortical impairments in regions that receive projections from such subcortical sources [7]. One may thus hypothesize that long-term hypoactivation of the cortex by subcortical structures leads to cortical atrophy.

Voxel-based morphometry (VBM) studies in PD typically comprised small subject groups and yielded variable and even conflicting results [8]. In the present study, we retrospectively collected 3-tesla MRI images of 77 PD patients that were included in previous studies performed at our center. This resulted in one of the largest VBM analyses comparing patients with PD and controls. Given the variation in disease duration of these patients, it also provided the opportunity to investigate cortical atrophy as an index of disease progression.

**Methods**

**Subjects**

T1-weighted images were retrospectively collected from all 7 studies performed in our center between 2007 and 2013 (104 controls, 111 PD patients). These studies concerned only 1 VBM study [9] and 6 different functional MRI experiments that included the acquisition of a T1-weighted anatomical brain image used for coregistration and the assessment of general brain characteristics. Since some subjects participated in more than 1 study, we only used the last scan made in order to assess the patient scans with the most pronounced characteristics of disease. As 34 PD patients and 8 controls participated in 2 or more studies, the control scans outnumbered the included PD scans. This provided the opportunity to make a selection of control subjects in order to get an optimally age-matched control group. To that end, we inspected the age distribution of both groups, after excluding the first of the possible double scans, and subsequently excluded 9 controls that were not in the same age group as the PD patients. This resulted in a total of 87 controls and 77 patients with PD who were included for further analysis. Patients met the criteria of the UK Parkinson’s Disease Rating Scale (UPDRS) [10]. The Mini-Mental State Examination (MMSE) was used as a screening tool for cognitive functions. An MMSE ≥24 was considered to represent unimpaired cognition. Subjects suffering from neurological disorders other than PD or psychiatric disorders were excluded. All studies were approved by the local Medical Ethical Committee. Participants gave their written informed consent.

**Image Acquisition**

Data acquisition was performed using a 3-tesla Philips MR system (Best, the Netherlands) with a standard 8-channel SENSE head coil. In 3 of the 6 studies (49 controls, 45 PD patients) T1-weighted 3-dimensional anatomical scans were acquired with field of view 232 × 256 × 170, TR = 9.0 ms, TE = 3.5 ms, flip angle 8°, 170 slices without slice gap and voxel size 0.9 × 1 × 1 mm. The other 3 studies (38 controls, 32 PD patients) acquired T1-weighted scans with field of view 256 × 160 × 204 mm, TR = 25 ms, TE = 4.6 ms, flip angle 30°, 160 slices with slice gap –1 and voxel size 1 × 1 × 1 mm. The two acquisition protocols were equally distributed over the groups.

**Image Analysis**

VBM is a method applied in MRI data sets, which offers the opportunity to compare the entire cortex across different groups without a priori assumptions on a subjective region of interest selection [11]. VBM enables the detection of differences in gray matter concentration on a local scale, while correcting for global shape differences [11]. Image processing and voxel-based statistical analysis were conducted using Statistical Parametric Mapping, version 8 (2009, Wellcome Department of Cognitive Neurology, London, UK; http://www.fil.ion.ucl.ac.uk/spm). Raw images were manually reoriented to put the middle of the anterior commissure at the coordinate x0,y0,z0 position. The images were processed using the unified segmentation approach with standard settings [12]. In this approach the images are segmented in gray matter, white matter and cerebrospinal fluid, bias corrected and spatially normalized to the Montreal Neurological Institute brain reference space, based on nonlinear registration with tissue probability maps. The segmented gray matter images were modulated and smoothed (full width at half maximum 8). We used the optimized VBM method in SPM8, because it is widely used and there is extensive experience with this method. An alternative might have been to use DARTEL. It has, however, been shown that DARTEL not necessarily results in better image registration in patients with Alzheimer’s disease, frontotemporal dementia and semantic dementia, while it may be less sensitive than the method we used [13]. On the other hand, DARTEL has been reported to be more sensitive for detecting hippocampal changes in acute depression [14].

