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Neural Correlates of Distinct Cognitive Phenotypes in Early Parkinson's Disease

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Abstract

Objective: Cognitive decline is common in Parkinson's disease (PD), but changes can occur in a variety of cognitive domains. The lack of a single cognitive phenotype complicates diagnosis and tracking. In an earlier study we used a data-driven approach to identify distinct cognitive phenotypes of early PD. Here we identify the morphometric brain differences between those different phenotypes compared with cognitively normal PD participants.

Methods: Six different cognitive classes were included (Weak, Typical, Weak-Visuospatial/ Strong-Memory, Weak-Visuospatial, Amnestic, Strong). Structural differences between each class and the Typical class were assessed by deformation-based morphometry.

Results: The different groups evidenced different patterns of atrophy. Weak class had frontotemporal and insular atrophy; Weak-Visuospatial/Strong-Memory class had frontotemporal, insular, parietal, and putamen atrophy; Weak-Visuospatial class had Rolandic operculum; Amnestic class had left frontotemporal, occipital, parietal and insular atrophy when compared to the Typical class. The Strong class did not have any atrophy but had significant differences in left temporal cortex in comparison to the Typical class.

Conclusions: Structural neuroimaging differences are evident in PD patients with distinct cognitive phenotypes even very early in the disease process prior to the emergence of frank cognitive impairment. Future studies will elucidate whether these have prognostic value in identifying trajectories toward dementia, or if they represent groups sensitive to different treatments.

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Keywords

Cognitive dysfunction; Cluster analysis; Parkinson's disease; Neuroimaging; Magnetic resonance imaging; atrophy

1. Introduction

Parkinson's disease (PD) is a neurodegenerative disorder characterized by motor and nonmotor impairments. Cognitive impairments are quite heterogeneous and can be observed during all stages of the disease, including the prodromal phase. Executive, visuospatial, and memory deficits are common, [1] and up to 80% of patients develop dementia [2]. However, the underlying neuroanatomical mechanisms of cognitive deficits in PD have yet to be clarified, especially early in the disease course when cognitive changes, if present, are subtle. Establishing underlying structural changes of cognitive impairments may be important for the early detection and management of these impairments, as well as prognosis.

Several cortical and subcortical regions have been implicated in cognitive impairments in PD. PD patients with dementia (PDD) and mild cognitive impairment (PD-MCI) have frontal, temporal, and parietal atrophy [3–7]. Atrophy to subcortical structures including the thalamus, putamen, hippocampus, amygdala, and caudate have also been reported [3,8–11]. However, results vary greatly in the studies of PD-MCI possibly due to the heterogeneity of PD-MCI, or specific to how MCI was defined and assessed in each study. For example, one study reported atrophy in only subcortical regions [8], while another study showed no structural differences in PD-MCI compared to healthy controls [12]. Another study demonstrated insular, frontal, and temporal atrophy only in the left hemisphere in PD-MCI compared to cognitive domains suggest that changes in distinct regions are involved in each domain in PD. Memory performance was related to medial temporal lobe structure volumes, whereas visuospatial deficits have been attributed to superior temporal, parietal, and occipital volume loss [14–17]. Further classification of PD-MCI based on affected cognitive domains may provide more consistent results in future studies.

These structural changes can also present early on in the disease alongside cognitive deficits. In early PD, memory deficits have been associated with cortical thinning in occipital, superior parietal and postcentral gyri [18]. In a study focusing on de novo PD, PD-MCI group had verbal memory deficits and temporal lobe, including the hippocampus; and parietal lobe with the precuneus and posterior cingulate gyrus atrophy compared to cognitively intact PD patients [19]. More interestingly, these changes can be seen even without pronounced cognitive deficits in PD. In a sample of generally cognitively intact PD patients, weaknesses in the cognitive profile were associated with distinct anatomical patterns: deficits in executive functioning were associated with decreased volumes within regions of the frontostriatal cognitive-control network, lower memory scores were associated with decreased volume in frontostriatal cognitive-control network regions and the medial

temporal lobe, and worse visuospatial performance was associated with temporo-occipital volume reduction [14].

