ABSTRACT

Background: The long duration response to levodopa in Parkinson’s disease outlasts the elimination of the drug by days to weeks. Though a substantive part of anti-parkinsonian motor benefit, it cannot easily be observed.

Objectives: To infer the magnitude of the long duration response during the first decade of Parkinson’s disease and to identify factors that influence it.

Methods: Serial defined off scores of 24 patients from a longitudinal study of the levodopa short duration response were used to establish their rate of motor progression. A line of notional untreated disability (as if drug treatment had never been given) with the same gradient was the basis for a calculation of the long duration response. Predictors of mean long duration response amplitude were identified using a multiple linear regression model.

Results: Over a mean treatment period of 16.6 ± 4.4 years, the annual progression of motor disability was 2.3% of the maximum motor disability. The long duration response composed 49% of the total levodopa response during the first decade of treatment, and this proportion was significantly higher soon after commencing levodopa (p = 0.001). Higher pre-treatment motor score (r = 0.62) and lower MMSE (r = 0.61) were the main predictors of a larger long duration response. There was little correlation between long and short duration responses.

Conclusions: Long duration responses contribute almost half of the total levodopa benefit during the first decade of treatment. An appreciation of both long and short duration components of drug symptomatic effects is important in clinical trial design to investigate possible neuroprotective treatments.

INTRODUCTION
The long duration response (LDR) to levodopa is a motor benefit in a patient with Parkinson’s disease (PD) that outlasts the elimination of the drug by days to weeks. Right at the beginning of the era of dopaminergic therapy, George Cotzias and his colleagues were aware of it. They noted that after prolonged use of D,L-dopa, it took 4 – 14 days for motor state to return to baseline when the drug was ceased.\(^1\) Two previous levodopa withdrawal studies have estimated the LDR response at about one third of the total levodopa response.\(^2,3\) In the first year of treatment, the majority of motor benefit comes from the LDR.\(^4\) Drugs other than levodopa, dopamine receptor agonists for instance, have LDRs as well.\(^5,6\)

The magnitude of the initial levodopa LDR can be estimated by comparing motor disability immediately before treatment with when the drug is withheld overnight after weeks to months of treatment.\(^7\) Thereafter, it is only possible to measure it by prolonged drug withholding, impractical because of the difficulties in managing the loss of motor benefit and the delay in restoring it. There is uncertainty about the duration of withdrawal needed to reveal fully a LDR. LDRs are the ‘dark matter’ of anti-parkinsonian motor benefit, substantive but not directly observable.

Using serial measurements of defined \textit{off} states in a cohort of PD patients studied longitudinally, we devised a method to estimate the LDR. By determining the gradient of progression of \textit{off} scores, it is possible to infer the level of disability if drug treatment had never been commenced and to quantify the motor benefit that is not captured by \textit{on} and \textit{off} phase assessments.

**METHODS**

Thirty-four patients with PD were recruited to a longitudinal study of the levodopa motor response that began almost 30 years ago. Detailed methodology including entry criteria are described in earlier publications.\(^8,9\) A modified Webster scale (12 areas of motor function scored from 0 to 3 to give a maximum disability score of 36)\(^10\) was the chief motor assessment. A motor score was recorded before levodopa was started and at optimum treatment response during the next 6 months, with the initial drug response defined as the difference. At 3-year intervals, a researcher conducted defined \textit{off} state levodopa test-dose assessments on surviving subjects. Levodopa was administered while fasting and after withholding of other medication, with the \textit{on} state defined as the maximum improvement over the subsequent 30 – 90 minutes. Amplitude of the short duration response (SDR) was calculated as \textit{off} minus \textit{on} score. The Folstein Mini Mental State Examination (MMSE)\(^11\) was performed at each assessment. Patients were classified for the presence of motor fluctuations during the first 5 years of levodopa treatment,\(^9\) and for motor subtype from
their modified Webster scale scoring. Levodopa equivalent daily doses (LEDD) were calculated using standard conversion factors. This study has institutional research ethics approval.

