Effects of a flexible galantamine dose in Alzheimer’s disease: a randomised, controlled trial

K Rockwood, J Mintzer, L Truyen, T Wessel, D Wilkinson

Abstract

**Objective**—To assess the efficacy and safety of galantamine in Alzheimer’s disease at 3 months using flexible dose escalation.

**Methods**—A randomised, double blind, placebo controlled trial in 43 centres in the United States, Canada, Great Britain, South Africa, Australia, and New Zealand. Patients with probable Alzheimer’s disease (n=386; 171 women) with a score of 11–24 on the mini mental state examination, and a score ≥12 on the cognitive subscale of the Alzheimer’s disease assessment scale (ADAS-cog) were randomised to placebo, or galantamine escalated over 4 weeks to a maintenance dose of 24 or 32 mg/day. The primary outcome measures were the change in ADAS-cog score and the clinician’s interview based impression of change plus caregiver input (CIBIC-plus) score. Activities of daily living (ADL) and behavioural symptoms were secondary outcomes. To compare the effects of highest levels of dosing, an observed cases (OC) analysis was undertaken, with classic intention to treat (ITT) and ITT with last observation carried forward (LOCF) as confirmatory analyses.

**Results**—At 3 months, galantamine (24–32 mg/day) produced a significantly better outcome on cognitive function than placebo (treatment difference=1.9 points on ADAS-cog, p=0.002) and a significantly better global response than placebo, as measured by CIBIC-plus (deterioration in 11–24 on the mini mental state examination, and a score ≥12 on the cognitive subscale of the Alzheimer’s disease assessment scale (ADAS-cog) were randomised to placebo, or galantamine escalated over 4 weeks to a maintenance dose of 24 or 32 mg/day. The primary outcome measures were the change in ADAS-cog score and the clinician’s interview based impression of change plus caregiver input (CIBIC-plus) score. Activities of daily living (ADL) and behavioural symptoms were secondary outcomes. To compare the effects of highest levels of dosing, an observed cases (OC) analysis was undertaken, with classic intention to treat (ITT) and ITT with last observation carried forward (LOCF) as confirmatory analyses.

**Conclusions**—Patients on galantamine, compared with those on placebo, experienced benefits in cognitive function and instrumental and basic activities of daily living. Flexible dose escalation of galantamine was well tolerated. (J Neurol Neurosurg Psychiatry 2001;71:589–595)

Keywords: Alzheimer’s disease; galantamine; nicotinic receptors; acetylcholinesterase inhibition; activities of daily living
more information becomes available on specific patterns of treatment effects with galantamine (perhaps, for example, early stabilisation of activities of daily living (ADL), lack of sleep disturbance, or improved attention) it may be that the clinical role of the nicotinic receptor might be better understood. As the tolerability of cholinesterase inhibitors is dose related, the present study was undertaken to test the efficacy and safety of galantamine using a flexible dose, in patients with mild to moderate Alzheimer’s disease, over 3 months.

**Methods**

**Patients**

Patients with a history of cognitive decline that had been gradual in onset and progressive for at least 6 months were included. Other inclusion criteria were:

- A diagnosis of probable Alzheimer’s disease according to the criteria of the National Institute of Neurological and Communicative Disorders and Stroke and Alzheimer’s Disease and Related Disorders Association (NINCDS/ADRDA).24
- Presence of mild to moderate dementia (a score of 11–24 on the mini mental state examination (MMSE)25 and a score ≥2 on the standard cognitive subscale of the Alzheimer’s disease assessment scale (ADAS-cog).26

Patients also had to have regular contact with a responsible caregiver. Those with concomitant diseases such as hypertension, congestive heart failure, non-insulin dependent diabetes mellitus, and hypothyroidism were included in the study provided the disease was controlled. Patients were excluded from the study if they had evidence of other neurodegenerative disorders; any cardiovascular disease thought likely to prevent completion of the study; clinically significant cerebrovascular, hepatic, renal, pulmonary, metabolic or endocrine conditions; clinically significant psychiatric disease, including moderate or severe or uncontrolled behavioural disturbances; urinary outflow obstruction; an active peptic ulcer; any history of epilepsy, or significant drug or alcohol misuse.

