SHORT REPORT

Coding of movement direction and amplitude in Parkinson’s disease: are they differentially impaired (or unimportant)?

Dean L Jones, James G Phillips, John L Bradshaw, Robert Iansek, Judy A Bradshaw

Abstract
A recent study suggested that the preparation of movement direction, but not amplitude, may be selectively impaired by Parkinson’s disease (PD). The authors examined the reprogramming of direction only, amplitude only, and direction and amplitude together, and included a control condition in which neither parameter was reprogrammed. The findings suggested that neither direction nor amplitude coding was differentially impaired in PD. Thus the structures affected by PD may not be uniquely involved in specifying only the direction or the amplitude of future movements; these structures probably have more complex higher-level roles.

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Two reaction time (RT) studies by Pullman et al.² suggest that Parkinson’s disease (PD) may cause particular difficulty in the specification of the direction rather than the amplitude, of future movements. The same patients and procedures were used, measures were included of circulating plasma levodopa (the precursor to dopamine), and direction and amplitude were separately and independently varied in the two studies. The authors suggest that the preparation of movement direction, but not of amplitude, may be related to dopamine status. Unfortunately they failed to include two other conditions, where neither, or both, parameters were precued. While this requirement was largely met by Stelmach et al.,³ whose findings suggested normal specification of both parameters, there was no complete group analysis in this study. There is some evidence that PD may cause difficulty in specifying movement amplitude. Thus large amplitude movements in PD fall short of target,⁴ and are associated with abnormally short initial EMG bursts in agonists,⁵ (see Berardelli et al.) A final reason for questioning the findings of Pullman et al.² comes from recent studies of cell activity in the basal ganglia, which have emphasised that such responses may not be related to any simple fashion to movement parameters.⁶⁻⁷

We therefore compared the programming of direction and amplitude parameters on their own and together, and when neither was prespecified. The task involved reprogramming an ongoing movement.⁸ Compared with the methods of Pullman et al.² and Stelmach et al.,³ this paradigm allows us to examine both the preparation of a new response and the inhibition of the current motor programme. Additionally, the above conditions (either, both, or neither parameter prespecified) can be achieved without confounding the number of stimuli related to each response. In the RT studies described above,¹⁻³ some conditions involved just one visual cue, while others required two or more cues. This is an important issue, since PD may be associated with impaired maintenance of visual attention⁹ and poor control of eye movements.¹⁰ The present reprogramming design associates the same visual stimulus with all four task conditions.

Method
Subjects
Ten male PD subjects and 10 male control subjects participated voluntarily. All were right handed, and were screened for evidence of dementia, using the Mini-Mental State Examination (MMSE),¹² other neurological impairments and use of neuroleptic medication. Clinical data for the PD subjects are shown in the table, including ratings of symptom severity¹³ and disease progression.¹⁴

The PD group had a mean age of 66.6 years, a mean premorbid IQ of 113 according to the New Adult Reading Test, NART,¹⁵ and a mean educational level of 11.1 years. The control group had a mean age of 65.7 years, a mean premorbid IQ of 116, and a mean educational level of 10.2 years. According to one-way analyses of variance

Clinical data for PD subjects

<table>
<thead>
<tr>
<th>Number</th>
<th>Age</th>
<th>Wechsler ratings</th>
<th>Hoehn and Yahr</th>
<th>Levodopa (mg day⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>62</td>
<td>18</td>
<td>III</td>
<td>700</td>
</tr>
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<td>650</td>
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<tr>
<td>5</td>
<td>65</td>
<td>9</td>
<td>II</td>
<td>375</td>
</tr>
<tr>
<td>6</td>
<td>64</td>
<td>10</td>
<td>III</td>
<td>500</td>
</tr>
<tr>
<td>7</td>
<td>68</td>
<td>6</td>
<td>II</td>
<td>400</td>
</tr>
<tr>
<td>8</td>
<td>64</td>
<td>5</td>
<td>I (R)</td>
<td>900</td>
</tr>
<tr>
<td>9</td>
<td>78</td>
<td>6</td>
<td>II</td>
<td>1850</td>
</tr>
<tr>
<td>10</td>
<td>78</td>
<td>16</td>
<td>II</td>
<td>500</td>
</tr>
</tbody>
</table>

*(Stage of PD according to Hoehn and Yahr;¹⁴ (L: Left/R: Right side most affected).
(ANOVA), the groups did not significantly differ in age, $F(1, 18) < 1$; premorbid intelligence, $F(1, 18) = 1.14$, $p > 0.25$; or years of education, $F(1, 18) < 1$.

**Apparatus and procedure**

Testing took place at the most severe point in a patient's symptom cycle, that is, shortly before the next medication was taken. Each subject made responses on a board with a row of five buttons (1–5). The subjects sat so that the row of buttons lay along their sagittal plane. Adjacent buttons were separated by 30 mm (centre to centre) and each button was 13 mm in diameter. The buttons required a distance of 7 mm and a minimum force of 60 g to be depressed. A computer recorded the time data.