**Statistical Analysis of Behavioral Parameters**

MMSE scores and age were not normally distributed and therefore tested for differences between the groups with the Mann-Whitney U test. Gender was compared using the $\chi^2$ test. A p value <0.05 was considered statistically significant.

**Image Statistical Analysis**

Total gray matter was calculated per subject and used as a covariate in the model to remove variance due to differences in brain size, because we were interested in regional gray matter differences. Also, the type of T1 scan was used as a covariate to correct for possible effects of differences in scanning parameters. Initial analysis assessed the presence of regional differences between patients and controls at voxel level $p < 0.001$ to see if there were any differences. Resulting clusters of regional difference were regarded statistically significant at $p < 0.05$ family-wise error (FWE) corrected with a cluster extent of 8 voxels. We compared healthy controls ($n = 87$) with PD patients ($n = 77$) on local gray matter density changes with ANOVA (flexible factorial design). To check wheth-

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er the perceived atrophy was not merely the consequence of normal aging we similarly compared the 25 youngest controls with the 25 oldest controls. In addition, we compared the 87 healthy controls with the 25 PD patients with the shortest duration of disease (PD_short) and the 25 patients with the longest disease duration (PD_long), respectively. These comparisons were expected to provide insight into disease progression. An additional comparison between the 25 PD_short and 25 PD_long patients complemented the two comparisons between patients and control subjects. The reason to introduce this binary split was to get a clear distinction between the subgroups with short and long disease duration, leaving out patients with intermediate disease duration. The large patient group indeed allowed such a selection.

**Results**

There were no significant differences between PD patients and healthy controls regarding age and gender, while the MMSE score was slightly but significantly lower in the patient group (table 1). None of the participants had an MMSE below 24.

**VBM of Patients with PD versus Controls**

Comparison of PD patients with healthy controls revealed significant reductions of regional gray matter density which was most pronounced in anterior temporal regions (fig. 1a; table 2). Adjacent to the cluster of left anterior temporal atrophy, inferior frontal atrophy was seen, while in the right hemisphere posterolateral frontal atrophy was seen at a more dorsal position. In posterior parts of the brain, cortical atrophy was identified in both ventrolateral and dorsal extrastriate visual areas of particularly the left hemisphere, together with significant right precuneus atrophy. At a relaxed threshold of p < 0.001 voxel level uncorrected, the revealed pattern of reduced right occipital and parietal gray matter density was similar to that in the left hemisphere. There were no regions identified with more gray matter atrophy in controls compared to PD patients.

**Progression of Cortical Atrophy in Patients with PD**

Statistical analysis showed that disease duration of the 25 PD_short patients (mean duration 3.3 years, SD 1.2) was indeed significantly shorter compared to the PD_long patients (mean 13.2 years, SD 4.1; p < 0.001). These two groups did not differ in age and MMSE. Mean age of PD_short was 61.8 years (SD 8.8) with a mean MMSE of 27.6 (SD 1.3) while it was 63.0 years (SD 7.1) for PD_long and MMSE 27.4 (SD 1.3) (p = 0.60 for each parameter). Neither did the two groups differ in gender (15 males in PD_short and 16 males in PD_short, p = 0.77). The groups had an equal distribution in the type of T1 scan (17 of the first sequence described in the Methods).

Compared to the control group, the 25 PD_short patients had cortical atrophy of both temporal poles (p < 0.001, voxel level uncorrected; fig. 1b). This difference in gray matter density did, however, not reach statistical significance at cluster level corrected for the entire brain volume (FWE). When the 25 PD_long patients were compared with the controls, significant gray matter reductions were seen at the two temporal poles and in left extrastriate cortical regions (p < 0.05, FWE corrected; fig. 1c).

