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Visuospatial and visuoperceptual impairment in relation to global atrophy in Parkinson’s disease*

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Parkinson’s disease (PD) patients differed from controls of similar age in visuospatial and visuoperceptual functions at diagnosis moment, and these deficits have been shown to be neuropsychological markers of evolution to dementia. The aim of this study was to relate these dysfunctions with measures of brain. The sample of this study consisted of 92 PD patients and 36 healthy subjects matched by age, sex and education. All subjects were evaluated with Judgment of Line Orientation, Visual Form Discrimination and Facial Recognition Tests and magnetic resonance imaging at 3 Tesla. We found significant differences between patients and controls in all three tests and in the mean of cortical thickness, gray matter volume and ventricular system. All visuospatial and visuoperceptual tests correlated with the measures of global atrophy suggesting that they are reflecting the brain degeneration associated to PD.

Keywords: Parkinson’s disease, visuospatial/visuoperceptual, MRI, cortical thickness, brain atrophy.

Alteraciones visuoespaciales y visuoperceptivas en la enfermedad de Parkinson y su relación con parámetros de atrofia global

Los pacientes con enfermedad de Parkinson (EP) comparados con controles de similar edad muestran alteraciones visuoespaciales y visuoperceptivas desde el momento del diagnóstico, y se ha demostrado que estas alteraciones

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Visuospatial impairment and atrophy in PD

actúan como marcadores neuropsicológicos de la evolución a demencia. El objetivo de este estudio es relacionar estas alteraciones con las medidas globales de atrofia cerebral. La muestra consta de 92 EP y 36 sujetos sanos emparejados por edad, sexo y educación. Todos los sujetos fueron evaluados mediante el Judgment of Line Orientation, Visual Form Discrimination y Facial Recognition Test y se obtuvieron datos de neuroimagen estructural con un aparato de Resonancia Magnética de 3 tesla. Se observaron diferencias entre los pacientes y los controles en los tres tests usados y en las medidas de grosor cortical total, volumen de sustancia gris y sistema ventricular. Todos los tests visuoespaciales y visuoperceptivos correlacionaron con medidas de atrofia global, lo cual sugiere que son indicadores útiles de la degeneración cerebral asociada a la EP.

Palabras clave: Enfermedad de Parkinson, imágenes de resonancia magnética, grosor cortical, alteraciones visuoespaciales y visuoperceptivas, atrofia cerebral.

Introducción

Parkinson’s disease (PD) is considered to be one of the most common neurological diseases associated with aging. The prevalence ranges from 1-3% in the general population (Galvin, Pollack, & Morris, 2006; Riedel et al., 2008). Classically, PD was regarded as a motor disease, but it is now recognized as a systemic neurodegenerative illness. Cognitive dysfunction and impairment as well as mood disorders and apathy have been increasingly viewed as important symptoms that accompany the disease even in preclinical stages. It has been stated that, after 20 years, 80% of PD patients evolve to dementia, whereas only 25% of them are recognized by clinicians in routine daycare (Svenningsson, Westman, Ballard, & Aarsland, 2012). The disease may course with impaired executive function (Williams-Gray, Foltynie, Brayne, Robbins, & Barker, 2007; Williams-Gray et al., 2009; Aarsland et al., 2009), visuospatial and visuoperceptual dysfunctions (Williams-Gray et al., 2007-2009; Aarsland, Brønnick, Larsen, Tysnes, & Alves, 2009) and verbal memory impairment (Aarsland et al., 2009) but relatively spared language, except for semantic fluency (Williams-Gray et al., 2007; 2009).

