Frontal vs. Posterior Cognitive Dysfunction: Does greater risk of dementia lead to differential gait in Parkinson’s disease?

CAROLINA R. A. SILVEIRA1,2 | ERIC A. ROY1,2 | QUINCY J. ALMEIDA1

1Movement Disorders Research and Rehabilitation Centre, Wilfrid Laurier University, 75 University Avenue West, Waterloo, Ontario, N2L 3C5, CANADA.
2University of Waterloo, Department of Kinesiology, 200 University Avenue West, Waterloo, Ontario, N2L 3G1, CANADA.

Correspondence to: Dr. Carolina R. A. Silveira, Parkwood Institute, 550 Wellington Road, London, ON, CANADA, N6C 0A7.
email: Carolina.Silveira@sjhc.london.on.ca

BACKGROUND: Gait impairment is suggested to predict the onset of dementia in Parkinson’s disease (PD). Interestingly, studies have shown that PD patients with cognitive deficits mediated by posterior brain areas are at greater risk of developing dementia than those with frontal deficits. However, it remains unknown whether PD patients with posterior cognitive deficits show differences in gait when compared to those with frontal deficits.

AIM: This study aimed to compare gait of individuals with PD showing “posterior”, “frontal”, or no cognitive impairment (NCI).

METHOD: Based on a sample of 64 individuals with PD, median scores were calculated for three neuropsychological tests relying on “frontal” and three relying on “posterior” brain areas. Individuals assigned to the Frontal or Posterior groups showed at least 2 out of 3 scores lower than the median in frontal or posterior tests, respectively. Those with 0 or 1 score lower than the median were classified as NCI. Participants walked under single and dual task conditions.

RESULTS: All groups walked slower, with greater variability, a wider base of support, and longer double support in the dual task condition.

CONCLUSION: PD patients with posterior cognitive deficits walk similarly to those with frontal deficits and those with normal cognition.

KEYWORDS: Parkinson’s disease | gait | cognition | dual task

INTRODUCTION

Deficits in gait and cognition are commonly observed in individuals with Parkinson’s disease (PD). Importantly, research has shown that PD patients with postural instability and gait deficits (PIGD) are at greater risk of developing dementia than those with tremor as the predominant motor symptom.1,2 and that gait characteristics at diagnosis predict decline in specific cognitive domains over time in PD.3 In this context, researchers have argued that changes in gait can precede and predict cognitive decline in PD and that the progression of gait and cognitive deficits could result from shared underlying mechanisms.4-7 Thus, gait has been proposed as a surrogate marker for dementia in PD. Although evidence exists that changes in gait predict the incidence of dementia in healthy older adults8,9 and individuals with mild cognitive impairment,10 research investigating gait as a marker for dementia in PD is still in its early stages. If changes in gait are found to reveal individuals at greater risk of...
dementia, then this information could contribute to early identification and intervention prior to dementia onset in PD.

Given that few longitudinal studies have examined the relationship between gait and cognitive decline in PD, a way of testing the potential of gait as a marker could be to compare gait characteristics of individuals with PD at a higher and lower risk of dementia and determine whether those at higher risk show distinct gait behavior. Previous research has demonstrated that two cognitive profiles at diagnosis differently predict the risk of dementia in PD. The first was characterized by deficits in cognitive domains that rely on frontal lobe functioning (i.e. executive functions) and respond to dopaminergic therapy, while the second was characterized by deficits in cognitive domains mediated by posterior brain areas (i.e. memory, language, and visuospatial functioning) that do not respond to dopamine. At 5 and 10-year follow up, it was found that a larger number of individuals with “posterior” deficits at diagnosis had developed dementia in comparison to those with “frontal” deficits. \cite{11,12} It was concluded that individuals with PD who show deficits in “posterior” cognitive domains at diagnosis of PD are at greater risk of dementia than those showing deficits in “frontal” cognitive domains. Thus, if gait is a marker for dementia in PD, then it could be hypothesized that differences in gait may exist between non-demented PD patients with “posterior” and “frontal” cognitive deficits.

To date, no study has directly compared gait of PD patients with predominantly “posterior” or “frontal” cognitive deficits. However, associations between gait and cognitive deficits in PD have been extensively reported. Interestingly, studies have consistently shown that slower gait speed and greater step-to-step variability were linked to frontal cognitive deficits. \cite{13,14} In contrast, the relationship between deficits in gait and cognitive domains mediated by posterior brain areas are inconsistent and more rarely reported. Although objective measures of gait stability, such as double support, \cite{15} as well as the severity of gait impairment and postural instability from the Unified Parkinson’s disease Rating Scale, \cite{16} have been associated with posterior cognitive deficits, these associations do not reveal whether differences in gait exist between groups at different risk for dementia.