**Notional untreated disability and LDR calculation**

In patients with at least two test-dose assessments, individual gradients of off state motor progression were calculated using linear least squares regression. A second line with an identical gradient was drawn from each subject’s pre-treatment motor score to represent their notional untreated disability (hypothetical progression as if anti-parkinsonian treatment had never been commenced). Figure 1 shows these parallel lines on a graph of mean results from all participants.

The LDR amplitude was calculated by subtracting the off score from the notional untreated disability score. This was done only for the 3 test-dose assessments performed within the first 10 years of treatment because of uncertainty about extrapolation beyond this point. The test dose for these assessments was levodopa 200mg/ carbidopa 50mg. Total levodopa response was defined as the sum of the SDR and LDR.

**Statistical methods**

A linear regression model of individual mean LDRs was employed to identify clinical variables that predict LDR amplitude. Age of disease onset, pre-treatment motor score, MMSE, LEDD, mean SDR, disease phenotype and presence of motor fluctuations were the factors of interest. We used the backward elimination method and ANOVA to compare model fit and we tested for interactions using product terms.

We also examined disease progression with a linear mixed-effects regression model of off state scores, using treatment duration as a fixed effect. We estimated random effects for subjects to account for repeated measures and differing durations of follow-up.

Descriptive statistics are reported as mean ± standard deviation. Continuous non-parametric variables were compared using the Wilcoxon signed rank test. Data analysis was performed using R statistical software (R Foundation for Statistical Computing, Vienna, Austria, 2017) and the packages nlme and ggplot2.

**RESULTS**

Twenty-four patients had two or more test-dose assessments, which allowed estimation of a gradient of off state progression. Their mean age at PD diagnosis was 61.1 ± 11.6 years; time from diagnosis to treatment initiation was 0.6 ± 2.1 years. The mean follow-up duration for these patients was 16.6 ± 4.4 years. Nine had a tremor-dominant phenotype, and 14 developed motor fluctuations. Only 5 patients were still alive at the end of the study period. Mean treatment duration for those who had died was 15.2 ± 5.0 years (range 5.2 - 20.3). At the final assessment of those performed during the first treatment decade, the mean MMSE score was 26 ± 5.4 and the mean LEDD was 628 ± 305mg. Although all patients
began with levodopa monotherapy, by the end of this 10-year period other drugs had also been used: bromocriptine (2), pergolide (2), deprenyl (4), benztrpine (1).

**Motor Progression & LDR amplitude**

Figure 1 shows the pooled initial and test-dose motor scores for all 34 patients originally enrolled in the study, with numbers of survivors at each assessment. The gradient of *off* phase deterioration on this graph is 2.0% p.a. of maximum motor disability. Figure 2 shows the 24 individual lines of best fit for progression of motor disability. According to the linear mixed effects regression model, annual progression in disability for this group was 2.3% of maximal motor disability.

Using the notional untreated disease trajectory, we calculated mean LDR amplitude at the first defined *off* state assessment as 4.8 ± 3.6, which is equivalent to 62% of the total motor response. The mean of all LDR amplitudes calculated for the first decade of treatment was 5.1 ± 4.2 or 49% of the total motor response. There was a significant reduction in the contribution of the LDR to total motor response between the first and the last of these estimations (p = 0.001, r=-0.49). There was no significant change in the absolute magnitude of the LDR.

**Predictors of LDR amplitude**

Only higher pre-treatment motor score ($r = 0.62, p = 0.001$) and lower MMSE ($r = 0.61, p = 0.001$) correlated with LDR amplitude. In a two-factor regression model, pre-treatment disability and MMSE predicted 65% of the variance in mean LDR. Although mean SDR (Figure 3) and motor fluctuations did not correlate with mean LDR, their addition as factors improved the regression model of mean LDR (four-factor model adjusted $R^2 = 0.74, F (21,19) = 4.50, p = 0.03$), and were thus included in the final model (Table 1). LEDD, age at diagnosis and disease phenotype were not significant predictors and there were no interactions.