Visuospatial and visuoperceptual deficits can be seen early in the disease course, but their main interest is their power to predict subsequent evolution to dementia (Williams-Gray et al., 2009; Svenningsson et al., 2012). Several studies have assessed the differential impairment of these functions in subcortical dementias, and PD has been stated to have a distinct pattern of alteration compared with other types of dementias, such as Alzheimer’s disease (Ala, Hughes, Kyrouac, Ghobria, & Elble, 2001; Cormack, Aarsland, Ballard, & Tovée, 2004; Tiraboschi et al., 2006; Saka & Elibol, 2009). For this reason, they have been regarded as screening tools for cognitive impairment and evolution to dementia in clinical settings (Williams-Gray et al., 2009; Helmes, 2013).
Previous neuroimaging research has focused on describing the progress of global atrophy in PD patients using total gray matter and ventricular measures. Gray matter changes have been related to cognitive decline and dementia (Ibarretxe-Bilbao et al., 2012; Zarei et al., 2013; Camicioli et al., 2011) and recent research has demonstrated that gray matter decrease holds a high discriminant power in accurately identifying demented PD patients (Zarei et al., 2013). Nevertheless, progressive gray matter reductions have already been described in early stages of the disease (Ibarretxe-Bilbao et al., 2012). Gray and white matter losses are related to enlargement of the ventricular system in PD patients. Apostolova et al. (2010) found that left and right ventricles were significantly larger in PD patients with dementia compared with healthy subjects and their non-demented peers. Also, it has been shown that ventricular volumes are significantly increased in PD patients with MCI and correlate with memory performance (Dalaker et al., 2011).

In sum, visuospatial and visuoperceptual impairment and atrophy measures have been associated with cognitive decline and exhibit a predictive value in the evolution to dementia, but hitherto and to our knowledge, the relationship between these two measures has not been studied. Thus, the aim of this study is to assess the neuropsychological performance in these cognitive functions and its relation with global structural changes in a large sample of non-demented PD patients.

**Methods**

**Subjects**

This study included 121 patients, recruited from an outpatient movement disorders clinic (Parkinson’s Disease and Movement Disorders Unit, Service of Neurology, Hospital Clinic, Barcelona, Spain), and 49 healthy subjects, recruited from the Institut de l’Envelleciment (Barcelona, Spain). Controls were matched for age, sex and years of education to their patient peers. This study was approved by the institutional ethics committee; all subjects agreed to participate voluntarily and written informed consent was obtained after full explanation of the procedures.

Inclusion criteria for patients consisted of meeting the diagnostic criteria for PD established by the UK PD Society Brain Bank (Daniel & Lees, 1993). The following exclusion criteria were considered: [1] presence of dementia according to the Movement Disorders Society criteria (Dubois, Burn, Goetz, & Aarsland; 2007), [2] Hoehn and Yahr scale score $\geq 3$, [3] young-onset PD, [4] presence of psychiatric and/or neurologic comorbidity, [5] low IQ score (scaled score score <8 on the Vocabulary subtest from the WAIS-IV -Wechsler Adults Intelligence Scale), [6] Mini-Mental State Examination (MMSE) score <24, [7] claustrophobia, [8] pathological findings non compatible with PD in MRI and [9] MRI artifacts. (demographic data of patients and healthy controls are summarized in table 1).
TABLE 1. DEMOGRAPHIC AND CLINICAL DATA OF THE SAMPLE.

<table>
<thead>
<tr>
<th>Demographics</th>
<th>Patients (n=92)</th>
<th>Controls (n=36)</th>
<th>t, χ²</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>63.98 ± 11.10</td>
<td>63.36 ± 10.53</td>
<td>-0.287 a</td>
<td>0.775</td>
</tr>
<tr>
<td>Sex (male/female)</td>
<td>37/55</td>
<td>19/17</td>
<td>0.521 b</td>
<td>0.300</td>
</tr>
<tr>
<td>Education, years</td>
<td>10.61 ± 5.36</td>
<td>11.36 ± 4.34</td>
<td>0.750 a</td>
<td>0.454</td>
</tr>
<tr>
<td>Age at onset</td>
<td>56.04 ± 12.19</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Evolution, years</td>
<td>8.39 ± 5.91</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients with</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>hallucinations</td>
<td>n=19</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: Values are mean ± Standard Deviation (SD); aStudent t test statistics; bChi squared statistics.