This study aimed to compare gait of non-demented individuals with PD showing predominantly “posterior”, “frontal”, or no cognitive impairment (NCI). Given the associations between gait instability and posterior cognitive deficits, it was expected that individuals with predominant posterior deficits would walk with a wider base of support and longer double support than those with frontal deficits and NCI. Since previous research has shown associations between step-to-step variability and frontal deficits, it was hypothesized that individuals with predominantly frontal deficits would walk with greater step-to-step variability than those with posterior deficits and NCI. As gait speed could be associated with frontal and/or posterior deficits, it was expected that the frontal as well as posterior groups would walk slower than the NCI group. Finally, it was hypothesized that, due to its greater demands on cognitive processing, differences between groups would have greater magnitude when participants performed dual task gait compared to single task gait.

METHODS
This study was approved by Wilfrid Laurier University (WLU) and University of Waterloo (UW) research ethics boards (process identification numbers: 19582 and 3922) and informed consent was obtained prior to participation.

Participants

Sixty-four individuals with PD were recruited from the Movement Disorders Research and Rehabilitation Centre database at WLU (Waterloo, Canada). Inclusion criteria were men and women diagnosed with idiopathic PD by a Neurologist and able to walk 10 meters unassisted. Participants were excluded if one of the following criteria was met: history of neurological diseases other than PD, uncontrolled diabetes, uncontrolled hypertension, history of cardiovascular disease, chronic obstructive pulmonary disease, uncorrected visual impairments, and a diagnosis of dementia. Participants' demographic information (age, sex, and years of education), general cognitive function (Montreal Cognitive Assessment), depressive symptoms (Geriatric Depression Scale), and motor disease severity (Unified Parkinson’s disease Rating Scale motor subscale) were collected at baseline.

Group Assignment

Based on performance of all 64 individuals with PD, median scores were calculated for each neuropsychological test relying predominantly on “frontal” or “posterior” brain functioning. Participants were assigned to Frontal (n=14; age=68.50 (7.83)) or Posterior (n=12; age=69.25 (8.50)) groups if they showed at least 2 of 3 scores lower than the median in frontal or posterior tests, respectively. Participants with 0 or only 1 test score lower than the median were classified as non-cognitively impaired (NCI n=22; age=67.59 (8.09)), and those with lower scores in both frontal and posterior tests were excluded (n=16). Groups' demographic and clinical information are displayed in Table 1.

Table 1 - Demographic and clinical information

<table>
<thead>
<tr>
<th>Group</th>
<th>Age (years)</th>
<th>Sex (M/F)</th>
<th>Education (years)</th>
<th>MoCA</th>
<th>GDS</th>
<th>UPDRS-III</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frontal n=14</td>
<td>68.50 (7.83)</td>
<td>8/6</td>
<td>14.00 (3.80)</td>
<td>25.78 (1.88)</td>
<td>5.35 (3.73)</td>
<td>24.25 (7.19)</td>
</tr>
<tr>
<td>Posterior n=12</td>
<td>69.25 (8.50)</td>
<td>10/2</td>
<td>16.58 (3.57)</td>
<td>25.00 (3.69)</td>
<td>6.66 (5.82)</td>
<td>23.00 (7.04)</td>
</tr>
<tr>
<td>NCI n=22</td>
<td>67.59 (8.09)</td>
<td>10/6</td>
<td>15.45 (3.01)</td>
<td>27.83 (2.19)</td>
<td>6.36 (4.30)</td>
<td>25.09 (9.30)</td>
</tr>
</tbody>
</table>

Legend: NCI – non-cognitively impaired; Education – number of years completed in school; MoCA – Montreal Cognitive Assessment; GDS – Geriatric Depression Scale; UPDRS-III – Unified Parkinson’s disease Rating Scale motor subsection; c Posterior different from NCI.

Cognitive Assessment

To assess cognitive domains relying on frontal brain functioning, three executive function tests were employed: Digit Span, Stroop Test, and Trail Making Test (TMT). Median values were calculated based on the total number of digit sequences correctly recalled in the forward and backward conditions of the Digit Span, interference scores of the Stroop test (number of items correctly named in the Color-Word condition minus items correctly named in the Color condition), and the difference between parts B and A of the TMT in seconds. Assessment of cognitive domains relying on posterior brain functioning was conducted using the Copy of the Intersected Pentagons from the Mini Mental State Examination (visuospatial), Semantic Verbal Fluency (language), and the Short Form
California Verbal Learning Test\textsuperscript{22} (memory). Median values were calculated based on the total score in the Copy of the Intersected Pentagons, total number of words generated in the Semantic Verbal Fluency (semantic category: animals), and number of correct words recalled (immediate recall) in the California Verbal Learning Test.

