Dementia with Lewy bodies is diagnosed according to consensus criteria when progressive cognitive decline is accompanied by two of three clinical features (probable DLB; one of three for possible DLB). These features comprise well defined visual hallucinations, fluctuating cognition, and spontaneous parkinsonism. Up to 78% of DLB patients have extrapyramidal syndrome (EPS). Drugs used in the management of dementia, specifically cholinesterase inhibitors and neuroleptics, may worsen the EPS. Conversely, medications traditionally used to treat EPS can cause or worsen neuropsychiatric side effects, and thus have only been tentatively used in DLB to date for fear of precipitating or exacerbating an already fragile neurobehavioural state.

The most effective oral therapy available for Parkinson’s disease is levodopa. Other dopaminergic therapies include dopamine agonists, anticholinergics, selegiline, and amantadine, but the side effects of these agents, including confusion, somnolence, and hallucinations, preclude their safe use in DLB. The benefit of L-dopa in the management of DLB is unclear, with most data available from retrospective, non-systematic series. Studies have reported no or marked improvement, while others concluded that the lack of a documented L-dopa response, or even a perceived need to treat with L-dopa, could be used as a distinguishing factor between PD and DLB. The aim of this study was therefore to ascertain prospectively whether DLB patients demonstrate L-dopa responsiveness and whether, over a 6 month therapeutic period, they were able to tolerate this medication. We compared L-dopa responsiveness in DLB patients with that elicited in PD patients with and without dementia.

**METHODS**

**Patients**

PD and DLB patients were recruited from hospital and community dwelling populations under the care of neurologists, old age psychiatrists, and geriatricians. The Newcastle and North Tyneside ethics committee approved the study, and all subjects gave informed consent to participate. Diagnosis was made by consensus among experienced clinicians using the consensus criteria for DLB and the UK Parkinson’s Disease Society Brain Bank criteria for Parkinson’s disease. Patients with PDD conformed to the Brain Bank criteria but also fulfilled the Diagnostic and statistical manual of mental disorders, 4th ed (DSM IV) criteria for dementia and clinical diagnostic criteria for probable DLB, developing more than 12 months after the onset of motor symptoms.

The Mini Mental State Examination (MMSE) was used to provide a global measure of cognition. Motor assessments comprised the Unified Parkinson’s Disease Rating Scale part III (motor subsection), measurement of finger tapping times, and walking time tests, as recommended for monitoring L-dopa response. Patients already receiving L-dopa treatment were fasted overnight and assessed prior to their first daily dose of L-dopa, in a practically defined “off” state. Patients were then given co-beneldopa dispersible in a dose equivalent to their regular morning L-dopa dose (usually equivalent to one third of their total daily dose) and the acute motor effect was monitored every 30 minutes over the next 2 hours. Patients were then continued on their regular L-dopa dose and reassessed either on or off L-dopa, as tolerated, 6 months later.

If patients were not previously taking L-dopa this was commenced after their baseline motor assessment at a dose.

**Abbreviations:** DLB, dementia with Lewy bodies; DSM IV, Diagnostic and statistical manual of mental disorders, 4th ed; EPS, extrapyramidal syndrome; MMSE, Mini Mental State Examination; PD, Parkinson’s disease; PDD, Parkinson’s disease with dementia; UPDRS, Unified Parkinson’s disease Rating Scale.
of cocarboxylase 12.5/50, three times daily and gradually titrated up according to dose response and side effect profile. These patients were fasted and assessed as described above in a practically defined off state 6 months after commencing treatment.

A positive L-dopa response was defined as a >20% improvement in score in two of three tests of motor function, while a <10% improvement was classified as L-dopa unresponsive. This was based on the results of a meta-analysis of assessments of motor response in PD. An improvement of <20% in UPDRS III score is also recognised as indicative of positive motor response to dopamine in drug naïve patients. The effects of L-dopa upon cognition, mood, and sleep in DLB will be the subject of another publication and are not discussed further here.

A single investigator (SM) performed all motor assessments, unblinded to diagnostic status. Inter-rater reliability for UPDRS scoring between the study rater and an experienced rater (DJB) was performed for 11 patients.

Statistical analysis
SPSS for Windows (version 11) was used for data analysis. All results were tested for normality using Kolmogorov-Smirnov test. For continuous variables, differences between groups were assessed using one way analysis of variance with post hoc Games-Howell analysis tests to determine group differences. For non-parametric data, Mann-Whitney U test on two independent samples was used. All statistical tests were two tailed and were regarded as significant at p<0.05. Inter-rater reliability for UPDRS ratings was assessed using a weighted kappa statistic that controls for chance agreement between raters, with a value of 1 indicating perfect agreement.

