Regional Gray Matter Atrophy in Patients with Parkinson Disease and Freezing of Gait

BACKGROUND AND PURPOSE: FOG is a troublesome symptom of PD. Despite growing evidence suggesting that FOG in PD may be associated with cognitive dysfunction, the relationship between regional brain atrophy and FOG has been poorly investigated.

MATERIALS AND METHODS: Optimized VBM was applied to 3T brain MR images of 24 patients with PD and 12 HC. Patients were classified as either FOG− or FOG+ (n = 12) based on their responses to a validated FOG Questionnaire and clinical observation. All patients with PD also underwent a detailed neuropsychological evaluation.

RESULTS: The VBM analysis in patients with FOG+ showed a reduced GM volume in the left cuneus, precuneus, lingual gyrus, and posterior cingulate cortex compared with both patients with FOG− and HC. We did not detect any significant change of GM volume when comparing HC versus all patients with PD (FOG− and FOG+). FOG clinical severity was significantly correlated with GM loss in posterior cortical regions. Finally, patients with FOG+ scored lower on tests of frontal lobe function.

CONCLUSIONS: Our findings provide the first evidence that the development of FOG in patients with PD is associated with posterior GM atrophy, which may play a role in the complex pathophysiology of this disabling symptom.

ABBREVIATIONS: 18F = fluorine 18; FOG = freezing of gait; FOG-Q = FOG Questionnaire; FWE = family-wise error; GM = gray matter; HC = healthy controls; PD = Parkinson disease; RCPM = Raven’s Colored Progressive Matrices; UPDRS = Unified Parkinson’s Disease Rating Scale; VBM = voxel-based morphometry

FOG is a common and very disabling symptom of PD in which patients experience an episodic inability to generate effective stepping despite the intention to walk. Recent studies identified FOG as 1 of the most relevant contributing factors to the worsening quality of life among patients with PD. Although typically occurring late in the progression of PD, FOG can be experienced in relatively earlier stages and has been reported in up to 26% of patients never treated with levodopa. FOG has been suggested as an independent cardinal sign of PD not correlated with rigidity or bradykinesia but associated with gait, balance, and speech dysfunctions. In the deprenyl and tocopherol antioxidative therapy of Parkinsonism cohort, early occurrence of FOG was associated with cognitive decline, and recent neuropsychological evidence has demonstrated a relation between cognitive dysfunction and FOG in patients with PD.

The mechanisms underlying FOG are not yet fully understood. Nonconventional imaging studies have yet to investigate the neural correlates of FOG showing an involvement of both posterior cortical and subcortical areas. However, to our knowledge, only 1 study has investigated the pattern of brain atrophy in patients with FOG, demonstrating reduced GM volume in the brain stem without differences in cortical areas between patients and HC. Hence, the aim of the present study was to further explore the differences in local GM volume in patients with PD with and without FOG by using VBM.

Materials and Methods

Patient Population

We investigated 24 patients (7 women and 17 men) with a diagnosis of PD according to the clinical diagnostic criteria of the UK Parkinson’s Disease Society Brain Bank. Inclusion criteria were the following: 1) age of 45 years or older, 2) a Hoehn and Yahr stage ≤2.5 while in “on medication state,” 3) disease duration <10 years, and 4) anti-parkinsonian treatment at a stable and optimized daily dosage during the 4 weeks before study entry. Exclusion criteria were the following: 1) dementia according to clinical diagnostic criteria for dementia associated with PD; 2) major depression according to the criteria of the Diagnostic and Statistical Manual of Mental Disorders-IV for current major depression; 3) clinically significant comorbidities, including serious cardiovascular or cerebrovascular disease; and 4) anticholinergic or neuroleptic treatment. Patients were classified as exhibiting FOG (FOG+) based on the following 2 conditions that both had to be fulfilled: 1) score >0 to item 3 of the FOG-Q, and 2) patient recognition of this condition when it was demonstrated to them by an experienced clinician miming the phenomenon. Patients not fulfilling either of the above conditions were classified as FOG−. A group of 12 age- and sex-matched HC with no previous histories of neurologic or psychiatric diseases and with normal brain MRI findings was also recruited to assess FOG-related effects in relation to the
healthy population. All participants gave written informed consent, which was approved by the local ethics committee according to the Declaration of Helsinki.

**Clinical, Motor, and Neuropsychological Assessment Analysis**

All patients were evaluated 60–90 minutes after their morning dose of levodopa and underwent a detailed clinical and cognitive evaluation.