Complementary to comparing the control subjects with successively the PD_short and PD_long patients, a direct comparison of PD_short with PD_long showed a progression of regional atrophy in the PD_long group at the lateral occipitotemporal junction, bilaterally, with a maximum in the right middle temporal gyrus (x 54, y –42, z –4; p < 0.001, voxel level uncorrected; fig. 1d; table 3). No differences were seen at the temporal poles. The clusters of advanced atrophy did, however, not reach significance at the cluster level corrected for the entire brain volume. This was also the case for changes in the right middle frontal gyrus (BA 45) and right supramarginal gyrus (BA 2).

**Comparison of Old and Young Controls**

In comparison with the 25 youngest controls (mean age 51.3 years, SD 4.3), the 25 oldest control subjects (mean age 68.1 years, SD 3.7) had a reduction of gray matter density in both the midcingulate cortex and superior surface of the left temporal lobe along the sylvian fissure (online suppl. fig. 1; for all online suppl. material, see www.karger.com/doi/10.1159/000363245). This differ-

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**Table 1. Subject characteristics**

<table>
<thead>
<tr>
<th></th>
<th>Controls (n = 87)</th>
<th>PD (n = 77)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>60.1 (7.2)</td>
<td>63.0 (10.5)</td>
<td>0.07</td>
</tr>
<tr>
<td>Gender, male</td>
<td>50</td>
<td>46</td>
<td>0.42</td>
</tr>
<tr>
<td>MMSE (0–30)</td>
<td>28.0 (1.0)</td>
<td>27.0 (1.3)</td>
<td>0.00</td>
</tr>
<tr>
<td>UPDRS III (0–56)</td>
<td>n.a.</td>
<td>24.3 (7.5)</td>
<td></td>
</tr>
<tr>
<td>Disease duration, years</td>
<td>n.a.</td>
<td>8.0 (5.1)</td>
<td>65.1</td>
</tr>
</tbody>
</table>

Data presented as means with standard deviation in parentheses. n.a. = Not applicable.

1 Lower subject number due to missing data.

2 Score range 24–30 in patients.
ence in gray matter density only reached a subthreshold significance level of \( p < 0.001 \) (voxel level, uncorrected) while the difference in age was indeed statistically significant \( (p < 0.01) \).

**Discussion**

In this large study with 77 PD patients, we found significant PD-related reductions of gray matter density in predominantly the anterior temporal cortex, bilaterally, compared to healthy controls \( (p < 0.05, \text{FWE corrected, k of 8 voxels}) \). This distribution of regional atrophy was different from the observed aging-associated pattern of atrophy in temporal perisylvian and cingulate cortices. The latter was consistent with previous studies on normal aging \[15\]. Progression of atrophy in PD was inferred from the comparison between patients with the shortest and longest disease durations. This analysis indicated that the early anterior temporal atrophy did not show major progression, whereas atrophy of particularly the posterior parts of the lateral temporal cortex and adjacent ex-

**Fig. 1.** (a) Clusters of significant regional reduction of gray matter density in patients with PD when compared to control subjects \((p < 0.05, \text{FWE corrected, voxel extent k of 8 voxels})\). In addition to the brain volume images, the atrophy at the ventral surface of the temporal poles is depicted on a coronal slice at \( y +2 \) mm relative to the vertical traversing the anterior commissure. (b) The single regions with reduction of gray matter density in the 25 patients with PD with the shortest disease duration compared to the 87 healthy control subjects \((p < 0.001, \text{uncorrected, k of 8 voxels})\) in a coronal section \((y +2 \text{ mm})\). (c) Regional reduction of gray matter density in the 25 PD patients with the longest disease duration compared to healthy controls \((p < 0.05, \text{FWE corrected, k of 8 voxels})\). (d) Regional reduction of gray matter density of the 25 PD patients with the longest disease duration when compared to the 25 patients with the shortest disease duration \((\text{voxel level} p < 0.001, \text{uncorrected, k of 20 voxels})\). In addition, the atrophy of the right fusiform gyrus, right middle temporal gyrus and left inferior temporal gyrus are depicted on a coronal slice at \( y -44 \) mm relative to the vertical traversing the anterior commissure. Clusters are rendered onto the volume or a section of a standard T1-weighted brain (Montreal Neurological Institute). \( t \) values are indicated by the color bars; \( R = \text{right side of the brain} \).
At first sight, atrophy at the temporal pole may seem at odds with the dominant characteristics of motor dysfunction, which define PD. On the other hand, the consistency of symptoms such as hyposmia that precede motor symptoms may point at an impaired function of striate visual cortex as an early sign of PD would seem consistent with the previous description of Braak et al. [4].