In a previous study, we used latent class analysis (LCA) to identify different cognitive phenotypes in early PD [20]. This technique is purely data-driven, not requiring a priori hypotheses, and extracts classes based on patterns of cognitive scores between patients. Based on LCA, several classes emerged including the cognitively strong, those with isolated memory, and visuospatial weaknesses. Demographics and disease characteristics were evaluated only after the classes were created. Cognitive performance was independent of motor and non-motor impairments, cerebrospinal fluid (CSF) biomarkers, and striatal dopamine transporter (DaT) binding ratios. In this study, we investigated the differences in structural integrity between these classes to determine underlying neuroanatomical alterations of PD patients with different cognitive profiles. Due to multiple networks and structures being implicated in cognitive impairment in PD [21,22], instead of regional volume analysis, a whole brain analysis approach without a priori determined regions of interest was implemented. We utilized deformation based morphometry (DBM), as this technique preserves the entirety of the imaging data and has a higher sensitivity for subcortical atrophy, increasing the likelihood of detecting early changes in de novo PD [23].

2. Material and Methods

2.1 Participants

122 de novo, medication-naïve PD subjects with 3 T baseline MRI from the PPMI database were included. Details of the aims and design of the PPMI have been published elsewhere [24] and can be found on the PPMI website (http://www.ppmi-info.org/study-design). The study was approved by the institutional review board at each site and all participants provided written informed consent. The research was completed in accordance with Helsinki Declaration.

2.2 Cognitive measures and classes

Classes reported in our previous study were used [20]. Cognitive tests included Judgment of Line Orientation (JLO), learning and recall scores of the Hopkins Verbal Learning Test-Revised (HVLT-R), Letter Number Sequencing (LNS), Symbol-Digit Modalities Test (SDMT), and semantic fluency with three prompts (animals, vegetables and fruits). Raw scores were Z-transformed against the mean of all cases. These scores served as cluster indicators. Clusters of variables were identified and labeled based on the characteristics of the variables within each cluster. Then clusters were compared to each other. The solution with the lowest Bayesian Information Criterion statistic, good class stability, and unique performance patterns across classes was selected as the best fit solution. This resulted in six cognitive classes in total; the Weak-Overall, Typical-Overall, Strong-Overall, Strong-Memory, Weak-Visuospatial, and Amnestic. The Weak-Overall class had average, and the Strong-Overall class demonstrated strong cognition. The Strong-Memory class performed above average in the memory tests, and had poor visuospatial functioning. The Weak-

We relabeled the Weak-Overall class as Weak, Typical-Overall as Typical, and Strong-Overall as Strong for ease in this study. To emphasize the visuospatial weakness in the Strong-Memory class, this class was relabeled as Weak-Visuospatial/Strong Memory class. The other class labels were not changed. 7 subjects in the Weak class, 30 in the Typical class, 9 in the Weak-Visuospatial/Strong-Memory, 21 in the Weak-Visuospatial, 26 in the Amnestic, 29 in the Strong class had available imaging data and were included. Although class identification was independent of norms, we further investigated the frequency of MCI for each class. Due to the limited number of cognitive tests included in the PPMI, Movement Disorders Society Task Force Level I criteria was used to screen for MCI [25]. Standard scores were used to determine MCI or high functioning. None of the participants had scores 1.5 standard deviation higher than the normal mean across all tests. However, as individuals in the Strong class performed above normal levels in some of tests (13.8% above norms in semantic fluency, 3.4% in SDMT, 34.5% in JLO, 24.1% in LNS, 20.7% in HVLT-R Learning, 31% in HVLT-R Recall), the Typical class was used as the reference group.

Disease characteristics were assessed by the Movement Disorders Society-Unified Parkinson's Disease Rating Scale. Disease severity was determined by Hoehn and Yahr staging (HYS). Modified Schwab and England Activity of Daily Life (ADL) was used to assess patients' independency level. Non-motor scales included the Geriatric Depression Scale (GDS), State-Trait Anxiety Inventory (STAI), REM Sleep Behavior Disorder Screening Questionnaire (RBDSQ), Epworth Sleepiness Scale (ESS), University of Pennsylvania Smell Identification Test (UPSIT) and the Scales for Outcomes in Parkinson's Disease-Autonomic (SCOPA-AUT).

