Motor response to apomorphine and levodopa in asymmetric Parkinson’s disease

M Rodriguez, G Lera, J Vaamonde, M R Luquin, J A Obeso

Abstract
The motor responses of 14 patients with Parkinson’s disease (six previously untreated and eight chronically receiving levodopa) with pronounced asymmetry in the severity of motor signs between the left and right sides of the body were studied. The effects of a short (60 minutes) and a long (16–22 hours) intravenous levodopa infusion as well as of subcutaneous apomorphine (1–6 mg bolus) were assessed. Four different tapping tests were used to measure motor function. For all pharmacological tests, the more affected side showed a shorter response duration, increased latency, and greater response magnitude than the less affected side. These differences were more pronounced in those patients receiving chronic levodopa treatment. As apomorphine is not dependent on dopamine storage capacity, these findings suggest that postsynaptic mechanisms play an important part in the origin of motor fluctuations in Parkinson’s disease.

(Please note: The abstract and full text have been shortened for clarity and readability.)

Daily fluctuations in motor performance are a common and troublesome complication of chronic treatment with levodopa for Parkinson’s disease. Chase’s group showed that the duration of the motor response (the “on” period) to an acute levodopa challenge is inversely correlated with disease severity. It was proposed that progressive degeneration of the nigrostriatal pathway leads to a decreased storage capacity for the dopamine synthesized from exogenous levodopa.1-3 Thus the basic pathophysiological mechanism underlying motor fluctuations in Parkinson’s disease could be a presynaptic defect in buffering the oscillations in plasma levodopa concentrations.4 Reduced storage capacity might be associated with higher synaptic dopamine concentrations, thus explaining the steeper dose–response curve reported in more severely affected patients.4 The relation between severity of disease and shortening of the response to levodopa has been less clear, however, in other studies5-7 throwing some doubt on the fundamental role of defective striatal dopamine storage capacity in the origin of motor fluctuations. The existence of postsynaptic dopaminergic abnormalities in severe Parkinson’s disease has also been considered in view of the difficulty in keeping patients constantly “on” during prolonged levodopa infusions8-9 and continuous treatment with dopamine agonists such as lisuride, apomorphine, and (+)-4-propyl-9-hydroxy-naphthoxazine (+PHNO).10-12

In this study we report the response to subcutaneous apomorphine boluses and prolonged and short infusions of levodopa in 14 patients with asymmetric Parkinson’s disease. This approach allowed us to assess and compare the motor response induced by a presynaptically and postsynaptically acting drug in relation to disease severity without the problems in interpretation derived from peripheral pharmacokinetic modifications.

Patients and methods

PATIENTS
Fourteen patients (three females and 11 males) with asymmetric Parkinson’s disease were studied. Parkinsonian signs were worse on the left side in 10 patients and the right side in more affected in four. The non-dominant side was more severely parkinsonian in eight patients. Six patients had never been treated with levodopa or any other antiparkinsonian drug (de novo) and eight showed daily motor fluctuations (“treated”) after a mean 4.9 (range 2-10) years of levodopa treatment. The mean daily dose of levodopa (plus a decarboxylase inhibitor) was 856±2 (range 600–900) mg for the eight patients on treatment. Two patients were also on treatment with bromocriptine (20 and 25 mg daily) and another was taking amantadine (300 mg daily). Both drugs were discontinued seven days before the study. The mean duration of illness was 4.5 (range 1–4) years for the “de novo” group, and 6.5 (range 2–10) years for the treated group. Before the study the mean score on the unified Parkinson’s disease rating scale (UPDRS) was 37.7 (range 26–56) for the entire group; “de novo” patients had a mean score of 31 (SD 4.7) (range 26–36) and treated patients scored 45.6 (7.9) (range 28–59) when “off”.

PHARMACOLOGICAL TESTS
Patients were in hospital during the study and received domperidone (60 mg daily) from three to five days before until the conclusion of the study. All patients were informed of the purpose and potential risks of the study. Levodopa was stopped the night before the tests in the eight patients on treatment. These patients resumed their normal therapeutic.
regimen immediately after the end of each test. The total period in hospital was 7–10 days.

Levodopa (Chiesi Farmaceutica, Italy) was diluted in sterile saline (9%) to a final concentration of 1 mg/ml and a pH of 5.4. Intravenous infusions were given through a forearm vein on the less affected side through a standard catheter. Apomorphine (10 mg/ml, Woelm, Germany) was given subcutaneously in the arm.