Neuropsychological Assessment

The neuropsychological assessment comprised the evaluation of all cognitive functions, and the assessment of visuospatial and visuoperceptual functioning was obtained through the following neuropsychological tests: [1] Benton’s Judgment of Line Orientation (JLO; Benton, Hannay, & Varney; 1975), [2] Benton’s Visual Form Discrimination (VFD; Benton, Sivan, & Hamsher; 1994), and [3] Benton’s Facial Recognition (FRT; Benton et al., 1994) tests.

Global measures of atrophy analysis

FreeSurfer software (version 5.1; available at http://surfer.nmr.harvard.edu) was used to assess global brain atrophy. Along the process, cortical surface 3D model of cortical thickness (CTh) is created using intensity and continuity information, as described in detail by Fischl & Dale (2000). Independent steps are performed in the initial preprocessing of images for each subject: removal of non-brain tissue, automated Talairach transformation, intensity normalization, tessellation of the gray matter / white matter boundary, automated topology correction and surface deformation to optimally place the gray matter / white matter and gray matter / cerebrospinal fluid (CSF) boundaries. The resulting representation of CTh is calculated as the distance between tissue boundaries. All surface models in our study were visually inspected for accuracy and were carefully manually corrected if necessary.

Segmentation of brain volume and cortical thickness parcellations were obtained based on an automatic procedure included in Freesurfer 5.1 (Desikan et al.,
Variables used in order to calculate global measurement of atrophy were: bilateral mean thickness, total cortex volume, total gray matter volume, and the lateral ventricular system (left and right lateral ventricles).

### Statistical Analyses

Statistical analyses of visuospatial and visuoperceptual functions, global atrophy, and neuropsychological and clinical variables were carried out using the statistical package PASW-20 (2011; SPSS, Inc., Chicago, IL). Z scores were obtained in order to assess patients’ performance in visuospatial and visuoperceptual tests. Clinical and demographical variables and brain atrophy group differences were analyzed with Student \( t \) test statistics or ANCOVA in case of volumetric global measures of atrophy. Cohen’s \( d \) was used to measure effect size in clinical and demographical variables (Ellis, 2009). Correlations analyses were performed among visuospatial and visuoperceptual tests and those with global atrophy measures. All the volumetric analyses were controlled for intracranial volume (ICV).

### Results

#### Clinical and demographical analyses

Raw scores significantly differed between PD patients and healthy subjects in MMSE, NPI (Cumming’s Neuropsychiatric Inventory) and BDI (Beck Depression Inventory-II), and all effect sizes indicated a large effect (see table 2). Age and education showed correlations with visuospatial and visuoperceptual tests, which were higher in all cases for PD patients than for controls. The highest correlations were seen between age and VFD, and education and FRT.

<table>
<thead>
<tr>
<th></th>
<th>Patients</th>
<th>Controls</th>
<th>( t )</th>
<th>( p ) value</th>
<th>( d )</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMSE</td>
<td>29.05 ± 1.11</td>
<td>29.69 ± 0.47</td>
<td>4.581</td>
<td>.0001</td>
<td>.75\textsuperscript{b}</td>
</tr>
<tr>
<td>NPI</td>
<td>5.67 ± 7.39</td>
<td>1.68 ± 3.13</td>
<td>-4.124</td>
<td>.0001</td>
<td>.70\textsuperscript{a}</td>
</tr>
<tr>
<td>BDI</td>
<td>10.87 ± 6.15</td>
<td>6.15 ± 5.64</td>
<td>-2.845</td>
<td>.0001</td>
<td>.80\textsuperscript{a}</td>
</tr>
<tr>
<td>UPDRS-III</td>
<td>17.60 ± 10.92</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H&amp;Y</td>
<td>1.89 ± 0.57</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LEDD</td>
<td>803.70 ± 494.18</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: MMSE: Mini-Mental State Examination; NPI: Cummings’ Neuropsychiatric Inventory; BDI: Beck Depression Inventory; UPDRS-III: Unified Parkinson’s Disease Rating Scale-III; H&Y: Hoehn & Yahr; LEDD: Levodopa Equivalent Daily Dose.