Apolipoprotein E e4 (APOEe4) status was included for subjects with available genotyping. CSF biomarkers included amyloid- β , α -synuclein, phosphorylated-tau (p-tau), and total tau. Striatal DaT binding ratios were derived for the left and right caudate and putamen by single photon emission CT (SPECT) imaging with [1231]FP-CIT radioligand. Binding was calculated by a standardized volume of interest template for each region compared to an occipital reference region.

2.3 MRI data acquisition

All imaging data were acquired from the PPMI database. High resolution T1-weighted MRI structural images were collected for the subjects across 5 different scanners. 15 subjects were scanned on a GE Signa 3.0T HDx scanner using the 3D fast SPGR sequence (8 subjects with a voxel size of $1 \times 1 \times 1.2 \text{ mm}^3$, matrix size: 256×256 , 152 sagittal slices; 7 subjects with a voxel size of $1 \times 1 \times 0.7 \text{ mm}^3$, matrix size: 256×256 , 248 sagittal slices). 3 subjects were scanned on a GE 3.0T Discovery scanner using the fast SPGR sequence with a voxel size of $1 \times 1 \times 1.2 \text{ mm}^3$, matrix size: 256×256 , 248 sagittal slices). 3 subjects were scanned on a GE 3.0T Discovery scanner using the fast SPGR sequence with a voxel size of $1 \times 1 \times 1.2 \text{ mm}^3$, matrix size: 256×256 and 152 sagittal slices. 8 subjects were scanned on a Philips Achieva 3.0T scanner using the MPRAGE sequence with a voxel size of $0.92 \times 0.92 \times 1.2 \text{ mm}^3$, matrix size: 288×288 and 152 sagittal slices (1 subject acquired 152 axial slices). 6 subjects were scanned on a SIEMENS 3.0T Verio scanner using the MPRAGE sequence with a voxel size of $1 \times 1 \times 1 \times 1 \text{ mm}^3$, matrix size: 240×256 , 176 sagittal

slices. 90 subjects were scanned on a Siemens 3.0T Trio scanner using the MPRAGE sequence (88 subjects with a voxel size of $1 \times 1 \times 1 \text{ mm}^3$, matrix size 240×256 , 176 sagittal slices; 1 subject with a voxel size of $0.98 \times 0.98 \times 1 \text{ mm}^3$, matrix size 256×256 , 160 sagittal slices and 1 subject with a voxel size of $1 \times 1 \times 1 \text{ mm}^3$, matrix size 240×256 , 196 sagittal slices). All T1 images were acquired with the following imaging parameters: TR, 2300 ms; TE, 2.98 ms; flip angle, 9°; and TI, 900 ms.

2.3.1 Deformation based morphometry (DBM)—DBM analysis was performed

using the computational anatomy toolbox (CAT) for SPM (http://www.neuro.uni-jena.de/ cat/). T1 images from 122 subjects were first re-sliced to a uniform voxel size of $1 \times 1 \times 1$ mm³ and then preprocessed with signal inhomogeneity correction and global intensity normalization. The preprocessed T1 image of each subject was then non-linearly normalized to the standard MNI-152 1.5 mm template to minimize the regional differences due to local deformations. The type and localization of the structural differences between the subject brain and the template brain were minimized in this non-linear normalization step and coded in the deformation field. The retrospective quality control framework in CAT were performed and all images were scored C- or above, lying in or above the satisfactory category. Percentage change of each voxel volume was finally quantified by the Jacobian determinant of the deformation field [26]. The deformation field map from each subject was further smoothed using an 8mm Gaussian kernel before the statistical analysis.