Pharmacological tests were performed in the morning in the following sequences: (1) The minimal effective dose (MED) of subcutaneous apomorphine. The MED was defined as the dose of apomorphine capable of reducing motor disability (unified Parkinson’s disease rating scale or tapping tests) by at least 50% during 10–15 minutes. This dose was determined for each patient on successive days starting with 0.50 mg of apomorphine and increasing the dose to reach the MED. The mean MED of apomorphine was 2.25 (SD 0.5) mg (range 1–3 mg). (2) Twice the MED of apomorphine This was given subcutaneously to each patient. (3) Short levodopa infusion Intravenous levodopa was infused at a constant rate of 250 mg/h until a motor improvement of 50% in both sides was seen. The infusion was stopped at that moment. Carbidopa (25 mg orally) was given three hours and one hour before starting the infusion. (4) Prolonged levodopa infusion A levodopa infusion at a rate of 60–100 mg/h was given for 16–22 hours. The infusion rate was adjusted to ensure that the patient remained “on” (bilaterally) until sleep time. Levodopa infusion was maintained subsequently during the night. Three patients were not “on” the next morning despite having received the levodopa infusion during the night. In these instances, the levodopa infusion rate was again increased and adjusted to guarantee the “on” state for a minimum of three hours before stopping the infusion. Carbidopa (25 mg orally) was given every three hours until the end of the infusion.

ASSESSMENT
Motor function was assessed separately in the right and left sides of the body by four motor tests. Patients were sitting comfortably on a chair maintaining the trunk in an upright position and as vertical as possible. The examiner counted the number of valid movements made in 30 seconds. The tests consisted of: (a) wrist tapping (patients touched repetitively a manual digital counter placed on a table); (b) finger counting (separation of the index finger from the thumb by a minimum of 3 cm); (c) arm tapping (patients tapped with the hand two points placed 30 cm apart in front of the subject); (d) foot tapping (repetitive foot raising in the vertical axis by at least 20 cm). These four tests were applied to monitor the motor response to levodopa and apomorphine. The motor state was continuously assessed by the examiner. The motor tests were performed at baseline and every 10 minutes after giving drugs, or at any time during the experiment when the observer believed that an important modification in the motor state had occurred.

The following items were analysed for each patient: (a) latency to the beginning of the motor response (defined as a minimum improvement in motor capability of 25%) in any of the motor tests; (b) The magnitude of the response (difference in motor scores between “off” and “on”); (c) The duration of the benefit (“on”) defined as the interval between the beginning of the response and the return to baseline of the motor scores after giving apomorphine. For the levodopa infusions, “on” duration was counted as the time elapsed between cessation of the infusion and return to the baseline score.

A paired Student’s t test was used to compare the more and less affected sides. A t test for unpaired comparisons was applied to analyse differences between “de novo” and treated patients. Values are expressed as means (1 SD).

Results
Each of the four tests used to monitor the motor responses followed a similar pattern and all discriminated significantly (p < 0.05) between the more and less affected sides in basal conditions (fig 1, A and B; fig 2). The wrist tapping test was chosen for presentation of the results to simplify comparison between sides.

MOTOR RESPONSE AND DISEASE SEVERITY
(n = 14)
Latency
The “on” latency (table 1) was significantly greater in the more affected side in response
Table 1. Comparison of the motor response to apomorphine and levodopa in 14 patients with asymmetric Parkinson’s disease

<table>
<thead>
<tr>
<th>Pharmacological test</th>
<th>More affected</th>
<th>Less affected</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Latency &quot;on&quot; (min)</td>
<td>Duration &quot;on&quot; (min)</td>
</tr>
<tr>
<td>Apomorphine (MED)</td>
<td>22.3(9)</td>
<td>54.9(30.1)</td>
</tr>
<tr>
<td>Apomorphine (twice MED)</td>
<td>16.6(6.7)</td>
<td>74.9(34.2)</td>
</tr>
<tr>
<td>Short levodopa infusion</td>
<td>28.4(14.8)</td>
<td>150.2(83.9)</td>
</tr>
<tr>
<td>Long levodopa infusion</td>
<td>—</td>
<td>130(88.8)†</td>
</tr>
</tbody>
</table>

Values are means (1 SD). *p < 0.05; **p < 0.01; ***p < 0.001 (differences between sides). †Detailed data for each test in each patient is available upon request. ‡Values express the duration of the "on" after stopping the infusion.

Figure 2. Motor response elicited by subcutaneous apomorphine (mean dose 2.25±0.5 mg) in 14 patients with asymmetric Parkinson’s disease. Each point represents the mean value for each tapping test. Apomorphine was given at time 0. The response had a greater magnitude but reduced duration in the most affected side.

