Usefulness of pallidotomy in advanced Parkinson’s disease

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Abstract
Objective—The combined effect of putaminal pallidotomy and optimal medical treatment was assessed in 22 patients with levodopa sensitive Parkinson’s disease.

Methods—Timed motor tests, video recordings, and computer assisted optoelectronic movement analysis were used for serial hourly assessments performed preoperatively and four and 12 months after operation. Tests were made while patients were on optimal medical therapy.

Results—There were no serious adverse events of surgery. Two of the 22 patients could not complete all the tests after operation. The proportion of dyskinesia periods decreased in the 20 patients and there was a proportional increase in normal or fairly normal occasions. “Off” periods were not significantly affected. In 12 of 13 patients with limb dyskinesia this symptom was completely abolished in the contralateral limbs There was also some degree of improvement axially and ipsilaterally. Tremor was moderately improved contralaterally. Bradykinesia remained unchanged. Results at 12 months follow up were similar to those at four months.

Conclusion—Pallidotomy produced a pronounced positive effect on dyskinesia and a moderate effect on tremor. Bradykinesia was not affected. Posteroventral pallidotomy may be useful in patients with Parkinson’s disease who have severe motor fluctuations and may allow an increase in levodopa dose to alleviate bradykinesia in “off” states.

Materials and methods

Patients
In a prospective, consecutive series 21 right handed patients and one left handed patient were included according to the following inclusion criteria: (a) idiopathic Parkinson’s disease; (b) documented positive early response to levodopa; (c) insufficient response to pharmacological treatment or severe adverse effects, primarily motor fluctuations; (d) age < 80 years. Medication was adjusted to maximise the antiparkinsonian efficacy and minimise side effects. Patients had been on stable doses of levodopa for at least two months before entry into the study.

Patients with secondary parkinsonism, multisystem degeneration, prior brain operations (except for Parkinson’s disease), and other degenerative or vascular brain diseases were excluded. Patients with advanced cortical atrophy and hydrocephalus on CT were excluded. In one of the 22 patients an asymptomatic preputinate cyst was found on preoperative MRI and another patient had an asymptomtomatic small calcified parasagittal left sided meningioma at the level of the bregma.

Although some of the patients had experienced transient toxic hallucinations, no patient was psychotic or depressive at the preoperative
investigations, according to the Unified Parkinson’s Disease Rating Scale (UPDRS) part I.30

The 22 patients (16 men, six women) had a mean age of 63·8 (range 43 to 78) years. Duration of Parkinson’s disease was 14·8 (range 7 to 22) years and duration of motor fluctuations 7 (range 2 to 11) years. The five patients without fluctuations had a shorter duration (mean 10 years) of disease than those with fluctuations (mean 16 years). The median score of the Hoehn and Yahr stage36 (in “best on”) was 3·0. The mean score (in “best on”) of activities of daily living (Schwab and England)37 was 72 (range 60 to 80)%.

The following laboratory tests were performed to rule out other serious diseases: full blood count, erythrocyte sedimentation rate, tests of hepatic, renal, and thyroid function, blood glucose, serum electrolytes, B12/folic acid, protein electrophoresis, CSF protein, cell count, and ECG.

All patients were on levodopa therapy. Slow release preparations (Madopar HBS or Sinemet SR) alone or in combination with standard levodopa were used in 16 patients. Many patients were on soluble levodopa for rapid effect in off stages or as a booster dose in the morning. Bromocriptine and selegiline were taken by 18 and 11 patients respectively. Two patients had apomorphine in subcutaneous injections and four other patients had tried apomorphine or levodopa infusions without benefit. One patient took anticholinergic drugs for tremor and three patients had amantadine. Three patients had prior experience with protein modulated diets. One patient had had a thalamotomy for tremor seven years before the study and another patient a pallidotomy for dystonia and tremor one year previously. Both patients underwent PVP in the contralateral hemisphere. Postoperatively, the patients were allowed to adjust their Parkinson’s disease medication if necessary.