2.4 Statistical analysis

Statistical analyses of demographics, disease characteristics and test scores were completed using SPSS V 23.0. Numerical data are reported as the mean (standard deviation) and categorical data as percentage unless stated otherwise. Since not all of the participants from our previous study were included, all variables were reassessed across groups. Demographics, disease characteristics, and test scores across groups were compared by analysis of variance (ANOVA) with Dunnett's post hoc test (Typical class as the reference) and chi-square tests. p <0.05 was considered statistically significant.

Statistical comparisons for brain atrophy were performed on Jacobian determinants of the deformation field between each cognitive class and the Typical class. The Typical class was chosen for comparison as this group had an average cognitive performance across domains. Two-sample t-tests were performed with age, gender, year of education, scanner type, and handedness as covariates. Statistical significance level was established at family-wise corrected p-value of 0.05 (cluster-wise corrected, |t|>2.4 with at least 1589 voxels in the cluster). Covariates were regressed out from the data to determine the effect sizes of regional volume differences. Cohen's d was calculated by dividing the difference between the means of two groups by the pooled standard.

Using the same analysis approach, comparisons between Weak-Visuospatial/Strong-Memory and Amnestic class were done to investigate the structural associates of memory performance (Supplementary Results).

3. Results

3.1 Demographics, biomarkers and disease characteristics

Demographics and disease characteristics are summarized in Table 1. Non-motor symptoms and biomarkers are shown in Supplementary Table 1. As all participants were de novo PD, disease severity only ranged between HYS 1 and 2. There were group differences in MDS-UPDRS Part I, RBDSQ, SCOPAAUT and MoCA scores. Weak class had higher MDS-UPDRS Part I (p=.039), RBDSQ (p=.019), and SCOPAAUT (p=.024) scores compared to the Typical class. None of the cognitive classes differed from the Typical class for the MoCA score.

3.2 Cognitive classes

Raw scores on cognitive tests are given in Table 2. All tests were different across the cognitive classes. Comparisons with Typical class showed that Weak class had lower scores on all tests (p=.000 for all tests) except for semantic fluency. Weak-Visuospatial/Strong-Memory class had lower score on JLO and higher scores in HVLT-R learning and recall compared to the Typical class (p=.000 for all). Weak-Visuospatial class had lower score in JLO compared to the Typical class (p=.000). Amnestic class had lower scores in HVLT-R learning and recall compared to the Typical class (p=.000). Strong class had higher scores in semantic fluency (p=.035), LNS (p=.007), HVLT-R learning and recall (p=.000 for both) compared to the Typical class.

3.3 Deformation based morphometry (DBM) results

Coordinates of the significantly different clusters in each cognitive class compared to the Typical class are summarized in Table 3. Weak class had the most extensive atrophy including regions of left frontotemporal, insular; and right frontal lobes with a very large mean effect size [Cohen's d=1.24 (0.22)] (Figure 1a). Weak-Visuospatial/Strong-Memory class had left frontotemporal, insular, parietal; right frontal and putamen atrophy with a large mean effect size [Cohen's d=1.00 (0.15)] (Figure 1b). Weak-Visuospatial class only had atrophy in the left Rolandic operculum with a large mean effect size [Cohen's d=0.81 (0.11)] (Figure 1c). Amnestic class had left inferior temporal, occipital; right motor cortex, parietal and insular atrophy with a large mean effect size [Cohen's d=0.86 (0.37)] (Figure 1d). Compared to the Strong class, Typical class had atrophy in left middle and inferior temporal gyri with a medium-large mean effect size [Cohen's d=-0.66 (0.06)] (Figure 1e). The only overlap between Weak-Visuospatial/Strong Memory and Weak-Visuospatial class when compared to the Typical class, was the atrophy in left precentral gyrus (Figure 2 illustrates the overlap, not the statistical contrast).