**DISCUSSION**

Overall, the LDR composed about half of the total levodopa motor response, comparable to the size of the SDR and somewhat larger than previous estimates. The percentage was significantly higher early in the disease course, starting at 62% for the first test-dose measurement and falling to 42% by the final reckoning. As shown by the multiple regression analysis and Figure 3, there is surprisingly little correlation between the sizes of LDRs and SDRs. The strongest predictors of a large LDR were pre-treatment disability and reduced MMSE score, each of these accounting for about 38% of the variation of the mean LDR. Greater initial motor deficit and early cognitive decline both imply a heavier burden of Lewy pathology in the brain, yet these patients appear to have a greater early LDR. Age at diagnosis had no significant effect.

Our method for calculating the LDR takes advantage of longitudinal defined *off* state measures of the SDR. It relies on three important assumptions. First, that disease progression is linear during the first part of the disease course. Secondly, that the rate of decline is 2-3%. Three other longitudinal studies, though not employing rigorous levodopa test-dose methods, produced linear plots of progression over 8-year periods, with estimates
of annual deterioration of between 1.4% and 3.1%. Third, that commonly used symptomatic treatments do not modify the underlying disease process. The validity of a notional untreated disability line that runs from the pre-treatment motor score in parallel to the trajectory of the defined off phase scores rests squarely on this last assumption. The best interpretation of available evidence is that levodopa, the dominant anti-parkinsonian drug in this study, has a powerful but purely symptomatic effect on PD with no influence on the underlying rate of progression.

Large clinical trials of possible neuroprotective agents have struggled to discern symptomatic motor benefit from an effect on the natural disease course. Several LDR considerations are relevant. Trial designs have incorporated questionable assumptions about wash-in and wash-out times for symptomatic effects. Wash-in is an estimate of the time taken for both SDR and LDR to fully develop. Wash-out, on the other hand, is mainly an estimate of time taken for a LDR to decay, since SDRs of most drugs can be predicted to follow their pharmacokinetic elimination curve after discontinuation. Another impediment to neuroprotective drug trials is the difference in magnitude between symptomatic and disease modifying effects. An agent that is capable of completely arresting the disease would cause a 2 – 3% deviation from the trajectory of a placebo control group over 1 year. A drug that retarded progression by 20% would cause the line to deviate by only 0.4% to 0.6% p.a. But the symptomatic effect of dopaminergic therapy runs at around 22% of the maximum disability score for the combined SDR and LDR. In a clinical trial of less than a decade, even modest variability of symptomatic effects may obscure a disease modifying one. Such variability could come from standard therapy or from an inherent symptomatic effect of a putative neuroprotective agent. At least the SDR can be measured by a defined off phase test dose method. There is no practical way of directly measuring variations of a LDR over time. Until a better biomarker for pathological progression in PD is found, proof of neuroprotective effect must depend on measurement of the rate of deterioration of motor disability scores. Simple, placebo-controlled study designs of long (5 – 10 years) duration may be the best approach. Standardisation of anti-parkinsonian drug therapy and avoidance of delayed drug start or withdrawal protocols should reduce the risk of confounding LDR effects.