Coordinates (standard brain Montreal Neurological Institute) refer to the voxels of maximum difference within clusters of significant difference (p < 0.05 FWE corrected, voxel extent of 8 voxels). BA = Brodmann area; s.c. = same cluster.

Table 2. Regional decreases in cerebral gray matter in patients with PD compared with controls

<table>
<thead>
<tr>
<th>Brain region</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>t value</th>
<th>Extent</th>
<th>p corrected</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Right hemisphere</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Temporal pole (BA 38)</td>
<td>32</td>
<td>14</td>
<td>−28</td>
<td>6.65</td>
<td>979</td>
<td>0.000</td>
</tr>
<tr>
<td>Inferior temporal gyrus (BA 20)</td>
<td>50</td>
<td>4</td>
<td>−42</td>
<td>6.52</td>
<td>s.c.</td>
<td>s.c.</td>
</tr>
<tr>
<td>Fusiform gyrus (BA 19)</td>
<td>26</td>
<td>−70</td>
<td>−14</td>
<td>5.47</td>
<td>18</td>
<td>0.012</td>
</tr>
<tr>
<td>Superior parietal lobule (BA 7)</td>
<td>8</td>
<td>−68</td>
<td>58</td>
<td>3.34</td>
<td>22</td>
<td>0.009</td>
</tr>
<tr>
<td>Rolandic operculum (BA 48)</td>
<td>46</td>
<td>−4</td>
<td>10</td>
<td>3.26</td>
<td>9</td>
<td>0.020</td>
</tr>
<tr>
<td>Dorsolateral prefrontal cortex (BA 46)</td>
<td>52</td>
<td>16</td>
<td>36</td>
<td>3.48</td>
<td>15</td>
<td>0.014</td>
</tr>
<tr>
<td><strong>Left hemisphere</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Temporal pole (BA 38)</td>
<td>−30</td>
<td>18</td>
<td>−32</td>
<td>5.40</td>
<td>253</td>
<td>0.000</td>
</tr>
<tr>
<td>Inferior frontal gyrus (BA 47)</td>
<td>−44</td>
<td>20</td>
<td>−2</td>
<td>5.39</td>
<td>s.c.</td>
<td>s.c.</td>
</tr>
<tr>
<td>Inferior temporal gyrus (BA 20)</td>
<td>−56</td>
<td>−6</td>
<td>−34</td>
<td>5.46</td>
<td>260</td>
<td>0.000</td>
</tr>
<tr>
<td>Hippocampus (BA 27)</td>
<td>−24</td>
<td>−10</td>
<td>−12</td>
<td>5.49</td>
<td>75</td>
<td>0.001</td>
</tr>
<tr>
<td>Fusiform gyrus (BA 19)</td>
<td>−28</td>
<td>−74</td>
<td>−16</td>
<td>5.78</td>
<td>81</td>
<td>0.001</td>
</tr>
<tr>
<td>Anterior lingual gyrus (BA 27)</td>
<td>−12</td>
<td>−36</td>
<td>−6</td>
<td>5.74</td>
<td>36</td>
<td>0.005</td>
</tr>
<tr>
<td>Extrastriate visual cortex (BA 19)</td>
<td>−50</td>
<td>−76</td>
<td>0</td>
<td>6.12</td>
<td>221</td>
<td>0.000</td>
</tr>
<tr>
<td>Superior occipital gyrus (BA 18)</td>
<td>−24</td>
<td>−92</td>
<td>26</td>
<td>5.44</td>
<td>49</td>
<td>0.003</td>
</tr>
<tr>
<td>Cuneus (BA 18)</td>
<td>−2</td>
<td>−84</td>
<td>34</td>
<td>5.21</td>
<td>31</td>
<td>0.006</td>
</tr>
</tbody>
</table>