4. Discussion

The current study sought to investigate the structural neuroanatomic correlates of different cognitive profiles in recently diagnosed, nondemented PD patients. We compared groups of PD patients with data-driven patterns of cognitive weakness to a comparison group of PD patients with average scores on cognitive tests. We were able to demonstrate subtle structural differences in these classes which might be important in understanding the earliest cognitive

disruption in PD. As our main concern was to investigate neural correlates of different types of cognitive weakness in PD, but not PD itself as a disorder, we chose not to include a healthy control group. By using a group of PD patients with average cognitive performance as a control group, we aimed to control for the effects of PD (neural correlates of motor or other non-motor changes) and focus solely on cognition. Additionally, we aimed to investigate whether neural correlates of cognitive weakness can be found early in the cognitive impairment process, thus we only included nondemented PD patients.

The Weak class had both the most extensive set of cognitive weaknesses and the most widespread pattern of atrophy. They demonstrated worse executive and visuospatial function, learning and memory scores than the Typical class. The analysis demonstrated atrophy in the lateral and inferior temporal regions, posteromedial and lateral frontal regions, insula and motor cortex in the Weak class. Lateral and inferior temporal regions are associated with semantic processing [27,28]. Posteromedial and lateral frontal regions are suggested to work together for attention and error monitoring [29]. Insula has been associated with sensorimotor processing and general cognition [30]. Lateral temporal and frontal regions, as well as the insula have also been associated with visuospatial functioning using lesion-symptom mapping in ischemic stroke patients [31]. In a previous study volumetric reduction in insular, superior frontal, and lateral temporal grey matter was observed in PD-MCI compared to PD with no cognitive impairment [13]. In this study, insular atrophy was associated with executive function and attention; lateral temporal atrophy was associated with executive function, attention and memory, superior frontal atrophy was associated with attention impairment. Our findings suggest that even in a sample of PD patients who have subtle cognitive weakness as opposed to the frank impairments seen in MCI, structural patterns observed in PD-MCI may be present. While interpreting the findings within this class, it is also important to consider that the sample size was limited to only seven PD patients in this cognitive class.

The Weak-Visuospatial/Strong-Memory class had worse visuospatial functioning coupled with better learning and memory performance compared to the Typical class. Compared to the Typical class, atrophy in prefrontal, lateral temporal, parietal, insular, motor cortex; and subcortical regions was found. Previous studies indicate that frontal, lateral temporal, insula, and basal ganglia volumetry is associated with visuospatial decline [31]. The involvement of putamen in visuospatial function was shown in nondemented PD patients with visuospatial weakness in a previous study [32]. As regions with volumetric reductions we found in this class are in line with previous studies on visuospatial functioning associations of regional grey matter volume, the structural differences between this class and the Typical class are likely to be driven by visuospatial weakness rather than learning and memory strength. However, the small number of patients in this class make it hard to interpret the results.

The only differences between Weak-Visuospatial class and the Typical class were weakness in visuospatial functioning, and atrophy in left rolandic operculum including the precentral gyrus and the insula. In PD, atrophy of the Rolandic operculum has been seen in patients with mild to moderate diseases [33]. In particular, left insular atrophy has been shown to correlate with visuospatial performance in PD-MCI [34]. Our findings indicate that the atrophy pattern in PD-MCI associated with visuospatial functioning present itself early on in

the disease without any significant visuospatial deficits, but only weaknesses. In addition to volumetry, structural connectivity of insula was also implicated in a visuospatial attention task [35]. Thus, insula has the potential to serve as a marker for future visuospatial decline. More interestingly, this class had less atrophy than Weak-Visuospatial/Strong Memory class when both classes were compared to the Typical class. This suggests that the pathophysiology of the cognitive weaknesses in these two groups are different. The overlap between the regions with atrophy in these two classes may be responsible for the visuospatial weakness seen in both classes. Traditionally known for its involvement in movement control, precentral gyrus also plays a role in motor imagery and spatial transformations [36]. Working memory assessed with recalling sensory materials and movement sequences after a time delay has also been associated with precentral gyrus activation [37]. Precentral gyrus atrophy is also observed in PD-MCI [38], and progression to MCI in these classes may be associated with atrophy in this region. A longitudinal study focusing on the progression of cognitive weaknesses to deficits in these classes can help clarify this hypothesis.