\( \textsuperscript{a} \)Cohen’s \( d \) statistics (Group 1=Patients/Group 2=Control subjects); \( \textsuperscript{b} \)Group 1=Control subjects/Group 2=Patients.
Clinical variables showed significant correlations with visuospatial and visuoperceptual tests, except for JLO and VFD with LEDD (Levodopa Equivalent Daily Dose) and healthy subjects with MMSE. However, visuospatial and visuoperceptual performance was still significantly different between patients and controls when controlling the possible effect of these variables (table 3 and table 4).

**Table 3. Pearson’s r correlations among demographical variables and visuospatial and visuoperceptual tests.**

<table>
<thead>
<tr>
<th></th>
<th>Age</th>
<th>Education</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HC</td>
<td>PD</td>
</tr>
<tr>
<td>JLO</td>
<td>-.471**</td>
<td>-.502**</td>
</tr>
<tr>
<td>VFD</td>
<td>-.243</td>
<td>-.416**</td>
</tr>
<tr>
<td>FRT</td>
<td>-.498**</td>
<td>-.432**</td>
</tr>
</tbody>
</table>

Note: JLO: Judgment of Line Orientation; VFD: Visual Form Discrimination; FRT: Facial Recognition Test; IQ: Intelligence Quotient; HC: Healthy Controls; PD: Parkinson’s Disease; **p<0.05.

**Table 4. Pearson’s r correlations among clinical variables and visuospatial and visuoperceptual tests.**

<table>
<thead>
<tr>
<th></th>
<th>Age at onset</th>
<th>H&amp;Y</th>
<th>MMSE</th>
<th>LEDD</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PD</td>
<td>HC</td>
<td>PD</td>
<td>PD</td>
</tr>
<tr>
<td>JLO</td>
<td>-.348**</td>
<td>-.277*</td>
<td>-.132</td>
<td>.405**</td>
</tr>
<tr>
<td>VFD</td>
<td>-.275**</td>
<td>-.238*</td>
<td>.130</td>
<td>.279**</td>
</tr>
<tr>
<td>FRT</td>
<td>-.221*</td>
<td>-.191</td>
<td>.235</td>
<td>.487**</td>
</tr>
</tbody>
</table>

Note: JLO: Judgment of Line Orientation; VFD: Visual Form Discrimination; FRT: Facial Recognition Test; H&Y: Hoehn & Yahr; MMSE: Mini-Mental State Examination; LEDD: Levodopa Equivalent Daily Dose; HC: Healthy Controls; PD: Parkinson’s Disease; *p<0.05; **p<0.01.

**Visuospatial and visuoperceptual performace**

Visuospatial and visuoperceptual tests showed significant differences between patients and healthy subjects, indicating that patients performed significantly worse than controls. VFD (z=-0.671) and FRT (z=-0.678) showed the greatest z scores, corresponding to a moderate difference between PD patients and controls, followed by JLO (z=-0.504; see figure 1). Correlations between visuospatial and visuoperceptual tests (JLO, VFD and FRT) showed that all tests exhibit high correlations between them in PD patients, but not in healthy subjects (see table 5). The highest
correlations were seen between JLO and FRT, followed by JLO and VFD and, lastly, VFD and FRT.

Note: JLO: Judgment of Line Orientation; VFD: Visual Form Discrimination; FRT: Facial Recognition Test.

Figure 1. Graphic representation of visuospatial and visuoperceptual z scores.

Table 5. Pearson’s R correlations between visuospatial and visuoperceptual tests.