RESULTS
A total of 91 patients were included in this study, of whom 27 had DLB. Baseline demographic data for these patients is shown in table 1, and compared with data for patients with PD (n = 31) and PDD (n = 33).

There was a male predominance in both the PD (24 men versus seven women) and PDD (23:10) groups, compared with the DLB group (15:12) (χ² = 12.5, p<0.001). The mean age of the PDD group (72.5 years) was less than either the PD (77.1 years, p<0.05) or DLB (77.5 years, p<0.05) groups, although, unsurprisingly, mean disease duration was longer in the PDD group (9.0 years compared with 4.9 years for PD and 2.7 years for DLB). MMSE scores in DLB and PDD groups were comparable (18.5 ± 20.2, respectively, p = 0.384) and both were less than the mean PD group score (26.0; p<0.0001). Cholinesterase inhibitor use was 73% and 96%, respectively, for the PDD and DLB groups.

Three PDD patients could not complete a fasting motor assessment due to severe motor disability, and three patients with PD did not require L-dopa. Baseline motor data for the remaining patients is presented in table 2. This combines patients assessed in a practically defined “off” state with the those who required but who had not yet commenced L-dopa. Given the longer disease duration and the fact that only six of the 27 DLB patients were taking L-dopa at baseline, it is not surprising that the mean L-dopa dose was significantly greater in the PDD group (p = 0.001; 590 mg per day, compared with 98 mg for DLB and 257 mg for PD groups).

Inspection of table 2 also reveals that the PDD group had the most significant motor impairment in all parameters assessed. There was no significant intergroup difference in the walking time step second product, although the large standard deviations made these data difficult to interpret. The greater product in the PDD group could be interpreted as reflecting their greater disease duration, however. The evidence for more severe EPS in PDD persisted using the UPDRS-5 subscale (calculated using the sum of total scores from five items of the full UPDRS III, previously shown to be independent of the severity of cognitive impairment). Inter-rater reliability testing confirmed a high level of agreement between assessors for 11 cases checked for UPDRS III (average measure of intraclass correlation of 0.9775; confidence interval 95%).

Acute L-dopa challenge
Acute L-dopa challenges were performed in 14 of the 27 DLB patients. Reasons for the other 13 DLB patients not receiving this challenge were: patient previously on L-dopa but medically incapable of completing repeated assessments over a 3 hour study period (n = 4); patient receiving L-dopa de novo but declined or was medically incapable of completing an overnight fast and acute, repeated motor assessments (n = 3); patient receiving L-dopa de novo but not tolerated, therefore acute fasting challenge not performed (n = 4); referring physician requested L-dopa not be administered (n = 1); and severe symptomatic orthostatic hypotension precluding introduction of L-dopa (n = 1). Of 14 DLB patients assessed acutely, two had previously received L-dopa prior to study entry, while 12 were commenced on L-dopa as part of this study. The mean dose of L-dopa administered for the challenge to the DLB group was 103 mg, and this did not differ significantly from the dose administered to the PDD (115 mg) and PD (101 mg) patients.

The magnitude of motor response to L-dopa was reduced in the PDD group for all parameters tested, compared with the PD and PDD patients (table 3). These differences reached statistical significance for the UPDRS III (p<0.05) and finger tapping (p<0.05), with the improvement in DLB significantly less in both tests compared with that seen in PDD (p<0.05). Although an improvement of >20% was seen in the walking time test for the DLB group, this is of uncertain significance due to the wide standard deviation. The UPDRS III, UPDRS-5, and finger tapping responses did not differ significantly between the PD and PDD groups, although all improvements were modest, presumably reflecting the relatively conservative L-dopa dose used for the challenges.

The L-dopa response of each patient was assessed, with “response” defined as a greater than 20% improvement over baseline on two of the three motor assessments (as detailed above). When the data were analysed in this way, only 36% of the DLB group was classified as L-dopa responsive, compared with 70% of the PDD group and 57% of the PD

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Table 1 Baseline demographics for DLB, PDD, and PD groups

<table>
<thead>
<tr>
<th>Group</th>
<th>Mean (SD) age (years)</th>
<th>Sex M:F</th>
<th>Mean (SD) DD (years)</th>
<th>≤ChEI</th>
<th>≤L-dopa</th>
<th>Mean (SD) MMSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>DLB (27)</td>
<td>77.5 (6)</td>
<td>15:12</td>
<td>2.7 (1.6)</td>
<td>26</td>
<td>6</td>
<td>18.5 (4.6)</td>
</tr>
<tr>
<td>PDD (33)</td>
<td>72.5 (5)*</td>
<td>23:10</td>
<td>9.5 (5.8)*</td>
<td>24</td>
<td>33</td>
<td>20.2 (5)</td>
</tr>
<tr>
<td>PD (31)</td>
<td>77.1 (6)</td>
<td>24:7</td>
<td>4.9 (4.6)</td>
<td>0</td>
<td>19</td>
<td>26.0 (2.5)*</td>
</tr>
</tbody>
</table>

DD, disease duration; ≤ChEI, number of patients taking a cholinesterase inhibitor at entry to the study; ≤L-dopa, number of patients taking L-dopa at entry to the study. *p<0.001, †p<0.0001.