Patients previously treated with any cholinomimetic agent for Alzheimer’s disease, except muscarinic agonists, were also excluded. Any other medication being taken to treat dementia had to be discontinued. The use of other concomitant medication was permitted, except that, where possible, drugs with a psychotropic action were discontinued 48 hours before cognitive evaluation. Drugs with anticholinergic effects or cholinomimetic effects were avoided.

The trial was performed in accordance with the Declaration of Helsinki and subsequent revisions and approved by ethics committees at each centre. The patient (or a representative), together with the carer, provided written informed consent to participate.

**Design**

This was a 3 month, parallel group, placebo controlled trial undertaken in 43 centres in six countries (Australia, Canada, Great Britain, New Zealand, South Africa, and the United States). After a 4 week, single blind, placebo run in phase, patients were randomised to receive galantamine or placebo in a 2:1 ratio using a computer generated code. The assignments were kept in sealed, opaque envelopes until the point of allocation. Patients treated with galantamine received 8 mg/day for 1 week, increasing to 16 mg/day for the 2nd week and to 24 mg/day (12 mg twice daily) for the 3rd week. During week 4, the galantamine or placebo dose could be increased (to 32 mg/day—that is, 16 mg twice daily for galantamine) at the discretion of the investigator, based on tolerance. By the end of the 4th week the investigator could reduce the dose (for galantamine from 32 mg/day to 24 mg/day). Patients continued with their final dose of galantamine or placebo for a further 2 months. Throughout the study, all individual doses of galantamine and placebo were otherwise identical single tablets taken twice daily.

The primary efficacy variables were the standard, 11 item cognitive subscale of the Alzheimer disease assessment scale (ADAS-cog/11)28 to assess cognitive function, and the clinician’s interview based impression of change plus caregiver input (CIBIC-plus)29 to assess overall clinical response. The CIBIC-plus was scored by a trained clinician based on separate interviews with the patient and the caregiver; the clinician was blinded to other assessments. Scores ranged from 1 to 7 (1=markedly improved with respect to baseline, 7=markedly worse).

Secondary efficacy variables were:

- The expanded (13 item) version of the standard ADAS-cog subscale (ADAS-cog/13), with a score range of 0–85.28
- The proportions of responders (defined as improvements in ADAS-cog/11≥4 points from baseline).29
- Neuropsychiatric inventory (NPI), which assesses 10 domains of behavioural symptoms (score range of 0–120).30
- Disability assessment for dementia (DAD) scale, based on an interview with the caregiver, to assess basic and instrumental activities of daily living (ADL), initiation, planning and organisation, performance, and leisure; the DAD has a score range 0–100.10

All of the efficacy assessments were performed at baseline and after 1 and 3 months.

Safety evaluations throughout the study comprised physical examinations, ECG, vital sign measurements, standard laboratory tests, and monitoring for adverse events (classified according to World Health Organisation preferred terms). For the first 2 weeks, the investigator contacted the patient or caregiver by phone at weekly intervals to record any adverse events. Safety was further evaluated at monthly clinic visits at weeks 3 and 4, and after 2 and 3 months. Sleep patterns were assessed after 1 and 3 months using the seven item Pittsburgh sleep quality index (PSQI), which measures subjective sleep quality, sleep latency, sleep duration, habitual sleep efficiency, sleep disturbances, use of sleeping medication, and daytime dysfunction; each item is scored from 0–3 (0 indicates no difficulty, 3 indicates severe difficulty).32

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Using data from an earlier phase II trial, we calculated that 94 patients were needed to complete the study in the placebo group, and twice this number in the galantamine group, to achieve 80% power (α=0.05) for detecting a 2.5 point difference in the change in ADAS-cog/11 score between patients treated with placebo and with each of the maximum daily doses of galantamine.