We recorded demographic data and disease history, and administered parts I-IV of the UPDRS and the FOG-Q. To evaluate cognitive function, we administered the following: 1) Mini-Mental State Examination, 2) the spatial (Corsi Block-Tapping) and verbal span to test spatial and verbal memory, 3) Rey Auditory Verbal Learning Test (15-word) to assess immediate and delayed recall of word lists, 4) the constructional apraxia test, 5) the RCPM test to evaluate abstract nonverbal reasoning, 6) Frontal Assessment Battery, a brief battery aimed at assessing frontal lobe functions, 7) phonological verbal fluency to estimate mental flexibility, 8) the Stroop Test (part II and III) to assess both color vision and sensitivity to interference, 9) Ten-Point Clock Test to evaluate spatial programming, and 10) cancellation attentional matrices to explore attention domain. Test scores were corrected for current normative values. Motor and cognitive functions were each evaluated by 2 different raters. Tests were administered by a trained neuropsychologist blinded to presence/absence of FOG.

**Imaging Parameters**

All subjects were scanned in the morning, and each patient was in the “on-medication” state (ie, 60–90 minutes after their first dose of levodopa). MR images were acquired on a 3T scanner (GE Healthcare, Milwaukee, Wisconsin) equipped with an 8-channel parallel head coil. Structural MR imaging data were acquired by using 3D T1-weighted sagittal images (Signa HDxt 3T twin-speed GE sequence inversion-recovery fast-spoiled gradient recalled, TR = 6988 ms, TI = 1100 ms, TE = 3.9 ms, flip angle = 10°, voxel size = 1 × 1 × 1.2 mm³). Gross anatomic abnormalities were ruled out by an experienced neuroradiologist who was blind to clinical diagnosis and evaluated MR imaging scans for each subject.

**Statistical Analysis of Clinical, Motor, and Neuropsychological Data**

Demographic, clinical, and neuropsychological features of patients with FOG+ and FOG− were compared by a t test for independent samples or Fisher exact test, as appropriate.

Given the small number of subjects in each group and to avoid type 1 errors, we evaluated between-group comparisons on cognitive measures with the Mann-Whitney U test, a nonparametric analysis. Computation was supported by the Statistical Package for the Social Sciences, Version 16 (SPSS, Chicago, Illinois) software. Significance threshold was set to P ≤ 0.05.

**VBM**

Data were processed and examined by using SPM8 software (http://www.fil.ion.ucl.ac.uk/spm), where we applied VBM implemented in the VBM8 toolbox (http://dbm.neuro.uni-jena.de/vbm.html) with default parameters incorporating the DARTEL toolbox in SPM8. This was used to obtain a high-dimensional normalization protocol. Images were bias-corrected, tissue-classified, and registered by using linear (12-parameter affine) and nonlinear transformations (warping) within a unified model. Subsequently, the warped GM segments were affine-transformed into Montreal Neurological Institute space and were scaled by the Jacobian determinants of the deformations to account for the local compression and stretching that occurs as a consequence of the warping and affine transformation (modulated GM volumes). Finally, the modulated volumes were smoothed with a Gaussian kernel of 8-mm full width at half maximum. The GM volume maps were statistically analyzed by using the general linear model based on Gaussian random field theory. Statistical analysis consisted of an analysis of covariance with total intracranial volume and age as covariates of no interest. Moreover, neuropsychological data differing between patients with and without FOG were entered as covariates of no interest in an additional analysis of covariance aimed at ruling out the possibility that GM differences were caused by these specific variables. For each group, we defined linear contrasts (1 as group of interest; −1 as a comparing group; 0 as group of noninterest) to test for differences in GM volumes between the groups: 1) each group versus control subjects (patients with FOG+ versus HC: [−1 0 1]; patients with FOG− versus HC: [0 −1 1]); 2) all patient groups (FOG+ and FOG−) versus HC: [−1 1 −1]; and 3) each group versus the other (patients with FOG+ versus patients with FOG−: [−1 1 0]; FOG−patients versus patients with FOG+: [1 −1 0]). Because we did not have an a priori hypothesis concerning a specific region of interest, we applied a whole-brain statistical threshold of P ≤ .05, FWE-corrected for multiple comparisons. Only clusters comprising 100 or more voxels were reported.