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group (table 4). Conversely, 29% of the DLB were classified as unresponsive, compared with 10% PDD and 21% PD patients (table 4).

Comparison between L-dopa responding and non-responding DLB patients revealed no difference in sex, disease duration, MMSE, UPDRS, five item UPDRS, or dose of L-dopa achieved. L-dopa responders were, however, younger (68.5 ± 77.1 years, p = 0.014).

**Chronic L-dopa use (tolerability)**

Of the 27 patients with DLB, one did not commence L-dopa at the request of the physician owing to fear of neuro-behavioural sequelae (UPDRS = 38) and one had severe symptomatic orthostatic hypotension. Thus, 25 of the 27 DLB patients were chronically treated with L-dopa. Six of these were already taking this medication at study entry. Of the 25 DLB patients, 82% (n = 21) warranted and tolerated long term L-dopa therapy. Nineteen patients with DLB were therefore commenced de novo on L-dopa for the purposes of this study, with a view to chronic dosing over at least 6 months. The six patients already taking L-dopa did not differ significantly in age, disease duration, MMSE, or baseline UPDRS from those not taking L-dopa at baseline assessment (data not shown).

Of the 19 patients, two were unable to tolerate L-dopa within the first 2 weeks due to worsening confusion (n = 1) and severe exacerbation of hallucinations (n = 1). A further two patients withdrew before the 3 month assessment, owing to gastrointestinal intolerance, nausea, abdominal pain, and bloating. All side effects resolved once the dopaminergic medication was stopped. Fifteen DLB patients (84%) were therefore able to tolerate L-dopa for the 6 month study duration. The mean daily L-dopa dose achieved was 323 (182) mg. None of the DLB patients either previously receiving or those commenced de novo on L-dopa developed dyskinesias during the study period.

**DISCUSSION**

The main conclusion from this study is that L-dopa, while generally well tolerated in modest doses by DLB patients, is not particularly efficacious in alleviating the parkinsonian syndrome seen in this disorder. Younger DLB patients may be more likely to experience a beneficial response. Thus, while 82% of 25 DLB patients were able to remain on a mean daily dose of 323 mg L-dopa for 6 months, formal L-dopa challenges indicated that only 36% of 14 patients tested could be classified as “responders”, according to previously published criteria. This contrasts with the 56 and 70% response rates in PD and PDD, respectively. Others have recently reported similar findings to the present study, with reduced motor improvement on formal levodopa testing in DLB patients. Variable dopaminergic responsiveness has been reported in observational studies of DLB, although the retrospective nature, lack of control groups and non-standardised assessment of motor function make these data difficult to interpret. The lack of a documented L-dopa response, or a perceived need to treat with L-dopa has been suggested to be of use as a distinguishing factor between PD and DLB. In contrast, all 12 DLB patients in another study were deemed to be L-dopa responsive, although this was not quantified. More recently, L-dopa therapy was initiated in 48% of 40 DLB patients at a mean dose of 525 mg/day, with a “good” motor response noted in 66%. Efficacy was maintained for a mean follow up of 3.2 years, although dyskinesias developed in 16% of the patients. This study was, however, a retrospective design and motor response was again not quantified.

There are several reasons why DLB patients might be expected to respond less well to dopaminergic treatments, including intrinsic striatal α-synuclein pathology and reduced caudal putamen dopamine D2 receptors. More recently, loss of dopamine D3 receptors on corticostriate projection neurones has also been proposed as a possible mechanism for L-dopa refractoriness. Alternatively, the emergence of so-called “non-dopaminergic” motor features, mediated by degeneration in other brainstem nuclei such as the pedunculopontine nucleus, would be predicted to give a reduced response to L-dopa therapy.