PREOPERATIVE AND POSTOPERATIVE INVESTIGATION

At the baseline visit, clinical characteristics were assessed through a neurological examination and careful history, together with an ophthalmological examination including visual fields and acuity. Magnetic cortex stimulation was carried out to screen for cortical lesions.11,12 A short psychometric test38 was administered. The mean score was 34·0 (SD 2·7) points (maximum score 40 points). None of the patients fulfilled the criteria for dementia.

Tests for motor performance were carried out during one day, from 8 00 am to 7 00 pm, at baseline and at four and 12 months postoperatively. Testing was carried out by a trained physiotherapist, who also checked that the patients followed their usual stable drug regimen during the test day.

The study protocol was approved by the local ethics committee. The patients gave their consent after information about the operation and possible adverse effects.

VARIABLES OF SURGICAL OUTCOME

Timed motor tests

Two tests for dexterity of each hand (pronation-supination test and peg board test) and a gait velocity test (3 × 10 m with turns) were performed. The three tests were repeated twice, at 11 00 am and 3 00 pm and the mean time for each test was calculated.

Video recordings

These were performed every other hour (six times) from 8 00 am to 6 00 pm. The following items from the UPDRS part III and IV30 were evaluated: resting tremor, dyskinesia (disability), rapid finger movements, rapid alternating hand movements, rising from a chair, posture, and gait.

The patients’ performance recordings were independently rated by one of us (FJ) and by a physiotherapist who had not taken part in the recordings or examinations. For each limb and axial structures, UPDRS items were scored 0 (absent) to 100 (most severe) using a visual analogue scale (VAS) without predefined steps.44,45 The VAS scale is both sensitive and reproducible and has equal power compared with a verbal rating scale.46 The VAS is an ordinal scale and each score was transferred into one of 10 groups (0–9, 10–19, 20–29…90–100). Every patient was recorded on video six times each test day and the median and maximal values were calculated for each UPDRS item.

Motor fluctuations

The fluctuations were calculated as percentage of the day and rated in the following categories: “on” time (good or fairly good mobility), “on plus” time (mobile with dyskinesiae) and “off” time (slow or absent mobility). These observations were made every 30 minutes from 8 00 am to 7 00 pm by the same physiotherapist who performed the other motor tests.

The posturolocomotion manual (PLM) test

This test has been designed to objectively quantify selected aspects of a subject’s motor performance during a standardised motor task, and can be performed on all freely moving patients with Parkinson’s disease of Hoehn and Yahr grade 4 or less.53 The rationale is intended to accurately determine the capacity of four different aspects of motor performance: the postural (P), the locomotive (L), and the manual (M) components, and overall movement time (MT). To this end subjects were fitted with six retroreflective markers positioned on the head, shoulder, elbow, hip, and both feet. The movements of these markers were recorded by a specialised high accuracy videocamera based system (MacReflex, Qualisys AB, Partille, Sweden). The patients were instructed to move an object as rapidly as possible from a marked spot on the floor 1·5 m forwards and place it on a shelf, individually adjusted to the level of the shoulder. These movements were repeated as rapidly as possible during a 30 second test period, and each side was tested. This test sequence was
repeated every other hour from 9.00 am to 7.00 pm; in total six times over the 10 hour period. At the subsequent analyses, four different durations of the standardised movement were determined; the movement time from lifting the object to placing it on the shelf (MT); the time taken to raise the body to an upright position (P phase), the time for the actual forward body movement (L phase), and the time taken to lift the arm (M phase). Also the number of lifts performed during each of 30 second test were determined and a simultaneity index (SI; defined as P + L + M/MT), which is intended to quantify how well the PLM phases are coordinated into a smooth motor act, were computed. For the present report, only the estimates of total MT and SI were evaluated.

Surgical Procedure

Fifteen PVPs were carried out in the left hemisphere and nine in the right. Two patients were reoperated on as described below. All patients underwent a stereotactic CT or surgery, or stereotactic MRI study, one to three days before surgery, using the Laitinen stereoadapter. The length of the intercommissural line was measured and the standard pallidal target was defined 2 mm in front of the midcommissural point, 5–6 mm below the intercommissural line, and 20–22 mm lateral to the midline of the third ventricle. This point represented the zero point—that is, the most ventral point of the target area to be stimulated and eventually coagulated.