The Amnestic class significantly differed from the Typical class on learning and memory. Previous studies in early PD reported that atrophy in the medial temporal lobe structures correlate with memory functioning similar to Alzheimer's disease [11,15]. Lateral temporal, parietal, occipital, insular and motor cortex atrophy in this class is in line with the atrophy pattern in PD-MCI [13]. This class is likely to have a higher rate of conversion to MCI, as alongside bradykinesia, years of education, subjective cognitive decline report, REM sleep behavior disorder; verbal memory retention is the only cognitive predictor of future cognitive decline in de novo PD patients [39]. Our results reveal that PD-MCI atrophy patterns are present in patients with weaknesses in only verbal learning and memory. The overlap between atrophic regions in previously reported PD-MCI and the Amnestic class suggests that this cognitive class alongside the Weak class are the two classes most likely to progress to PD-MCI. This should be evaluated longitudinally by characterizing PD patients without any cognitive impairment but rather weaknesses in learning and memory or overall cognitive performance during the baseline. These patients may have a higher risk of developing cognitive impairment more rapidly.

The Strong class performed better on executive function, learning and memory tests; and did not show any atrophy patterns compared to the Typical class. Compared to the Strong class, the Typical class had atrophy in left middle and inferior temporal gyri. These gyri have been associated with several cognitive domains including language, semantic memory, and visual perception [27]. Thus, atrophy in these regions, seen in the Typical class, might lead to difficulty in encoding and then recalling a list of words from certain semantic categories which may easily elicit visual representations in the normal brain. Additionally, structural differences were limited to the left hemisphere. The left hemisphere is more involved with language and motor control in both right-handed and left-handed individuals [40]. Therefore classes identified by scores on both verbal and nonverbal measures may provide different results and should be further investigated in future studies.

One of the major limitations in our study was the restricted number of cognitive tests which were used to determine the cognitive profiles. PPMI is a multi-center study aiming at

establishing biomarkers for PD, which includes data for multiple features related to PD. While the accessibility of this dataset provides an opportunity to conduct research on large numbers of patients in a readily available dataset, the number of cognitive tests is restricted within each cognitive domain. This potentially limits the reliability of the cognitive profiles included in our study. For example, the Movement Disorders Society Level II criteria for MCI requires at least two independent tests for each domain [25]. Therefore future studies would benefit from including more cognitive tests evaluating each domain with at least two independent tests to establish cognitive classes. Our imaging findings should also be interpreted in the light of this limitation, which limit the reliability of the structural findings. Another limitation is the number of patients included in the groups. There were only seven patients in the Weak class. Although this low number reduces the reliability of the findings within this class, high effect sizes (|d|>0.8) suggest that the results may reflect a true finding rather than an artefact. In conclusion, our findings indicate that different, data-driven, cognitive profiles in PD are related to specific structural patterns. These patterns can be observed within the first two years of the disease, and before any treatment has been administered. Future studies should include a more comprehensive neurocognitive battery to assess each cognitive domain extensively, using both verbal and nonverbal measures in larger patient samples. Increasing the statistical power may identify a stronger correlation between the cognitive profiles and associated structural changes. Additionally, studies focusing on structural and functional connectivity may help understand the role of specific networks involving these regions in cognitive impairments in PD as the symptoms of this disease are due to disrupted networks rather than atrophy in single regions. Prospective validation of the correlation between the cognitive profiles and underlying structural and functional changes in de-novo PD patients should be performed longitudinally as well. Due to the cross-sectional nature of our study, we cannot know whether these atrophy patterns associated with cognitive classes are independent of PD progression and solely reflect cognitive weaknesses. Future studies evaluating the PD patients with different cognitive phenotypes in a longitudinal manner, can reveal whether specific cognitive deficits are due to the baseline structural differences.

5. Conclusions

This study shows that cognitive weaknesses in de novo PD without dementia have distinct atrophy patterns. The atrophy pattern in each cognitive class is not due to the PD but rather to cognitive performance. Latent class analysis determining cognitive class can help establish imaging biomarkers in early PD patients.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

• Distinct atrophy patterns are associated with different cognitive classes.