Nutt et al. concluded that the presence of a long and a short duration levodopa effect implies either two different compartments that take up and release dopamine, or two different effector systems for levodopa with different time courses. The most plausible explanation of a dopaminergic response in the absence of dopaminergic drugs is that the separate compartment of dopamine release for the LDR is the surviving nigral neurons. A residual population of nigral cells is important, probably essential, to the dopaminergic drug response. Roughly 50% can be lost and the dopaminergic nigrostriatal system has sufficient functional reserve to forestall motor deficit. Thereafter, signs of parkinsonism appear and progress up to the point that drug treatment is commenced. But once an LDR is established, Figure 1 shows that more than 5 years will pass before the level of pre-treatment disability is reached again. Concomitant dopaminergic pharmacological treatment must somehow restore the capacity of these neurons to synthesise and release dopamine when short duration effects wane or drugs are temporarily withheld. Perhaps an alternative source of dopamine receptor stimulation from tablets allows nigral cells temporarily to reduce their
energy expenditure because of lowered demand on their metabolic or firing states. That the LDR is not immediately re-established by levodopa after a short drug holiday implies a compensatory mechanism that takes time to recover.23 Levodopa withdrawal studies suggest a relationship between the LDR and the number of surviving nigral neurons. There is a LDR in advanced PD but it decays more rapidly than in less severely affected patients.24 The LDR of levodopa can be sustained by the dopamine receptor agonist apomorphine.25

There are some methodological uncertainties in our approach to the LDR. Disease progression may not be linear. Most time relationships in the natural world are not linear, and there is some evidence that late progression of PD is exponential.26 Notional untreated disability is a logical concept, though it is not certain that progression would follow such a predictable parallel course without drug treatment. The SDR measures that form the basis for the ascertainment of LDR have their own margin of uncertainty. Defined off states at the start of the day may incorporate a sleep benefit element. A minority of patients were taking dopamine receptor agonists with longer durations of action than levodopa.

Rightly has the main focus of research into PD moved away from established pharmacological treatments on to molecular pathophysiology, its patterns of involvement beyond the motor system, and the keys that this might hold to modifying disease progression. Yet that singular feature of PD, its dopaminergic motor response, remains a mysterious thing. We have tried here to show how an appreciation of both long and short duration components of the levodopa symptomatic effect is essential to clinical trial design to identify neuroprotective drugs.

**Figure 1**

Mean modified Webster scores for all participants. Trapezium-ended box: upper pole, pretreatment motor score; lower pole, optimum initial treatment response. White rectangular
boxes: mean levodopa SDR amplitudes. Dashed line of best fit for mean off phase scores. Dotted line for notional untreated disease trajectory. Black rectangular boxes: calculated LDRs; gray shaded boxes show possible LDRs for later assessments.

Figure 2.
Dashed lines of best fit for each subject’s off motor scores. Solid line with gradient of 2.3% p.a. obtained from the linear mixed effects regression model of all off scores. The lengths of lines are proportional to treatment durations.

Figure 3. Each data point represents mean SDR and LDR for each subject’s first decade of treatment. There is little correlation ($r = 0.12, p = 0.58$). 95% confidence interval shaded in gray.
Table 1. Multiple regression factors that predict mean LDR amplitude

<table>
<thead>
<tr>
<th>Disease factors</th>
<th>b coefficient</th>
<th>95% confidence interval</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-treatment motor score</td>
<td>0.67</td>
<td>0.43: 0.91</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>MMSE</td>
<td>-0.44</td>
<td>-0.62: -0.27</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Motor fluctuations</td>
<td>2.86</td>
<td>0.42: 5.30</td>
<td>.024</td>
</tr>
<tr>
<td>Mean SDR amplitude</td>
<td>-0.80</td>
<td>-1.38: -0.22</td>
<td>.009</td>
</tr>
</tbody>
</table>

The adjusted $R^2$ of this model was 0.74

REFERENCES


DOCUMENTATION OF AUTHOR ROLES

Kanae Nagao
Research project: organization and execution
Statistical analysis: execution, review and critique
Manuscript: writing of first draft, review and critique

Ding C
Research project: organization and execution
Statistical analysis: execution, review and critique
Manuscript: writing of first draft, review and critique

Ganga G
Research project: execution
Manuscript: review and critique

Alty JE
Research project: execution
Manuscript: review and critique

Clissold BG
Research project: execution
Manuscript: review and critique

McColl CD
Research project: execution
Manuscript: review and critique

Reardon KA
Research project: execution
Manuscript: review and critique

Shiff M
Research project: execution
Manuscript: review and critique

Kempster PA
Research project: conception and organization
Statistical analysis: review and critique
Manuscript: writing of first draft, review and critique
FULL FINANCIAL DISCLOSURES OF ALL AUTHORS FOR THE PAST YEAR