To correlate GM volume changes and FOG-Q score, we performed a correlation analysis by using the multiple regression function of SPM8. Correlation analyses were performed inside and outside specific ROIs. We considered ROIs to be the identified regions that showed the most significant GM change in the comparisons between groups. The resulting region-of-interest image was transformed into a binary mask that was applied explicitly to compute regression analysis within PD groups by using a significance level of P ≤ .05, FWE-corrected for multiple comparisons. For exploratory purposes, we subsequently performed correlation analyses on the whole-brain level outside the ROIs (using a statistical threshold of P ≤ .001, uncorrected). The coordinates of voxels exhibiting the greatest group effects were transferred from Montreal Neurological Institute space to Talairach space by using M. Brett’s transformation (http://www.mrc-cbu.cam.ac.uk/Imaging/contents.html).

**Results**

**Clinical Findings**

The demographic characteristics of the patients are shown in the On-line Table. Among 24 patients with PD, 12 were classified as FOG+ and 12 patients, as FOG−. No significant differences were observed between patients with FOG+ and FOG− in age and sex; disease duration; Hoehn and Yahr stage; UPDRS I, III, IV scores; symptom-dominant side; and dopaminergic treatment. As expected, the FOG-Q and UPDRS II scores were significantly higher in FOG+ compared with patients with FOG−; in particular, the UPDRS II score differed significantly between patients with FOG+ and those with FOG− due to the presence in the scale of 2 gait-related items (item 14: freezing; item 15: walking). The 2 groups did not differ on the tremor UPDRS subscore. The total postural instability gait disturbances UPDRS subscore significantly differed between the 2 groups. However, because freezing (item 14) is relevant for the postural instability gait disturbances
subscore, the differences between the 2 groups disappeared when not considering the freezing score (our grouping variable) in the analysis (On-line Table).

**Neuropsychological Data**

Patients with FOG+ performed significantly worse than patients with FOG− on several cognitive tests. In particular, patients with FOG+ compared with patients with FOG− exhibited lower scores on the Frontal Assessment Battery, phonological verbal fluency, and Ten-Point Clock Test, with the first 2 exploring frontal-executive domains and the last mainly investigating spatial programming skill, which requires both executive and visuospatial functions. Furthermore, patients with FOG+ exhibited reduced mean immediate recall of the Rey Auditory Verbal Learning Test 15-word score (Table 1).

**VBM Findings**

There were no differences in global GM, WM, or CSF volume between groups (patients versus HC; patients with FOG+ versus patients with FOG−) (On-line Table). The analysis of regional volume differences revealed that patients with PD as a group (FOG+ and FOG−) showed no significant GM differences compared with HC at a threshold of \( P < .05 \), FWE-corrected. However, a significant difference in GM volume was detected when the FOG+ and FOG− groups were compared. In particular, the FOG+ group showed reduced GM volume in a cluster including the left cuneus, precuneus, lingual gyrus, and posterior cingulate cortex compared with the FOG− group (Fig 1 and Table 2) (\( P < .05 \), FWE-corrected). Moreover, patients with FOG+ exhibited reduced GM volume in the left cuneus and lingual gyrus compared with HC (Table 2) (\( P \leq .05 \), FWE-corrected), whereas, no differences in

| Cognitive Tasks       | FOG+ (n = 12) (mean ± SD) | FOG− (n = 12) (mean ± SD) | U Value | P Value |
|-----------------------|---------------------------|---------------------------|---------|---------|
| MMSE                  | 28.27 ± 2.22              | 27.94 ± 1.58              | 67.00   | .799    |
| Corsi Span            | 4.27 ± 0.64               | 4.77 ± 0.92               | 44.00   | .314    |
| Verbal Span           | 3.54 ± 0.66               | 4.02 ± 0.70               | 45.50   | .211    |
| Immediate recall (Rey)| 34.54 ± 8.44              | 42.64 ± 9.30              | 33.00   | .044*   |
| Delayed recall (Rey)  | 7.65 ± 2.90               | 9.34 ± 2.73               | 40.00   | .118    |
| CA                    | 10.95 ± 0.71              | 10.95 ± 1.77              | 56.50   | .525    |
| RCPM                  | 26.00 ± 4.98              | 27.4 ± 3.21               | 53.50   | .449    |
| FAB                   | 12.41 ± 1.74              | 15.67 ± 1.27              | 3.00    | .001*   |
| Verbal fluency        | 28.52 ± 7.95              | 43.61 ± 8.87              | 16.00   | .001*   |
| Stroop (part II)      | 37.20 ± 10.26             | 35.87 ± 10.21             | 65.50   | .976    |
| Stroop (part III)     | 18.42 ± 4.58              | 22.38 ± 5.71              | 37.00   | .079    |
| TPCT                  | 4.42 ± 2.71               | 7.67 ± 2.67               | 24.00   | .005*   |
| Attentional matrices  | 43.21 ± 9.11              | 48.36 ± 5.44              | 43.50   | .169    |

Note: CA indicates constructional apraxia; FAB, Frontal Assessment Battery; MMSE, Mini-Mental State Examination; TPCT, Ten-Point Clock Test

* Scores are age- and education-adjusted.