<table>
<thead>
<tr>
<th></th>
<th>JLO</th>
<th>VFD</th>
<th>FRT</th>
</tr>
</thead>
<tbody>
<tr>
<td>**HC</td>
<td>1</td>
<td>.237</td>
<td>.260</td>
</tr>
<tr>
<td>**PD</td>
<td></td>
<td>.467**</td>
<td>.493**</td>
</tr>
</tbody>
</table>

Note: JLO: Judgment of Line Orientation; VFD: Visual Form Discrimination; FRT: Facial Recognition Test; HC: Healthy Controls; PD: Parkinson’s Disease; **p<0.01

Global atrophy measures

Mean group differences in global atrophy analysis showed significant decreases in PD patients (see table 6) in the left lateral ventricular system, cortex volume, total gray matter volume and mean cortical thickness.
Global atrophy correlations analyses (see table 7) showed, except for JLO and right inferior lateral ventricle, significant results between JLO, VFD and FRT and mean thickness bilaterally, lateral ventricular system, total gray matter volume and cortex volume.

**Table 6. Mean values and standard deviations of global atrophy measures for PD patients and healthy subjects. Mean differences between groups assessed with Student t test statistics or F in ANCOVA.**

<table>
<thead>
<tr>
<th></th>
<th>Patients</th>
<th>Controls</th>
<th>F/F</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
</tr>
<tr>
<td>Mean Thickness(^a)</td>
<td>2.44</td>
<td>0.11</td>
<td>2.49</td>
</tr>
<tr>
<td>Cortex Volume(^b)</td>
<td>428325.38</td>
<td>48131.76</td>
<td>442560.56</td>
</tr>
<tr>
<td>Total Gray Matter Volume(^b)</td>
<td>601545.31</td>
<td>64185.97</td>
<td>611615.79</td>
</tr>
<tr>
<td>Left Lateral Ventricle(^b)</td>
<td>12783.52</td>
<td>7770.29</td>
<td>9886.78</td>
</tr>
<tr>
<td>Left Inferior Lateral Ventricle(^b)</td>
<td>512.58</td>
<td>354.91</td>
<td>372</td>
</tr>
<tr>
<td>Right Lateral Ventricle(^b)</td>
<td>11549.65</td>
<td>6464.13</td>
<td>9388.94</td>
</tr>
<tr>
<td>Right Inferior Lateral Ventricle(^b)</td>
<td>383.96</td>
<td>264.12</td>
<td>328.25</td>
</tr>
<tr>
<td>Total Ventricular System(^b)</td>
<td>25229.71</td>
<td>14297.25</td>
<td>19975.97</td>
</tr>
</tbody>
</table>

*Note:* \(^a\) All values are mm. \(^b\) All values are mm\(^3\). SD: Standard Deviation. \(^t\): Student t test statistics. \(^F\): ANCOVA (volumetric measures of global atrophy were controlled by total intracranial volume). *\(p<0.05\) **\(p<0.01\)

**Table 7. Pearson’s \(r\) correlations between visuospatial and visuoperceptual tests and global measures of atrophy.**

<table>
<thead>
<tr>
<th>Test</th>
<th>JLO</th>
<th>VFD</th>
<th>FRT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(r)</td>
<td>(r)</td>
<td>(r)</td>
</tr>
<tr>
<td>Mean Thickness</td>
<td>.294**</td>
<td>.226*</td>
<td>.314**</td>
</tr>
<tr>
<td>Cortex Volume</td>
<td>.402**</td>
<td>.304**</td>
<td>.269**</td>
</tr>
<tr>
<td>Total Gray Matter Volume</td>
<td>.411**</td>
<td>.282**</td>
<td>.303**</td>
</tr>
<tr>
<td>Left Lateral Ventricle</td>
<td>-.224*</td>
<td>-.257**</td>
<td>-.316**</td>
</tr>
<tr>
<td>Left Inferior Lateral Ventricle</td>
<td>-.184*</td>
<td>-.253**</td>
<td>-.280**</td>
</tr>
<tr>
<td>Right Lateral Ventricle</td>
<td>-.296**</td>
<td>-.305**</td>
<td>-.368**</td>
</tr>
<tr>
<td>Right Inferior Lateral Ventricle</td>
<td>-.151</td>
<td>-.190*</td>
<td>-.284**</td>
</tr>
<tr>
<td>Total Ventricular volume</td>
<td>-.266**</td>
<td>-.290**</td>
<td>-.354**</td>
</tr>
</tbody>
</table>