Nagao K  
Stock Ownership in medically-related fields: none  
Consultancies: none  
Advisory Boards: none  
Partnerships: none  
Honoraria: none  
Grants: none  
Intellectual Property Rights: none  
Expert Testimony: none  
Employment: Monash Health; private neurological consulting practice.  
Contracts: none  
Royalties: none  
Other: none

Ding C  
Stock Ownership in medically-related fields: none  
Consultancies: none  
Advisory Boards: none  
Partnerships: none  
Honoraria: none  
Grants: none  
Intellectual Property Rights: none  
Expert Testimony: none  
Employment: Monash Health; private neurological consulting practice.  
Contracts: none  
Royalties: none  
Other: none

Ganga G  
Stock Ownership in medically-related fields: none  
Consultancies: none  
Advisory Boards: none  
Partnerships: none  
Honoraria: none  
Grants: none  
Intellectual Property Rights: none  
Expert Testimony: none  
Employment: Monash Health; private neurological consulting practice.  
Contracts: none  
Royalties: none  
Other: none

Alty J  
Stock Ownership in medically-related fields: Clearsky Medical Diagnostics Ltd and Dr Carsten Grimm Medical Consultancy Ltd.  
Intellectual property rights: none  
Consultancies: none  
Expert testimony: none  
Advisory Boards: Merz  
Employment: Leeds Teaching Hospitals NHS Trust (LTHT)  
Partnerships: none  
Contracts: Principal Investigator at LTHT for the following NIHR portfolio Parkinson's research studies: CTH301, PDSTAT, PDCOMM, Vision in PD, PROSPECT and co-investigator for OPTIPARK and PD Families.  
Honoraria: Allergan, MediConf UK Ltd, Merz  
Royalties: Taylor & Francis Group
Grants: Britannia, Merz, Allergan, Medtronic, Bial, Ipsen
Other: none

Clissold BG
Stock Ownership in medically-related fields: none
Consultancies: none.
Advisory Boards: none
Partnerships: none
Honoraria: none
Grants: none
Intellectual Property Rights: none
Expert Testimony: none
Employment: Monash Health, private neurological consulting practice.
Contracts: none
Royalties: none
Other: none

McColl CD
Stock Ownership in medically-related fields: none
Consultancies: Therapeutic Goods Administration (Australia).
Advisory Boards: none
Partnerships: none
Honoraria: none
Grants: none
Intellectual Property Rights: none
Expert Testimony: none
Employment: Australian Capital Territory Health; private neurological consulting practice.
Contracts: none
Royalties: none
Other: none

Reardon KA
Stock Ownership in medically-related fields: none
Consultancies: none
Advisory Boards: none
Partnerships: none
Honoraria: none
Grants: none
Intellectual Property Rights: none
Expert Testimony: none
Employment: St Vincent’s Hospital, Calvary Healthcare Bethlehem; private neurological consulting practice.
Contracts: Principal investigator for RESILIENT trial investigating BYM338 in inclusion body myositis.
Royalties: none
Other: none

Schiff M
Stock Ownership in medically-related fields: none
Consultancies: none
Advisory Boards: none
Partnerships: none
Honoraria: none
Grants: none
Intellectual Property Rights: none
Expert Testimony: none
Employment: Private psychiatric consulting practice.
Contracts: none
Royalties: none
Other: none

Kempster PA
Stock Ownership in medically-related fields: none
Consultancies: none
Advisory Boards: none
Partnerships: none
Honoraria: none
Grants: none
Intellectual Property Rights: none
Expert Testimony: none
Employment: Monash Health, private neurological consulting practice.
Contracts: none
Royalties: none
Other: none