Corrected.

**Fig 1.** A, Areas of GM tissue loss deriving from the direct comparison between patients with FOG+ and those with FOG−. The peak differences can be found within the left cuneus, precuneus, lingual gyrus, and posterior cingulate cortex. Results are superimposed on representative axial and coronal sections of the customized GM template, at a threshold of \( P < .05 \), FWE-corrected. B Mean GM volume loss (± standard error of the mean) extracted from the region (Statistical Parametric Mapping coordinates x: −8, y: −82, z: −2) showing a maximal statistical difference between groups. Double asterisks indicate significance at the whole-brain statistical threshold corrected for multiple comparison, \( P \leq .05 \), FWE-corrected.
GM volume were identified in the comparison between HC and patients with FOG−. Moreover, we did not detect other significant GM differences between groups according to linear contrasts (HC versus PD and FOG− versus FOG+) previously defined.

Correlation Analyses
Across all patients with PD, a significant negative correlation was detected between FOG-Q scores and GM loss in the left occipital region (Statistical Parametric Mapping coordinates x: −3, y: −81, z: 16; r value: 6.43; r value: −0.58; cluster [k] = 43; P = .05, FWE-corrected; Fig 2). No significant correlations were detected in voxels outside the ROIs.

Discussion
The aim of our study was to investigate structural brain differences between patients with PD with and without FOG. Our findings revealed that patients with FOG+ exhibited left parietal (precuneus), occipital (cuneus and lingual gyrus), and posterior cingulate cortex atrophy compared with both patients with FOG− and patients with HC. Moreover, consistent with previous studies, we showed worse performance on several cognitive tests in patients with FOG+ compared with patients with FOG−. Our imaging findings were not driven by the observed cognitive differences between patients with FOG+ and patients with FOG− and a significant negative correlation was identified between GM loss of the left occipital cortex and FOG-Q scores.

Gait is traditionally seen as an automatic motor task that requires minimal higher cognitive input. However, a growing body of evidence has recently drawn attention to the importance of cognition in normal walking, which relies not only on the sensorimotor system but also on cerebral networks subserving attention, executive functions, and visuomotor integration.21,22 Mental loading and divided attention may increase the occurrence of FOG episodes23; however, visual and auditory cues can usually overcome FOG.24 Furthermore, Helmich et al25 demonstrated an increased dependence on visual-information processing during a motor imagery task in PD, suggesting that patients with PD increasingly rely on visual cues to control locomotion.26

Our findings are consistent with previous radiotracer studies demonstrating hypometabolism or hypoperfusion of temporoparietal and occipital cortices in patients with FOG. In particular, a [18F]-dihydroxyphenylalanine and FDG-PET study comparing patients with PD with and without FOG revealed that those with FOG had both reduced [18F]-dihydroxyphenylalanine uptake in the caudate and posterior putamen and reduced FDG uptake in posterior parietal areas.27 Moreover, hypometabolism of parietal, temporal, and occipital sensory association cortices was found to limit the effect of deep brain stimulation of the subthalamic nucleus in patients with FOG with PD, suggesting that the metabolic activity of these areas could contribute to the emergence of this disabling symptom.28 A recent fMRI study during imagined walking29 has shown less activation in the superior parietal lobule and anterior cingulate cortex of patients with PD compared with HC. When the patients were divided into FOG+ and FOG− groups, only patients with FOG+ showed decreased activation in the frontal and posterior parietal cortices. This pattern has been confirmed in another recent study.29

Consistent with previous neuropsychological evidence,6,30 patients with FOG+ performed worse than those with FOG− on several cognitive tests. In particular, patients with FOG+ displayed predominant executive-visuospatial impairment as evidenced by reduced scores on the Frontal Assessment Battery, phonemic verbal fluency, and Ten-Point Clock Test. These data suggest a prominent dysfunction of the dorsolateral prefrontal cortex, which has wide connections with posterior cortical areas, namely the parietal and occipital cortices.31 In the occipital cortex, the “ventral stream” (occipitotemporal pathway) is crucial for the identification of objects, whereas the “dorsal stream” (occipitoparietal pathway), which consists of reciprocal projections to the prefrontal and premotor cortices, is involved in mapping the spatial