Table 2. Duration (min) of the motor response in asymmetric de novo patients (n = 6) and patients treated chronically with levodopa (n = 8)

<table>
<thead>
<tr>
<th></th>
<th>More affected</th>
<th>Less affected</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>De novo</td>
<td>Treated</td>
</tr>
<tr>
<td>Severity: Tapping test</td>
<td>53(24)</td>
<td>43(26.2)</td>
</tr>
<tr>
<td>Pharmacological tests</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Apomorphine (MED)</td>
<td>74.1(37)</td>
<td>40.5(11.1)</td>
</tr>
<tr>
<td>Apomorphine (twice MED)</td>
<td>96.6(34)</td>
<td>52(17.5)*</td>
</tr>
<tr>
<td>Short levodopa infusion</td>
<td>203.1(86.7)</td>
<td>110.5(39.1)*</td>
</tr>
<tr>
<td>Long levodopa infusion</td>
<td>226.6(80.8)</td>
<td>93.7(62.4)*</td>
</tr>
</tbody>
</table>

Values are means (1 SD). M.E.D = minimal effective dose. *p < 0.05; **p < 0.01. †Wrist tapping test. Patients touched repetitively for 30 seconds a manual digital counter placed on a table in front of them.

to the MED of apomorphine (p < 0.001). There was no significant difference between sides after giving twice the MED of apomorphine, although the actual value was higher in the more affected side. The short levodopa infusion induced an earlier response in the more affected side (table 1). The slow infusion rate used at the beginning of the prolonged levodopa infusion precluded the measurement of "on" latency for this test.

Magnitude of the response
In all patients the magnitude of the response was greater in the more affected side (p < 0.01) in response to both apomorphine and levodopa (fig 1 A and B). This difference was significant (p < 0.01) for all pharmacological tests (fig 2).

Duration
In every patient, all four pharmacological tests described consistently showed a similar pattern—that is, the duration was longer in the less affected side than in the more affected side. Individual data are not shown for the sake of brevity but are available on personal request.

Apomorphine—The mean duration of the response (54.9 (SD 30.1) min) in the more affected body side after subcutaneous apomorphine (MED) injection was significantly reduced (p < 0.001) compared with the less affected side (90.4 (SD 44.9) min) (table 1; fig 2). Twice the MED of apomorphine (2.6 mg) elicited a motor response with a mean duration of 74.9 (SD 34.2) minutes in the more severely affected side and of 120.7 (SD 61.3) minutes in the less affected side (p < 0.001; table 1).

Short levodopa infusion—The mean duration of the motor response was significantly shorter (p < 0.001) in the more affected side (150.2 (SD 83.9) min) compared with the less severe side (199.7 (SD 71.4) min) (table 1).

Prolonged levodopa infusion—The duration of the antiparkinsonian action of levodopa after discontinuation of the infusion lasted for 130 (SD 88.8) minutes. In the more affected side and 191.8 (SD 93.4) minutes in the less affected side. This difference was statistically significant (p < 0.01; table 1).

De novo v treated patients
The magnitude of the response was greater (p < 0.01) in the more affected side of both de novo and treated patients. The duration of the motor response to apomorphine boluses and levodopa infusions was significantly longer in the less affected side in both de novo and treated patients (table 2). Comparison of the more affected side of de novo and treated patients did not reveal statistically significant differences in the tapping tests (table 2). Similar results were found for the less affected side suggesting a similar degree of parkinsonism for each side in both groups of patients. The magnitude of the response was also similar (fig 2) but the duration of the response to the pharmacological tests was shorter in treated patients (table 2; fig 3).
Discussion

Studies of pathology and PET scan indicate that clinical asymmetry of motor signs in Parkinson's disease reflects differences in the degree of degeneration of the substantia nigra and striatal dopamine loss. The study of patients with asymmetric disease allowed us to compare the effect of different severity of disease (dopaminergic denervation) upon the motor response in the same subjects and exclude confounding factors such as duration of levodopa treatment, duration of disease and problems of peripheral kinetics. We found a highly significant correlation between severity of parkinsonism and latency, magnitude, and duration of the motor response. Thus, the more affected side showed a greater magnitude of improvement and a shorter response duration. This relation occurred and was maintained when the same patients were challenged with apomorphine, a short levodopa infusion, or a long levodopa infusion. We therefore confirm the results of previous studies with levodopa infusions as a test regarding the shortening of the response and increasing magnitude with greater severity of disease and extend the findings to apomorphine.