- Atrophy can be found before the development of cognitive decline.
- Longitudinal studies can determine whether these atrophy patterns affect cognition



Figure 1.

Local clusters of atrophy in the a) Weak class, b) Weak-Visuospatial/Strong-Memory class, c) Weak-Visuospatial class, d) Amnestic class, e) Strong class. All comparisons were done against the Typical class. Color bar indicates the amount of difference between the two classes, a greater t value indicates a larger difference.



Figure 2.

Overlap of atrophy between Weak-Visuospatial/Strong Memory and the Weak-Visuospatial class. Red color indicates the atrophy in the Weak-Visuospatial/Strong-Memory class compared to the Typical class; green color indicates the atrophy in the Weak-Visuospatial class compared to the Typical class; yellow color indicates the atrophy overlap between these two classes.

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Demographics and disease characteristics

	Weak (n=7)	Typical (n=30)	Weak-Visuospatial/Strong-Memory (n=9)	Weak-Visuospatial (n=21)	Amnestic (n=26)	Strong (n=29)	Statistical comparison
Age	66.29 (7.65)	65.43 (4.78)	59.89 (9.2)	63.43 (8.15)	64.81 (6.74)	61.14 (7.12)	F(5, 116)=2.039, p=.078
Gender, male (%)	100	76.7	44.4	66.7	80.8	69	$\chi^{2}(5)=7.971$, p=.158
Years of education	13 (3.32)	14.7 (2.22)	15 (1.87)	15.24 (2.59)	14.73 (2.95)	15.9 (2.5)	F(5,110)=1.506, p=.194
Handedness, right (%)	71.4	93.3	77.8	85.7	88.5	86.2	$\chi^{2}(5)=13.229, p=.211$
Family history of PD (%)	28.6	16.7	11.1	19	30.8	17.2	$\chi^{2}(5)=2.937$, p=.710
APOEe4 carrier (%)	28.6	26.7	44.4	28.6	26.9	6.9	$\chi^{2}(5)=10.420, p=.404$
Disease duration, months	7.17 (8.28)	4.86 (4.74)	5.25 (6.23)	7.81 (6.96)	9.27 (8.19)	5.42 (7.42)	F(5,110)=1.487, p=.200
Dominantly affected side, right (%)	14.3	33.3	22.2	52.4	38.5	41.4	χ ² (5)=11.177, p=.344
HYS, 1 (%)	33.3	44.8	37.5	28.6	23.1	61.5	$\chi^{2}(5)=9.707$, p=.084
MDS-UPDRS total score	39.83 (16.87)	30.72 (13.76)	29.88 (11)	35.38 (14.51)	34.08 (12.1)	28.35 (11.86)	F(5, 110)=1.330, p=.257
MDS-UPDRS Part I	2.67 (2.16)	0.93 (1.07)	1.88 (1.81)	1.05 (1.2)	1.58 (1.98)	0.77 (0.99)	F(5,100)=2.643, p=.027*
MDS-UPDRS Part I Patient Questionnaire	5 (4.24)	4 (3.93)	3.88 (3.09)	3.95 (2.82)	5.08 (3.15)	4.5 (2.92)	F(5,110)=.462, p=.804
MDS-UPDRS Part II	6.33 (3.56)	4.69 (3.89)	5.5 (4)	5.91 (4.17)	5.65 (4.05)	5.65 (3.42)	F(5,110)=.462, p=.867
MDS-UPDRS Part III	25.83 (10.94)	21.1 (9.32)	18.63 (5.76)	24.48 (8.89)	21.77 (8.8)	17.42 (7.85)	F(5,110)=2.079, p=.073
MoCA	26.29 (2.5)	27.47 (1.96)	28.44 (1.59)	26.48 (2.54)	26 (2.62)	27.59 (2.04)	F(5,116)=2.788, p=.020*
PD: Parkinson's Disease Rat	ing Scale: MoC∆	· Montreal Cognitiv	e Assessment				

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All variables are stated as mean (standard deviation) or percentage where applicable.

Statistical significance is marked with *.