To maximise the symptoms at the time of surgery, all parkinsonian medication was stopped at least 12 hours before pallidotomy. For surgery, the stereoadapter was positioned on the head while the patient was in a semisitting position on the operating table. Around it, Laitinen’s stereoadapter was mounted rigidly on the skull under local anaesthesia. Thereafter the patient was laid supine on the table. The CT/MRI coordinates of the pallidal target were transferred to the stereoadapter and the stereoadapter was removed.

Through a frontal burr hole lying 2–3 cm from the midline, slightly in front of the coronal suture, a 1.8 mm thick monopolar electrode with a 2 mm long non-insulated tip was introduced towards the target under impedance monitoring. Electrical stimulations were conducted with 6 Hz, 10 mA and 60 Hz, 5 mA at 2 mm intervals, between the ventral most edge of the pallidum, represented by the zero target point, and the level of the intercommissural line estimated to lie 6 mm dorsal to that point. During stimulation, care was taken to detect any capsular or visual response. In addition, speech and articulation were carefully monitored. The patient was asked to perform bicycling movements, alternating hand movements, and finger agility movements. The orientation and short term memory of the patient were checked. These same tasks were also performed during coagulation at each of the points.

If stimulation did not give rise to any unde-

sirable reactions an incremental radiofrequency lesion was produced by heat coagulation at 75–83°C during 30–60 seconds in steps of 2 mm between the zero point and a point 6–8 mm dorsal. After being applied in 15 patients, this technique was slightly changed in the remaining seven patients: the electrode was first stopped 6 mm dorsal to the zero target point and then stimulation and coagulation were conducted in a dorsal to ventral direction. In all patients a stereotactic CT or MRI study was conducted 4–10 months after surgery. The scanning plane was made identical to the preoperative radiological study, both for the CT and MRI. Thus the lesion site could be assessed in relation to the preoperative target point—that is, to the reference points of the third ventricle (the intercommissural line, midcommissural point, and mid sagittal plane of the third ventricle).

Statistical Methods

Preoperative and postoperative variables of the timed tests were evaluated by a paired t test. Ordinal values (for example, from the video recording), were analysed using the Wilcoxon signed rank test. PLM tests were analysed with the Mann-Whitney U test and the Kruskal-Wallis test. Repeated measures analysis of variance (ANOVA) was used to determine the significance of changes in motor fluctuations over time. A significance level of P ≥ 0.05 was used.

Results

Adverse Events During Operation and Follow Up

In patients Nos 3, 15, and 17 capsular responses at stimulation led to a replacement or slight withdrawal of the electrode and to a coagulation of a smaller volume than initially intended. In 22 patients had any corticospinal dysfunction at examination or on motor cortex stimulation four or 12 months postoperatively.

In nine patients (40%), visual responses were obtained at stimulation between the zero target (most ventral point) and up to 4 mm above the zero target. In these patients, either the electrode was withdrawn and coagulation was made more dorsally (in patients in whom the electrode was first introduced to the zero target), or the coagulation and surgery were interrupted (in patients in whom stimulation and coagulation were started at points dorsal to the zero target). In either case, the lesion was generally placed slightly more dorsally or was smaller than initially planned. In patients Nos 20 and 22 slight confusion or dysarthria occurred during and directly after coagulation respectively. Steroids were given and the symptoms resolved after a few hours.

At the four month follow up, two patients could not be tested for the following reasons: patient No 1 with some cognitive deficit preoperatively developed psychotic symptoms a few months after the operation and was excluded from further analyses. Patient No 12 with severe dysarthria and dysphonia preopera-
Figure 1  Thin slice MRI showing the appearance of the lesion in the right pallidum one day after surgery (A) and four months after surgery (B arrow).