Coordinates (standard brain Montreal Neurological Institute, MNI) refer to voxels of maximal difference within clusters of significant difference (p < 0.05 FWE corrected, voxel extent of 8 voxels). BA = Brodmann area; s.c. = same cluster.

Table 3. Progression of cerebral gray matter loss in PD (PD_short > PD_long)

<table>
<thead>
<tr>
<th>Brain region</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>t value</th>
<th>Extent</th>
<th>p uncorrected</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Right hemisphere</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Middle temporal gyrus (BA 21)</td>
<td>54</td>
<td>−42</td>
<td>−4</td>
<td>3.9</td>
<td>56</td>
<td>0.24</td>
</tr>
<tr>
<td>Fusiform gyrus (BA 19)</td>
<td>36</td>
<td>−40</td>
<td>−20</td>
<td>4.3</td>
<td>82</td>
<td>0.16</td>
</tr>
<tr>
<td>Supramarginal gyrus (BA 2)</td>
<td>62</td>
<td>−26</td>
<td>46</td>
<td>3.7</td>
<td>40</td>
<td>0.32</td>
</tr>
<tr>
<td>Middle frontal gyrus (BA 45)</td>
<td>60</td>
<td>32</td>
<td>16</td>
<td>3.8</td>
<td>64</td>
<td>0.21</td>
</tr>
<tr>
<td><strong>Left hemisphere</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inferior temporal gyrus</td>
<td>−54</td>
<td>−46</td>
<td>−14</td>
<td>3.5</td>
<td>32</td>
<td>0.38</td>
</tr>
<tr>
<td>Inferior temporal gyrus</td>
<td>−54</td>
<td>−58</td>
<td>−4</td>
<td>3.5</td>
<td>75</td>
<td>0.18</td>
</tr>
<tr>
<td>Middle temporal gyrus (BA 21)</td>
<td>−56</td>
<td>−56</td>
<td>20</td>
<td>3.3</td>
<td>21</td>
<td>0.48</td>
</tr>
<tr>
<td>Extrastriate visual cortex (BA 19)</td>
<td>−54</td>
<td>−72</td>
<td>4</td>
<td>3.5</td>
<td>28</td>
<td>0.41</td>
</tr>
</tbody>
</table>

Coordinates (standard brain Montreal Neurological Institute, MNI) refer to voxels of maximal difference in clusters surviving a threshold of p < 0.001 uncorrected with a voxel extent of 20 voxels. BA = Brodmann area.
temporal atrophy particularly spread to occipitotemporal and posterior parietal regions.