The results presented here are at variance with the findings of Kempster et al who described a similar response duration to levodopa (250/25 mg orally) in both sides of patients with asymmetric disease. Differences in the method used to quantify the motor response during the pharmacological tests and in the severity of the patients may account for this discrepancy. Admittedly, measuring drug-induced responses in milder parkinsonian patients is not an easy task. We thus used four different motor tests, which assessed movement capacity in different body parts, and consistently found a shortening of the response in the more affected body side. These tapping tests are well known to be more discriminatory than the general scales usually applied to evaluate Parkinson's disease. Furthermore, it is unlikely that variations in the assessment of the duration of the motor response would always go in the same direction—that is, the more affected side consistently showing a shorter response in every patient and for all pharmacological tests. We did not take into consideration other clinical signs, such as tremor or dyskinesias (in treated patients), because they are very sensitive (and extremely variable) to uncontrollable factors such as anxiety, tiredness, and degree of mental and physical activity. In conclusion, we do not believe that our method was either insensitive for the purpose of the study or biased towards underestimating the duration of response in the more affected side. Indeed, before the study we had expected to find similar results with levodopa to those of Kempster et al and differences in the response to apomorphine, reflecting the importance of postsynaptic mechanisms in the pathophysiology of motor fluctuations. It may be that in the more severe patients studied by Kempster et al the response to levodopa, which presumably reflects mainly presynaptic mechanisms, was more similar due to a lesser degree of asymmetry. Whether or not this interpretation is correct may be tested in future studies.

The major finding of this study is that the correlation between severity of disease and motor response was similar for apomorphine and levodopa (both short and prolonged infusions). The decline in motor improvement after stopping the long levodopa infusion may reflect presynaptic dopamine storage. It follows that a reduced storage capacity should be accompanied by a shorter response after discontinuation of the infusion, as in fact occurred in this and previous studies. On the other hand, decreased storage capacity would lead to enhanced dopamine dispositions after a bolus of levodopa, which should provoke a greater improvement and also a longer response. This was not the case in the patients of Gancher et al or in our patients, suggesting that dopamine storage capacity is not the only factor involved in the origin of motor fluctuations. Indeed, our results with apomorphine strongly indicate a paramount role of postsynaptic mechanisms in the origin of motor fluctuations in Parkinson's disease. Apomorphine is a direct acting dopamine agonist (D-1 and D-2 receptors) and is capable, by itself, of reversing parkinsonism. Apomorphine does not require biotransformation to be active and is not stored in the brain. Accordingly, the response to apomorphine essentially reflects the degree of striatal dopaminergic responsiveness mainly as a consequence of denervation. The nature and intimate characteristics of the changes induced by the loss of nigrostriatal dopaminergic activity are not clear. In untreated patients with Parkinson's disease there seems to be hypersensitivity of D-2 receptors in keeping with the findings in animals with nigrostriatal lesions. Dopamine (D-2) receptor density remains within normal values despite disease progression. The data concerning D-1 receptors are still highly controversial. Different studies have reported normal, reduced, or increased activity and number of D-1 receptors in the striatum of patients with Parkinson's disease. One suggested possibility is that motor complications could be

Figure 3 Comparison of the improvement elicited by an intravenous bolus of apomorphine on the more and less affected sides of eight treated patients (TP) and six de novo (DN) patients. The response is shortest in the more affected side of TP and longest in the less affected side of DN patients.
more related to pathological modifications in the affinity state (high-low) of dopamine receptors.19 20 In this regard, Hornykiewicz’s group has recently described sensitisation of D-1 striatal receptors in rats with a 6-hydroxydopamine-induced lesion, monkeys treated with 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine, and in eight patients with Parkinson’s disease.21,22 This alteration is not believed to be simply due to up-regulation of D-1 receptors but to alteration of the signal transduction mechanism controlling adenyl cyclase activity.25 26 It is conceivable therefore, that denervation-induced pathological modifications of the affinity state and also of the relative proportion of D-1/D-2 striatal receptors could be directly implicated in the origin of the short duration response in Parkinson’s disease. It was noted that acute apomorphine and levodopa treatment was accompanied by a short duration response in both de novo and treated patients. This finding indicates that the mechanisms underlying the short duration response and by extension, motor fluctuations, are qualitatively similar in both groups.27 28 Chronic levodopa treatment probably aggravates this phenomenon, particularly by reducing the duration of the response.

In conclusion, the findings suggest that the degree of substantia nigra lesion and the consequent dopaminergic deficit govern the characteristics of the response to acute challenge with potent dopaminergic drugs. The main mechanism subserving daily motor fluctuations in Parkinson’s disease may not be defective storage of dopamine (from exogenous levodopa) but modifications in striatal responsiveness.

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