Patient No 17 developed acute psychosis two months after operation but recovered rapidly on small doses of clozapine without changing Parkinson's disease medication. At 12 months, results from another four patients were missing: patient No 11 developed a malignant glioma eight months after surgery, patient No 13 had a serious traffic injury when riding a motor cycle (!), patient No 2 had cardiac disease, and patient No 10 was not available for administrative reasons. Only in patient No 12 could the symptoms be directly attributed to the operation. Patient No 13 had a permanent but small and subclinical visual scotoma postoperatively.

REOPERATIONS
Dyskinesia recurred postoperatively in patient No 4 and tremor in patient No 9 and had not improved at the evaluation four months postoperatively. Furthermore, their lesions were considered to be small on CT or MRI. Surgery was repeated on the same side 10 and 11 months later. Both patients were evaluated at four months after the first operation and at
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Table 1  Fluctuations in motor performance calculated as percentage of the day

<table>
<thead>
<tr>
<th></th>
<th>Baseline</th>
<th>4 months</th>
<th>12 months</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean (%)</td>
<td>(95% CI)</td>
<td>Mean (%)</td>
</tr>
<tr>
<td>&quot;Off&quot; time</td>
<td>23.8</td>
<td>(14.1–33.5)</td>
<td>27.8</td>
</tr>
<tr>
<td>&quot;On&quot; time</td>
<td>19.5</td>
<td>(10.1–26.6)</td>
<td>26.4</td>
</tr>
<tr>
<td>&quot;On +*&quot; time</td>
<td>27.8</td>
<td>(14.5–70.5)</td>
<td>25.8</td>
</tr>
</tbody>
</table>

Observations were every 30 minutes from 8.00 am to 7.00 pm. "On" = good or fairly good mobility, "On plus" = mobile with dyskinesia, "Off" = slow or absent mobility.

Table 2  Median and maximum values of the six video recordings each test day on the operated side (contralateral to pallidotomy)

<table>
<thead>
<tr>
<th></th>
<th>Postoperatively</th>
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<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Baseline</td>
<td>4 months</td>
<td>P value</td>
<td>12 months</td>
</tr>
<tr>
<td>Limb dyskinesia</td>
<td>13</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>15</td>
<td>10</td>
<td>0.008</td>
<td>0.001</td>
<td>10</td>
</tr>
<tr>
<td>Maximum</td>
<td>30</td>
<td>10</td>
<td></td>
<td></td>
<td>10</td>
</tr>
<tr>
<td>Axial dyskinesia</td>
<td>12</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>15</td>
<td>10</td>
<td>0.031</td>
<td>0.001</td>
<td>10</td>
</tr>
<tr>
<td>Maximum</td>
<td>40</td>
<td>20</td>
<td></td>
<td></td>
<td>25</td>
</tr>
<tr>
<td>Tremor at rest</td>
<td>10</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>25</td>
<td>10</td>
<td>0.148</td>
<td>0.004</td>
<td>10</td>
</tr>
<tr>
<td>Maximum</td>
<td>55</td>
<td>30</td>
<td></td>
<td></td>
<td>10</td>
</tr>
<tr>
<td>Dexterity in hands</td>
<td>19</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>52.5</td>
<td>65</td>
<td>0.228</td>
<td>0.018</td>
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<tr>
<td>Maximum</td>
<td>80</td>
<td>75</td>
<td>0.307</td>
<td>0.086</td>
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<tr>
<td>Rapid alternating movements of the hands</td>
<td>19</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>50</td>
<td>50</td>
<td>0.908</td>
<td>0.549</td>
<td></td>
</tr>
<tr>
<td>Maximum</td>
<td>70</td>
<td>60</td>
<td>0.006</td>
<td>0.041</td>
<td></td>
</tr>
<tr>
<td>Arising from chair</td>
<td>11</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>10</td>
<td>10</td>
<td>0.945</td>
<td>0.125</td>
<td></td>
</tr>
<tr>
<td>Maximum</td>
<td>40</td>
<td>20</td>
<td>0.082</td>
<td>0.898</td>
<td></td>
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<tr>
<td>Posture</td>
<td>19</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>20</td>
<td>20</td>
<td>0.654</td>
<td>0.500</td>
<td></td>
</tr>
<tr>
<td>Maximum</td>
<td>40</td>
<td>30</td>
<td>0.178</td>
<td>0.322</td>
<td></td>
</tr>
<tr>
<td>Gait</td>
<td>19</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>20</td>
<td>22.5</td>
<td>0.628</td>
<td>0.758</td>
<td></td>
</tr>
<tr>
<td>Maximum</td>
<td>50</td>
<td>50</td>
<td>0.004</td>
<td>0.245</td>
<td></td>
</tr>
</tbody>
</table>