The temporal pole is implicated in a wide range of functions, particularly characterized by the attribution of an emotional content to stimuli in various perceptual domains [22]. Such emotion-related processing not only concerns its perception, but is intrinsically related to emotional expression, including simple vegetative responses such as changed heart rate or releasing tears, as well as more complex motor behavior [23, 24]. In this respect, functional coherence between limbic system structures such as anteromedial temporal regions, adjacent insula, orbitofrontal cortex and ventral striatum is logically subserving ‘emotion-based’ sensorimotor transformations [23]. The amygdala complex, embedded anteromedially in the temporal lobe, seems to play an important role in the integration of internal emotional states and actual stimulus conditions in guiding e.g. the ventral striatum in the onset of behavioral responses [23, 25]. A functional consequence of coherent temporal–striatum involvement in such circuitry is that its dysfunction, associated with temporal pole atrophy, may not only lead to olfactory deficit, but also to PD premotor symptoms such as depression, anxiety and autonomic impairment [26]. In this respect, a typical PD symptom such as hypomimia may not only be regarded as a manifestation of motor dysfunction, but may well reflect a more complex impairment of emotional processing, including emotion-motor transformation. The role of the temporal pole in emotion perception has been well described [22]. For instance, it supports our empathizing with other people [27], while bilateral removal of the temporal poles in monkeys leads to a disturbance in recognizing, interpreting and reacting to social signals [28].

It is intriguing to notice that the left-lateralized frontotemporal atrophy in the referred meta-analysis of Pan et al. [8] fits the observation that PD-related deficit in the uncinate fascicle particularly occurred in the left hemisphere [29]. Although Kim et al. [29] described more widely distributed white matter changes in the PD brain, changes in the crucial connection between the temporal pole and orbitofrontal cortex underscore the anterior temporal involvement in PD pathology.

A biochemical relation between anteromedial temporal dysfunction and PD might be inferred from the dopamine innervation in normal human subjects, as revealed by $^{18}$FDOPA positron emission tomography. While striatum uptake of this tracer is the most evident feature of this functional imaging method, $^{18}$F-DOPA uptake in limbic structures such as the entorhinal cortex and amygdala is more
than 4 times higher than that of the neocortex of the occipital lobe [30]. This implies that impaired dopamine innervation from the upper brainstem may have a larger impact on the referred limbic than neocortical regions. This argument, however, was not supported by the assessment of extrastriatal uptake of this tracer, as limbic uptake was not specifically reduced in early PD stages [31].

**Posterior Cortical Atrophy**

In addition to anterior temporal atrophy, regional clusters of PD-related atrophy particularly comprised posterior temporal, extrastriate visual and posterior parietal cortical regions. Moreover, the progression of atrophy during disease took place in predominantly these posterior regions, which suggests that it might initially follow the course of the inferior longitudinal fasciculus [32] or, more stepwise, the functional ventral visual pathway in reversed direction [33]. Such a progression of atrophy towards posterior temporal and extrastriate visual cortical regions would be consistent with a specific PD-related dysfunction in the visual domain. The latter includes hallucinations, which have been associated with impaired visual object processing in ventrolateral extrastriate visual areas [34]. The posterior parietal progression seems consistent with an important profile of neuro-psychological dysfunction seen early in the course of the disease, which may predict cognitive decline [35]. The association between functional impairment of both visual and parietal cortex regions in PD [36] and observations that visual stimuli may have a stronger impact on gait control in this disease [37, 38] underscore that more knowledge on visual cortex changes is required for understanding distinct PD symptoms.

**Atrophy in Motor-Related Areas**

While prefrontal and parietal foci of atrophy were found in our study, it may seem quite remarkable that no atrophy was seen in the striatum and cortical regions more specifically involved in motor processing. In a network view of cerebral processing, one might logically expect that degeneration of a crucial node or hub, such as the substantia nigra pars compacta, would inflict changes in distant parts of such a network (e.g. the cortico/basal ganglia/thalamocortical loops). This view is supported by the distribution of decreased metabolism including the premotor cortex in patients with PD [36]. One may, in this respect, speculate that atrophy in these regions may occur at more advanced stages of PD following a preceding decrease in metabolism.

**Conclusion**

VBM in a large group of PD patients showed that cortical atrophy was most pronounced at the temporal poles and the occipitotemporal junction, while it spread toward the lateral temporal cortex and adjacent extrastriate cortex in advanced PD. This distribution of cortical pathology highlights the presence of ‘premotor’ symptoms in PD.

**Acknowledgments**

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**References**