	Weak (n=7)	Typical (n=30)	Weak-Visuospatial/Strong-Memory (n=9)	Weak-Visuospatial (n=21)	Amnestic (n=26)	Strong (n=29)	Statistical comparison
Semantic fluency	12.91 (3.12)	15.40 (2.95)	17.07 (2.42)	14.63 (3.24)	13.47 (3.17)	17.71 (3.90)	F(5,116)=6.20, p=.000*
SDMT	24.86 (6.57)	41.03 (8.74)	48.44 (11.78)	36.71 (8.59)	35.73 (6.5)	44.14 (6.93)	F(5,116)=10.729, p=.000*
JLO	19.43 (3.60)	27.27 (2.2)	20.67 (1.41)	19.33 (2.39)	25.69 (3.03)	28.07 (1.65)	F(5,116)=53.111, p=.000*
TNS	4.71 (1.38)	9.63 (1.71)	10.56 (3.64)	9.76 (2.14)	10.08 (2.43)	11.62 (2.65)	F(5,116)=10.160, p=.000*
HVLT-R-Learning	17.86 (3.58)	23.1 (2.31)	31 (3.39)	24.05 (2.25)	16.73 (3.40)	29.76 (1.81)	F(5,116)=88.157, p=.000*
HVLT-R-Recall	4 (2.38)	8.03 (1.25)	11.11 (0.93)	7.76 (1.81)	4.54 (1.77)	10.79 (1.08)	F(5,116)=65.017, p=.000*
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Orientation; LNN: Line Number Sequencing; HVLI-R: Hopkins Verbal Learning Test-Revised. Lle SDMT: Symbol-Digit Modalities Test; JLO: Judgment of

All raw scores are stated as mean (standard deviation).

Statistical significance is marked with *. Cells highlighted in bold show significant difference between that cognitive class and the Typical class.

Table 3

Significant clusters of atrophy in each class compared to the Typical class

			MNI coordinates				
Region	Extent (Number of voxels)	t-value	x	у	z		
Weak							
L Middle Temporal Gyrus	10030	4.778	-48	2	-19		
L Inferior Temporal Gyrus	10030	4.498	-44	-39	-16		
R Precentral Gyrus	4366	4.393	42	-18	41		
R Posteromedial Frontal Cortex	4366	4.104	6	5	59		
R Inferior Temporal Gyrus	3063	3.619	39	-9	-34		
L Rolandic Operculum	1647	3.561	-45	-22	23		
L Insular Lobe	1647	3.245	-35	-6	15		
Weak Visuospatial/Strong-Memory							
L Superior Orbital Gyrus	3676	4.683	-12	62	-9		
L Middle Frontal Gyrus	3676	4.501	-33	41	38		
R Middle Frontal Gyrus	2934	4.549	33	48	35		
R Superior Frontal Gyrus	2934	2.735	25	50	8		
R Putamen	2286	4.482	34	-9	11		
R Precentral Gyrus	2286	3.073	54	0	26		
L Inferior Frontal Gyrus (pars opercularis)	4122	4.249	-56	11	38		
L Insular Lobe	4122	4.001	-33	-6	15		
L Middle Frontal Gyrus	4122	3.408	-45	12	56		
LSupramarginal Gyrus	1890	3.360	-54	-31	39		
L Inferior Temporal Gyrus	1890	2.783	-63	-42	-12		
Weak-Visuospatial							
L Rolandic Operculum	2458	4.520	-48	-6	17		
Amnestic							
L Inferior Temporal Gyrus	3277	4.955	-45	-3	-37		
L Lingual Gyrus	1681	3.866	-6	-79	6		
R Supramarginal Gyrus	7944	3.788	61	-31	29		
R Precentral Gyrus	7944	3.743	58	0	36		
R Insular Lobe	7944	3.666	37	-6	21		
	Strong *						
L Middle Temporal Gyrus	2548	-3.735	-50	-72	18		
L Inferior Temporal Gyrus	2548	-3.393	-56	-61	-7		

L: left; R: right.

 * Atrophy was found in the Typical class when compared to the Strong class.