n Refers to number of patients with the symptom.

Table 3  Median and maximum values of the six video recordings on the non-operated side (ipsilateral)

<table>
<thead>
<tr>
<th></th>
<th>Postoperatively</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Baseline</td>
<td>4 months</td>
<td>P value</td>
<td>12 months</td>
</tr>
<tr>
<td>Limb dyskinesia</td>
<td>13</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>15</td>
<td>10</td>
<td>0.001</td>
<td>0.001</td>
<td>10</td>
</tr>
<tr>
<td>Maximum</td>
<td>15</td>
<td>10</td>
<td></td>
<td></td>
<td>10</td>
</tr>
<tr>
<td>Tremor at rest</td>
<td>4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>17.5</td>
<td>15</td>
<td></td>
<td></td>
<td>20</td>
</tr>
<tr>
<td>Maximum</td>
<td>40</td>
<td>40</td>
<td></td>
<td></td>
<td>20</td>
</tr>
<tr>
<td>Dexterity</td>
<td>19</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>65</td>
<td>57.5</td>
<td>0.994</td>
<td>0.535</td>
<td></td>
</tr>
<tr>
<td>Maximum</td>
<td>80</td>
<td>70</td>
<td>0.450</td>
<td>0.027</td>
<td></td>
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<tr>
<td>Rapid alternating movements of the hands</td>
<td>19</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>50</td>
<td>50</td>
<td>0.413</td>
<td>0.510</td>
<td></td>
</tr>
<tr>
<td>Maximum</td>
<td>60</td>
<td>50</td>
<td>0.094</td>
<td>0.063</td>
<td></td>
</tr>
</tbody>
</table>

n Refers to number of patients with the symptom.

The lesion, which was estimated to be 4–6 mm in transversal diameter with a rostrocaudal length of 6–8 mm, was <95% CI (46·8–80·2 mm³) in nine patients (41%) and >95% CI (>101·8 mm³) in four patients (19%). There was no clear correlation between the size of the coagulated area and the size of the final lesion (fig 1A and B). The estimated location of the lesion coincided with the coagulated pallidal area in all patients. In nine patients the coagulated area lay more dorsal to the intended zero point target (ventralmost edge of pallidum) because of an optic response at stimulation as reported previously.

MEDICATION
Daily doses of levodopa during the study were: baseline, 952 (SD 365) mg; four months, 913 (SD 377) mg; 12 months, 931 (SD 373) mg. The corresponding doses of bromocriptine were: baseline, 18·2 (SD 14·3) mg; four months, 19·7 (SD 15·1) mg; 12 months, 21·4 (SD 15·4) mg. There were no significant changes in these treatments during the study period. Selegiline and the other Parkinson’s disease medications were also unchanged.

FOLLOW UP AT FOUR MONTHS
Motor fluctuations
A repeated measures ANOVA (motor fluctuations and follow up time) was performed to study differences in fluctuations in motor performance before and after operation. There was a significant interaction effect between the two factors (F = 7·2; P < 0·003). Table 1 gives the means. In summary, the proportion of "on plus" occasions decreased considerably at four months and there was a proportional increase in the proportion of "on" occasions (table 1). The proportion of "off" occasions did not vary significantly during the study period.