*Note:* JLO: Judgment of Line Orientation; VFD: Visual Form Discrimination; FRT: Facial Recognition Test. *\(p<0.05\) **\(p<0.01\).
Discussion

In this study, we have shown that PD patients had significantly lower performance in Benton’s tests than healthy subjects, and they exhibited significant correlations with measures of global atrophy and clinical variables. Thus, these results suggest that visuospatial and visuoperceptual performance reflects the structural deterioration that PD patients show from early stages of the disease.

Previous studies have shown that visuospatial and visuoperceptual dysfunctions are already present in non-demented PD patients (Foltynie, Brayne, Robbins, & Barker, 2004; Muslimovic, Post, Speelman, & Schmand, 2005; Aarsland et al., 2009; Elgh et al., 2009), and they have been related to gray matter volumetric changes using voxel-based-morphometry (VBM), mainly in posterior cortex and bilateral parietal regions (Pereira et al., 2009), but this relationship has not been studied using screening measures of global atrophy.

From a classical point of view, the visual system has been postulated to be divided into two streams (Ungerlieder & Mishkin, 1982; Goodale & Milner, 1992), namely, a dorsal pathway, related to spatial aspects of visual processing, and a ventral stream, which refers to the perceptual features of visual processing. However, a recent review has postulated the implication of widespread regions, suggesting the involvement of functional subdivisions, including parieto-premotor, parieto-temporal and parieto-prefrontal pathways. In addition, the authors of this review have highlighted the relevance of common areas between these pathways, and the possible convergence of this information in posterior temporal cortical regions (Kravitz, Saleem, Bake, & Mishkin; 2011). In agreement with this study, we have found that visuospatial and visuoperceptual tests show high intercorrelations, suggesting that they have similar substrates.

Visuospatial and visuoperceptual performance and global atrophy have increasingly been proposed as predictors of cognitive impairment and progression to dementia in PD patients (Apostolova et al., 2011; Williams-Gray et al., 2007-2013; Zarei et al., 2013). Recently, in a 10-year population-based longitudinal study, it has been shown that the pentagon item from MMSE, as a measure of these functions, has a predictive value in the evolution to dementia (Williams-Gray et al., 2013). In addition, Compta et al. (2013) also found that visuospatial and visuoperceptual dysfunction was a significant baseline dementia predictor in an 18-month follow up study. The authors also report the evolution of cortical thinning in PD patients to be a biomarker of dementia-risk and disease progression. Our results suggest that Benton’s visuospatial and visuoperceptual tests could be regarded as useful screeners of cognitive decline and global brain atrophy. However, longitudinal studies implying both measures are needed to assess their value in the prediction of dementia.

On the other hand, the correlations between neuropsychological tests and age and education were also significant in PD subjects in all cases, with a moderate to
large size effect, indicating that these results could probably be affected by these covariates. Indeed, age (Fjell et al., 2012; Williams-Gray et al., 2013) and education (Cohen et al., 2007) have been reported to have a differential role and an effect themselves on the progression of the disease. Whereas early onset PD patients show more rapid cognitive decline and poorer prognosis (Reid, Hely, Morris, Loy, & Holiday; 2011), advanced age and late PD onset has been associated with higher dementia risk (Dubois, Pillon, Sternic, Lhermitte, & Agid; 1990). Also, higher levels of education have been stated to modulate the cognitive performance in PD (Cohen et al., 2007); thus, this set of evidence suggests these two variables could exert a confounding effect in neuropsychological performance and that further research is needed to assess it.

In conclusion, our results indicate that global atrophy is related to visuospatial and visuoperceptual deficits detected by routine clinical practice in the assessment of cognitive performance in non-demented PD patients.

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Adaptación y validación del test Torre de Londres en mexicanos adultos mayores de 60 años