Table 2: Location and Talaraich coordinates of significant clusters of GM atrophy

| Contrast          | Region              | BA | X     | Y     | Z     | T Value | Cluster Size (kE) | P_{FWE-cor} |
|-------------------|---------------------|----|-------|-------|-------|---------|------------------|------------|
| FOG+ < FOG−       | L posterior cingulate | 30 | −1    | −62   | 9     | 3.90    | 785              | .025a      |
|                   | L lingual gyrus     | 18 | −7    | −78   | 2     | 3.77    |                  |            |
|                   | L cuneus            | 18 | −12   | −78   | 14    | 3.54    |                  |            |
|                   | L precuneus         | 23 | −4    | −61   | 20    | 3.40    |                  |            |
| FOG+ < HC         | L cuneus            | 30 | −2    | −68   | 6     | 4.53    | 374              | .015a      |
|                   | L lingual gyrus     | 18 | −7    | −61   | 1     | 4.27    |                  |            |

Note: kE indicates cluster size; FWE-corr, FWE-corrected; BA, Brodmann area; L, left.

* Significant.
relationships of objects, thereby supporting visually guided movements.32,33

Our cognitive and neuroimaging findings are in agreement with previous VBM studies in patients with PD reporting an association between cognitive impairment and GM reduction in several areas. These include the prefrontal, temporal, parietal, and occipital lobes, with the latter areas more involved in patients with PD with dementia.34,35 A more recent VBM study has reported GM cortical reduction in parietal and occipital areas in patients with PD who performed poorly on visuospatial/visuoperceptual tests.30 Moreover, previous work has shown that ventricular enlargement is associated with early cognitive impairment in patients with PD.37 Although suggested by our cognitive results and previous VBM studies, we did not find GM loss in anterior cortical areas in the whole PD population relative to HC. This apparent inconsistency might reflect the relatively early stage of disease in our PD population: The executive dysfunction detected might not yet be reflected in neuroanatomic changes detectable with VBM, which could emerge with disease progression. Nevertheless, our results are consistent with several imaging studies reporting an association between reduced cerebral glucose uptake in extensive posterior cortical areas and cognitive impairment in patients with PD.38–40

Taken together, our imaging and neuropsychological findings highlight the role of cognitive neural networks in the pathophysiology of FOG in PD. It has been demonstrated that patients with FOG have a pathologic gait pattern,41 even between FOG episodes. However, the degree of these gait alterations has not been correlated with the asymmetry in PD motor signs.42 This lack of correlation suggests that more complex networks are likely involved in their pathogenesis. As reported, previous studies have shown an association between frontal dysfunction and FOG in patients with PD.6,7,30 Recent studies43,44 have demonstrated, by evaluating the influence of space perception on gait in patients with PD, that visuospatial ability appears to be more greatly affected in patients with PD who experience FOG. Furthermore, external visual, mental, and auditory cues (ie, extrinsic drivers) can usually improve FOG. On the basis of previous results as well as those currently presented, we speculate that a specific executive-visuospatial dysfunction may represent the cortical substrate that, in concert with the above-mentioned pathologic gait pattern, could make a subset of patients with PD prone to freeze.

In PD, automaticity is impaired; therefore, automatic movements, such as gait, depend largely on the frontal voluntary system involved in producing internally driven movements (ie, intrinsic drivers). On one hand, our study confirmed that FOG might be related to frontal networks dysfunction; on the other hand, it showed that FOG is associated with reduced GM volume in occipitoparietal pathway areas that likely underlie a dysfunction on spatial perception and discrimination, as also evidenced by neuropsychological results. Visuospatial and visuoperceptual abilities might physiologically play a key role in providing compensation through external cues when intrinsic drivers, relying on frontoexecutive networks, are impaired. Therefore, we further speculate that FOG could be due to a dysfunction of frontoexecutive-based intrinsic drivers,45 not compensated by adequate occipitoparietal-based external cues (ie, external drivers). This speculation is supported by correlation analyses demonstrating that a decreased GM volume in the left occipital cortex was associated with higher FOG-Q scores.

Conclusions

Our data provide further insight into the mechanisms underlying the complex and disabling symptom of FOG in PD. We believe that cognitive impairment, namely executive-visuospatial dysfunction, and structural GM loss in posterior cortical regions may be associated with the development of FOG. However, the present study has some limitations worth noting. First, we relied on self-report of FOG in our patients, which may be influenced by biased recall. While ideal, clinical assessment of FOG represents significant challenges because the behavior is episodic and unpredictable, often not appearing during evaluations. Second, we assessed a relatively small sample of patients, and further studies including larger populations are needed to confirm these preliminary observations.

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