POSITION AND SIZE OF THE LESIONS
The lesion size as measured on CT (four months after operation) was 65·5 mm³ (95% confidence interval 95% CI) (44·9–86·1) in the 22 patients. Two patients with very small lesions were reoperated on and the corresponding group mean volume was 72·3 mm³ including these operations. Fifteen of the 22 patients also underwent a stereotactic MRI study four months after surgery. The mean volume of the MRI lesion was 72·8 mm³ (95% CI 46·8–101·8 mm³).

four and 12 months after reoperation. Results have been calculated on each of these follow up combinations. The results did not affect the group mean.
Video recordings
Because of technical errors, one patient (No. 11) was not video recorded at four months. Table 2 summarises the median and maximum values of the UPDRS items at baseline and four months after operation. The most prominent finding was a reduction of dyskinesiae on both the operated and non-operated sides (tables 2 and 3). In 12 patients (out of 13) who had dyskinesiae preoperatively, these had completely vanished four months after operation. The only patient (No. 20), who did not improve doubled her levodopa dose postoperatively, which might explain the negative result in this case. The maximum values of axial dyskinesias and tremor were significantly decreased postoperatively and there was a tendency, however not significant, towards improvement in the median values (table 2). The maximum values of rapid alternating movements of the hands and gait were significantly decreased postoperatively, but not the median values. The items rising from a chair and posture did not change after the operation. Patients with freezing were not improved.

Timed motor tests
Preoperatively, the mean time of the pegboard test was 44 (range 28–60) seconds and of the pronation-supination test 52 (range 17–173) seconds on the operated side. The mean time of the gait velocity test was 27 (range 16–41) seconds. No significant changes were found at four months compared with preoperative scorings, neither on the operated nor on the non-operated side.

Optoelectronic movement analyses
A total of 230 test sequences containing 879 standardised lifting movements were analysed at baseline and 231 tests/1028 lifts at four months. In patient No 2 no recording could be made at the four month follow up. There were no significant differences between the right/left side and between operated/non-operated sides, in recorded MT and SI estimates for individual patients at any of the observation occasions (Kruskal-Wallis and Mann-Whitney tests). In addition, no significant changes in MT or SI could be detected postoperatively, either when the overall data including all tests during the day were assessed or if a subgroup containing data with the selected longest MT of the day were analysed (Mann-Whitney test). However, when compared with the preoperative values, there was a significant increase in the average number of performed lifting movements per test sequence (Mann-Whitney, P = 0.02).

Follow up at 12 months
Motor fluctuations
Table 1 gives the mean percentage of motor fluctuations. Compared with the preoperative investigation, the proportion of “on plus” occasions decreased considerably and there was an increase in the proportion of “on” occasions. The proportion of “off” occasions did not change. Compared with the results at four months, there were no changes.

Video recordings
Table 2 summarises the median and maximum values of the UPDRS items at baseline and 12 months postoperatively. The results were similar to those at four months. There was a reduction in limb dyskinesias in both the operated and non-operated sides (tables 2 and 3). On the operated sides, the maximum values of tremor, axial dyskinesia, and rapid alternating movements of the hands were significantly decreased postoperatively. There was a tendency (non-significant), towards improvement in the median values of tremor and axial dyskinesia (table 2). There were no significant differences before and after operation in the items gait, arising from chair, and posture. Patients with freezing were not improved.

Timed motor tests
No significant changes were found at 12 months compared with preoperative scores or to the scores at four months, neither on the operated nor on the non-operated side.

Optoelectronic movement analyses
A total of 230 test sequences containing 879 standardised lifting movements were analysed. There were no significant differences between the right/left side and between operated/ non-operated sides in recorded MT and SI estimates at individual patient data level (Kruskal-Wallis and Mann-Whitney tests). In addition, no significant changes in MT or SI could be detected 12 months postoperatively, even when the overall data including all tests during the day or a subgroup containing data with the selected longest MT of the day were analysed (Mann-Whitney test). The increase in the average number of performed lifting movements per test sequence found at four months could not be confirmed.