Gait Assessment

Participants walked on a 10-meter long Zeno\textregistered Walkway System (ProtoKinetics, Havertown, PA, USA) under single and dual task conditions (3 trials per condition). Three trials per condition were performed to examine the effects of task novelty (1\textsuperscript{st} trial) and adaptation (subsequent trials), while making the protocol feasible to participants with different severities of gait impairment. In the single task condition participants were instructed to walk at their normal pace, whereas in the dual task condition they were instructed to walk and count the number of times that two pre-assigned digits were spoken on an audio track.\textsuperscript{23} Participants started walking 1 meter prior to stepping onto the carpet and continued walking 1 meter after stepping off the carpet to account for gait acceleration and deceleration phases. In addition, the first and last steps were removed from the analysis to eliminate the effect of surface transition. Gait speed, step length and step time variability (coefficient of variation=(standard deviation/mean)x100), base of support (step width), and percentage of time spent in double support ((total double support time/stride time)x100) were calculated using the ProtoKinetics Movement Analysis Software (PKMAS) version 507c7c.

Statistical Analysis

One-way analysis of variance (ANOVA) or a non-parametric test (Kruskal Wallis) was used to compare demographic, clinical, and neuropsychological information between groups. Repeated Measures ANOVA tested differences in gait between groups in task conditions and experimental trials [Group x Condition x Trial]. Repeated Measures ANOVA was used to compare participant performance in the secondary task while sitting versus walking [Group x Condition]. Significant differences were examined using Tukey’s HSD post-hoc test or non-parametric pairwise comparisons (Mann-Whitney U) and alpha levels were kept at $<0.05$.

RESULTS

Demographic and clinical information

Overall, groups had similar age, years of education, severity of depressive symptoms, and motor disease severity. However, group differences were found in the MoCA ($F(2, 45)=4.71; p=0.013; \eta^2_p=0.17$), where the Posterior group presented a worse general cognitive status than the NCl group ($p=0.017$).

Cognition

Neuropsychological testing was used primarily to classify individuals into experimental groups based on median performance per test. However, performance in neuropsychological tests was also compared after group assignment in order to confirm whether distinct patterns of deficits emerged. Group differences were found for the Stroop
test ($F(2, 45)=5.86; p=0.005; \eta^2_p =0.20$), TMT ($\chi^2(2)=14.21; p=0.001$; mean rank score Frontal=35.29, Posterior=25.17, NCI=17.27), Digit Span ($F(2, 45)=13.75; p<0.001; \eta^2_p =0.37$), Copy of the Intersected Pentagons ($\chi^2(2)=18.90; p<0.001$; mean rank score Frontal=26.29, Posterior=12.92, NCI=29.68), and Semantic Fluency ($F(2, 41)=7.09; p=0.002; \eta^2_p =0.25$). Post-hoc analysis showed that the Frontal group performed worse than both Posterior and NCI groups in the Stroop (Posterior $p=0.038$; NCI $p=0.005$), TMT (Posterior $p=0.035$; NCI $p<0.001$), and Digit Span (Posterior $p=0.001$; NCI $p=0.0001$) tests, whereas no differences between Posterior and NCI groups were found in these tests. The Posterior group performed worse than both Frontal ($p=0.004$) and NCI ($p<0.001$) groups in the Copy of Intersected Pentagons, and worse than the NCI group ($p=0.001$) in the Semantic Fluency test. Group performances in the neuropsychological tests are presented in Table 2.

### Table 2 - Group performance in neuropsychological tests

<table>
<thead>
<tr>
<th>Group</th>
<th>Stroop</th>
<th>TMT</th>
<th>Digit Span</th>
<th>Pentagons</th>
<th>Semantic Fluency</th>
<th>CVLT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frontal n=14</td>
<td>-33 (10.74)</td>
<td>93.85 (65.78)</td>
<td>13.92 (2.43)</td>
<td>9.78 (0.42)</td>
<td>20.30 (3.19)</td>
<td>6.78 (1.57)</td>
</tr>
<tr>
<td>Posterior n=12</td>
<td>-24.08 (8.07)</td>
<td>50.91 (37.79)</td>
<td>18.66 (3.67)</td>
<td>8.83 (1.11)</td>
<td>16.30 (3.30)</td>
<td>5.83 (1.85)</td>
</tr>
<tr>
<td>NCI n=22</td>
<td>-22.95 (8.05)</td>
<td>31.77 (20.88)</td>
<td>19.40 (3.26)</td>
<td>9.86 (0.63)</td>
<td>22.66 (5.36)</td>
<td>7.18 (1.43)</td>
</tr>
</tbody>
</table>