Interestingly, a similar loss of dopaminergic responsiveness has also been suggested to occur in PDD, and clinical data suggest that non-dopaminergic motor features may, indeed, be over-represented in both PDD and DLB. In the present study, however, the PDD group showed a 70% overall response to the acute L-dopa challenge, with only 10% of the 30 patients tested being classified as non-responders. Although UPDRS III scores were highest in this group, compared with the PD and DLB patients, the mean disease duration was still relatively short at 9.0 years. The contrast in L-dopa responsiveness between DLB and PDD patients in this study highlights a potentially important clinical difference between these conditions. Further work is required to determine the pathophysiological mechanisms underpinning these observations.

| Table 2 Baseline motor data for DLB, PDD, and PD patient groups |
|-------------------|-----------------|-----------------|-----------------|
| Group (n)         | L-dopa (%)      | Daily L-dopa (%) | UPDRS III (%)   |
|                   | L-dopa naive (%)| L-dopa mg (SD)  | 14.3 (6.4)      |
| DLB (27)          | 21              | 98 (188)        | 23.5 (14.4)     |
| PDD (30)          | 0               | 590 (414)       | 18.7 (73.6)*    |
| PD (28)           | 9               | 257 (241)       | 15.7 (5.6)      |
|                   |                 |                 | 40.6 (9.4)      |
|                   |                 |                 | 801 (1717)      |

Data shown as mean (SD). *p < 0.05, **p < 0.001, ***p < 0.0001. FT, finger tapping; WT, walking test.

| Table 3 Group changes in motor parameters with acute L-dopa challenge |
|------------------|---------------------|---------------------|
| Group (n)        | L-dopa (mg (SD))    | % Δ UPDRS III (%)   |
| DLB (14)         | 103.3 (51.6)        | 13.8*               |
| PDD (30)         | 115.5 (45.8)        | 23                  |
| PD (28)          | 101.8 (21.4)        | 20.5                |

*p < 0.05. FT, finger tapping; WT, walking test.

| Table 4 Responder and non-responder frequency with acute L-dopa challenge |
|-------------------|---------------------|---------------------|
| Group (n)         | UPDRS III (%)       | FT (%)              |
| DLB (14)          | 5                   | 16                  |
| PDD (30)          | 16                  | 21                  |
| PD (28)           | 14                  | 14                  |

Responder: defined as a group of patient manifesting ≥ 20% improvement in two or more of the three motor tests after introduction of L-dopa therapy. Non-responder: defined as <10% improvement in two or more of the three motor tests. FT, finger tapping; WT, walking test.
dose titration employed in the one third of the PD group that were introduced to L-dopa de novo. This, in turn, led to the administration of a lower than normal L-dopa dose for the acute challenge (mean 101 mg), whereas most other studies have advocated a supranormal dose challenge (of at least 250 mg). The reason for this choice was to make the acute response between DLB and the other groups as comparable as possible. It was felt that using higher L-dopa doses in frail and cognitively impaired patients would run an unacceptably high risk of exacerbating or causing neuropsychiatric complications. A further potential explanation is the fact that the PD patients performed particularly well on the finger tapping test at baseline, and thus there may have been a “ceiling effect” in response to L-dopa on this measure.

There are several drawbacks to this study. It was an open label design, therefore any results are subject to a degree of bias, while some of the benefits of L-dopa use may reflect a placebo response. Many of the patients recruited were elderly and frail. Physical and cognitive disability was profound in some and this may have rendered the interpretation of tests results unreliable. Test conditions were not always standardised, and while the finger tapping test and UPDRS are less subject to environmental influence, assessments were often carried out in the patient’s home, making the 6 metre walk test prone to increased variability. Furthermore, only 14 of 27 (52%) DLB patients were able to undergo an acute L-dopa challenge, potentially limiting the generalisability of our results. The doses of L-dopa used throughout this study were relatively conservative and we could not exclude a greater motor response with higher doses. It is noteworthy that all but one of the DLB patients were taking cholinesterase inhibitors when exposed to L-dopa, so we are unable to comment upon the tolerability of dopaminergic treatment in DLB patients not taking cholinesterase inhibitors. Although we found that younger DLB patients may respond better to L-dopa, the relatively small numbers do not permit a firm conclusion to be made in this regard.

Despite these drawbacks, the study has a number of advantages. To the best of our knowledge, this is the first time that DLB patients have been prospectively assessed both acutely and chronically for L-dopa responsiveness and tolerability. A single assessor performed all the motor tests, thereby eliminating interobserver variability. The inclusion criteria were broad, so the study population recruited and conclusions drawn are likely to represent the “real life” situation faced by clinicians in their day to day practice.

In conclusion, we suggest that L-dopa can be employed in the management of dementia with Lewy bodies.

ACKNOWLEDGEMENTS

Our thanks to E N Rowan for her valued statistical review and to A Nicholson and J Bowman for help with database formation.

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Competing interests: none declared

REFERENCES