Discussion
In this prospective long term study we confirmed other reports of a beneficial effect of PVP on tremor and dyskinetic movements.16-20 24-26 38 39 The effect on tremor was moderate but very pronounced on dyskinesiae, especially contralaterally but also on axial structures and ipsilaterally. The patients had a better quality of “on-off” fluctuations postoperatively with a significant increase in the number of more normal “on” phases and reduction in the number of “off” phases with dyskinesiae.

The number of “off” phases, however, did not change significantly and contrary to most earlier reports,16-20 24 25 40 41 we could not find any consistent improvement in bradykinesia or gait disturbance over time. Our negative results in this respect are in line with those in a small group of five patients with nine operative procedures.26 As in a recent report, we were also unable to find any improvement in freezing of gait.25 The PLM method, which has proved its value as a sensitive indicator of changes in
bradykinesia in Parkinson's disease,\textsuperscript{11} 14-21 did not show any significant changes in MT or SI of the standardised tasks in the present study. The reasons for this may be twofold. Firstly, the defined task involved a movement which includes a visually guided reaching task, and it may well be that this motor act is less influenced by the dyskinetic components, which are clearly reduced by the PVP. Secondly, the algorithms of the PLM test are mainly focused on the duration of the motor act and timing of the subparts. Pseudorandom deviations in space from the "ideal" movement path, which is one characteristic of dyskinetic behaviour, might not be optimally monitored and quantitatively assessed by the PLM method. The present finding, of a significant postoperative increase in the average number of lifting movements per test sequence, could indicate a more efficient performance with less time spent on movements unrelated to the particular task.

Previous reports\textsuperscript{16-20, 38 39} have shown that PVP has a great positive effect on almost all symptoms and signs in Parkinson's disease; bradykinesia, rigidity, tremor, gait and voice disturbance, balance, freezing, and also on ADL and psychiatric functions. However, the preoperative and postoperative data on the patients and methods are insufficient to make such far reaching conclusions,\textsuperscript{21, 22} and the long lasting benefit of pallidotomy still remains to be proved.\textsuperscript{27 28 45}

Recent results from elsewhere\textsuperscript{25} can be compared in many aspects with those in the present study. Important information is given in these other two studies, that the motor improvement was significantly better in the worst off condition (12 hours off medication) than in the best on condition after medication. One study\textsuperscript{25} indicated that there was no significant difference in blinded and non-blinded scorings of motor performance preoperatively and postoperatively in the on states. The best on conditions are more like the hourly rated Parkinson's disease conditions in the present study than the 12 hours off medication states.

The right or optimal size and location of the pallidal lesion are factors of probably great importance for the outcome of the operation, but the optimal lesion site and size remain uncertain.\textsuperscript{45} The locations of the PVPs in the present case series were in close vicinity to the target suggested by Sven-nilson et al.\textsuperscript{15} and Laitinen et al.\textsuperscript{17} This is illustrated in fig 2 (patient No 21), where a right sided PVP by Laitinen one year before inclusion can be compared with a left sided PVP in this study. The anatomical location of the lesions is similar in both hemispheres. Dogali et al.\textsuperscript{14} used microelectrode registration to locate the pallidal target and claimed that all lesions were within one mm of the intended target. However, on the coronal MRI illustrating the pallidotomy (figure 2a on page 755 of Dogali et al.) it is clear that this lesion is too far anterior compared with the target suggested by Laitinen and Leksell. The optic chiasm, which is visible on the slice, lies ventral and slightly anterior to the anterior commissure, but the intended target according to Laitinen and Leksell lies in the coronal plane, 9–11 mm behind the level of the anterior commisure. Perhaps such a different location of the pallidal lesion might explain the better effect on bradykinesia in the study by Dogali et al.

It is clear that this motor act is less influenced by the dyskinetic components, which are clearly reduced by the PVP. Secondly, the algorithms of the PLM test are mainly focused on the duration of the motor act and timing of the subparts. Pseudorandom deviations in space from the "ideal" movement path, which is one characteristic of dyskinetic behaviour, might not be optimally monitored and quantitatively assessed by the PLM method. The present finding, of a significant postoperative increase in the average number of lifting movements per test sequence, could indicate a more efficient performance with less time spent on movements unrelated to the particular task.

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