Legend: Stroop – difference between the number of items correctly named in the Color-Word and Color conditions (negative value denotes greater interference); TMT - difference between parts B and A of the Trail Making Test in seconds (larger value denotes worse performance); Digit Span - total number of digit sequences correctly recalled in the forward and backward conditions (larger value denotes better performance); Pentagons - total score on the Copy of Intersected Pentagons (larger value denotes better performance); Semantic Fluency - total number of words correctly generated (larger value denotes better performance); CVLT - number of correct words recalled (immediate recall) in the short form of the California Verbal Learning Test (larger value denotes better performance). $^a$ Frontal different from Posterior and NCI; $^b$ Posterior different from Frontal and NCI; $^c$ Posterior different from NCI.

**Gait**

A main effect of Condition was found in the analysis of all gait-related variables, showing that when performing the dual task all groups walked slower ($F(1, 45)=72.45; p<0.001; \eta^2_p =0.61$), with greater step length ($F(1, 45)=12.17; p=0.001; \eta^2_p =0.21$) and step time ($F(1, 45)=11.06; p=0.001; \eta^2_p =0.19$) variability, a wider base of support ($F(1, 45)=16.60; p=0.001; \eta^2_p =0.26$), and longer double support ($F(1, 45)=56.08; p<0.001; \eta^2_p =0.55$) compared to single task walking. However, no group differences were identified in either single or dual task walking conditions. Figure 1 shows the average values between groups for each outcome measure.
Figure 1. Groups showed similar behavior during single and dual task walking.

Performance in the secondary task was similar between groups while seated, confirming that all participants were able to perform this digit monitoring task. A group by condition interaction ($F(2,45)=5.43; p=0.007; \eta^2_p =0.19$) showed that only the Frontal group presented significantly more errors when walking compared to sitting ($p=0.029$).

DISCUSSION

To our knowledge, this is the first study to directly compare gait behavior of non-demented PD patients with either predominantly frontal, posterior, or no cognitive impairment. Contrary to the study’s hypotheses, results showed that all groups walked
similarly during single and dual task conditions. Despite previous investigations showing associations between gait and cognitive deficits in PD, 3, 13-15, 24 the gait of participants in the Frontal and Posterior groups did not differ from those in the NCI group. The lack of differences between groups is also in contrast with findings that the PIGD motor subtype is a predictor of cognitive decline in PD. 1, 2 In the current study, participants were categorized based on cognitive profile in order to identify individuals with deficits previously linked to an increased risk of dementia in PD. However, early research has shown that gait characteristics at diagnosis are better predictors of cognitive decline over time than cognitive characteristics. 3 Therefore, it might be that categorization based on cognitive profile alone does not fully capture the relationship between gait, cognition, and risk of dementia in PD. It is also important to note that the relationship between gait and cognition in PD may depend on stratification based on predominant motor symptoms (tremor vs. PIGD), since associations between gait and cognitive decline in PD have been found in those with predominant PIGD but not predominant tremor symptoms 1-2. Thus, in addition to cognitive profile, PIGD subtype should be taken into consideration when examining the relationship between, gait, cognition, and risk of dementia in PD.

With respect to performance in the secondary task, it was found that participants in the Frontal group had worse performance while walking compared to sitting. This result suggests that individuals with Frontal deficits experienced greater interference in cognitive processing or that these individuals prioritized gait in order to sustain performance during the dual task condition. Notably, the secondary task utilized in this study relied on working memory, a cognitive function largely processed in the frontal areas of the brain. 25 Thus, the nature of the secondary task likely contributed to individuals in the Frontal group suffering greater interference than those in the Posterior and NCI groups. Future research should examine how secondary tasks tapping into different cognitive domains influence gait in PD patients with distinct cognitive profiles.

Findings from this research may have been influenced by its cross-sectional design, as it would be preferable to characterize gait and cognitive profiles at the time of diagnosis and follow individuals longitudinally in order to test the study hypotheses. In addition, sample size and assignment criteria may have attenuated differences between groups. Thus, it is suggested that future studies with larger samples use normative data for neuropsychological tests in order to identify and categorize individuals with impaired cognitive function.

CONCLUSION

The present study demonstrated that, in a cross-sectional design, PD patients with a cognitive profile linked to increased risk of dementia could not be distinguished based on gait.

REFERENCES


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