The subjects completed trials, each of which consisted of a series of 12 button presses, with their right hand as quickly and as accurately as possible. The movements between buttons alternated between being proximal-to-distal and distal-to-proximal. Each next button to be depressed was illuminated by a small red light emitting diode (in the base of the button) when the present button was released. Only one button was illuminated at any one time. On a typical trial, the subject depressed the starting button (button number 3) to begin, which caused the adjacent button (for example, button 4) to be illuminated. After pressing that newly illuminated button, the original button (3) again lit up. In the no change condition, this back and forth movement (between buttons 3 and 4) continued for the entire trial of 12 button presses. However, under the other three conditions of cued change, a new button was illuminated on either the fifth, sixth or seventh movement. The new button could be: for example, A) twice the amplitude but in the same direction as the usual movement (for example, button 5, when the subject expected to go from 3 to 4); B) in the reverse direction but of the same amplitude as the usual movement (for example, button 2, when the subject again expected to go from 3 to 4), or C) both twice the amplitude and in the reverse direction (for example, now button 1). On depression of the new button, the next one illuminated returned subjects to the button from which they had previously departed (3), so that they could continue alternating as before, for the remaining movements in the trial. Thus where both direction and amplitude were changed on the fifth movement, an example trial was as follows: 3–4–3–4–3–1–3–4–3–4–3–4.

On half the trials the orientation of the board was reversed, so that the number of proximally and distally directed movements was the same for all conditions. There were 12 trials per condition, and the 4 conditions were presented in random order. The mea-

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**Figure** Mean Movement Time (MT) before change and on change in movement for all reprogramming conditions, for Parkinson's disease and control subjects. Standard error bars are included.
sures of interest for each trial were the movement time (MT) for the change movement, compared with the MT for the movement immediately before the change, that is, the pre-change movement from button 3 to button 4. For trials with no change, a MT value was obtained for a movement in the sequence at the point corresponding to when a change would have occurred, had it been a change condition, that is, equiprobably at the fifth, sixth or seventh button tap.

Results
The extent of reprogramming is indicated by the increase in MT comparing pre-change and change conditions. To examine the extent of reprogramming we performed a mixed model 2 × 4 × 2 analysis of variance considering Group (PD, control), Condition (no change, direction only change, amplitude only change, direction and amplitude change) and Movement Type (pre-change, change movement). A difficulty reprogramming specific movement parameters would be indicated by disproportionate increases in MT (pre-change vs change) for patients compared with controls. The MT for pre-change and change movements in all conditions is presented in the figure.

Significant main effects indicated that PD subjects moved slower (PD, 388 ms vs controls 327 ms), \( F(1, 18) = 6.51, p < 0.05 \); that reprogramming occurred (pre-change, 267 ms vs change 449 ms), \( F(1, 18) = 129, p < 0.001 \); and that the specific conditions affected MT (no change, 281 ms; direction only, 378 ms; amplitude only, 366 ms; direction and amplitude, 407 ms), \( F(3, 54) = 66.94, p < 0.001 \). However, MT for each group was not differentially affected by the change in movement or by the condition, as indicated by non-significant interactions of Group by Condition \( F(3, 54) = 1.1, p > 0.05 \); and Group by Movement Type, \( F(1, 18) < 1 \). Note, from the figure, that both groups tend to slow down (that is, "fatigue") over time. There is, for instance, a slight increase in MT over the two positions even when no change of movement was required. Most importantly, however, this increase was not different for patients compared with controls, as shown by the lack of a significant interaction of Group by Condition by Movement Type, \( F(3, 54) < 1 \).

We can therefore conclude that PD and control subjects did not differ with respect to the effect on MT of the various changes in movement. PD subjects move slower, and can reprogram their movements, but do not appear to have problems reprogramming any specific movement parameter.

Discussion
In this experiment, where a reprogramming technique was employed, PD subjects, while overall slower than controls, did not experience disproportionate difficulty in altering the direction and/or amplitude characteristics of ongoing movements. These results, while consistent with those of Stelmach et al in the RT paradigm, apparently conflict with those of Pullman et al. In our study hand movements were visually guided (permitting PD subjects to use preferred, external rather than internally generated cues), while in the studies of Pullman et al the entire arm was hidden from view. Use of such visual information, however, is unlikely to explain the present results, since Stelmach et al also used movements where the hands and arms were out of sight, and their results generally concurred with the present findings.

Our data and those of Stelmach et al suggest two possible explanations. Firstly, the basal ganglia may be involved in coding both direction and amplitude, a position which is compatible with certain data on cell activity in these structures. Alternatively, the basal ganglia may not be primarily involved in specifying parameters of future movement, but may instead play some other role, probably at a higher level of motor function. This latter position has been taken by other, later, investigators of cell function in the basal ganglia. Clearly a broader view of basal ganglia function must be taken, to accommodate such complex concepts as motor programming, "internal" (or nonsensory) cueing, and sequential movement. According to this view, RT studies that demonstrate programming deficits in PD probably reflect problems in high level preparation processes, rather than problems in the simple specification of movement parameters. Note must be taken of the strong interconnections between the basal ganglia and the motor cortices. In particular, the supplementary motor area (SMA) probably participates in many higher motor functions, and is probably involved in motor programming and the dysfunction caused by PD.

We very much appreciate the helpful comments of Dr Malcolm Horne on our findings. We also sincerely thank Dr David Andrewes and the Parkinson's Disease Association of Victoria for access to additional patients, and Drs. Dick, Bob Wood, Mike Durham, Frank Devlin, and Truong Nguyen for designing and maintaining the apparatus and software. This work has been presented in part at a Meeting of the International Neuropsychological Society, Gold Coast, Australia, 1991.


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