All randomised patients who took at least one dose of trial medication were included in the analyses of baseline characteristics and safety data. As we were chiefly concerned with assessing the treatment effects of the maximum daily doses of galantamine (24–32 mg/day), a traditional “observed cases” (OC) analysis at 3 months was the primary efficacy analysis; this included patients who provided postbaseline data for ADAS-cog/11, CIBIC-plus, or DAD variables at the designated assessment times. More conservative 3 month, intention to treat (ITT) analyses were also performed to examine the robustness of the efficacy results. This included classic ITT analysis (according to assigned treatment and using the last observation available for each patient regardless of whether they took trial medication), and the traditional last observation carried forward (LOCF) analysis (using the last postbaseline observations available for each patient who received treatment). All results discussed are based on OC analysis unless otherwise stated.

Baseline characteristics of the three groups were compared using two way analyses of variance (ANOVA) for continuous variables and generalised Cochran-Mantel-Haenszel tests for categorical variables. Changes from baseline in efficacy variables, vital signs, and body weight were assessed using two tailed, paired t tests. Comparisons of variables between the galantamine and placebo groups employed the following methods: ANOVA for changes from baseline in ADAS-cog subscales, DAD, PSQI, and vital signs; generalised Cochran-Mantel-Haenszel tests for ADAS-cog/11 response rates; and Van Elteren tests for CIBIC-plus. The critical level of significance was set as p≤0.05. The statistical software used in these analyses was SAS version 6.12.

Table 1  Baseline characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Placebo (n=125)</th>
<th>Galantamine 24–32 mg (n=261)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demography: Males/females</td>
<td>58/67</td>
<td>113/148</td>
</tr>
<tr>
<td>Age (y)*</td>
<td>74.6 (0.68)</td>
<td>75.2 (0.45)</td>
</tr>
<tr>
<td>Clinical: Weight (kg)*</td>
<td>68.5 (1.37)</td>
<td>66.1 (0.86)</td>
</tr>
<tr>
<td>Smokers</td>
<td>9 (7.2)</td>
<td>21 (8.0)</td>
</tr>
<tr>
<td>Other active medical conditions</td>
<td>112 (89.6)</td>
<td>235 (90.0)</td>
</tr>
<tr>
<td>ApoE ε4 genotype: Homozygous</td>
<td>14 (13.0)</td>
<td>38 (16.7)</td>
</tr>
<tr>
<td>Heterozygous</td>
<td>56 (51.8)</td>
<td>111 (48.9)</td>
</tr>
<tr>
<td>Total MMSE score*</td>
<td>19.6 (0.32)</td>
<td>19.7 (0.24)</td>
</tr>
<tr>
<td>ADAS-cog/11 score*</td>
<td>24.7 (0.85)</td>
<td>25.6 (0.65)</td>
</tr>
<tr>
<td>Total NPI score*</td>
<td>9.4 (1.01)</td>
<td>9.2 (0.66)</td>
</tr>
<tr>
<td>Total DAD score*</td>
<td>73.0 (1.91)</td>
<td>69.1 (1.42)</td>
</tr>
<tr>
<td>Time since cognitive problem diagnosed (y)*</td>
<td>3.22 (0.19)</td>
<td>3.8 (0.20)</td>
</tr>
<tr>
<td>Time since probable Alzheimer’s disease diagnosed (y)*</td>
<td>0.69 (0.1)</td>
<td>0.71 (0.07)</td>
</tr>
</tbody>
</table>

Data are number (%) of patients, except those marked * which denotes mean (SE).

Results

Five hundred and thirty four patients were screened for the study and 386 were randomised to trial medication, of whom 75% completed the study (fig 1). During week 4, 64 patients remained on the 24 mg/day dose of galantamine, whereas 165 were escalated to the 32 mg/day dose, of whom 40 (24%) reverted to the lower dose during that week. Of the 125 patients remaining on the higher dose by the end of week 4, 103 (82%) completed the study.
Table 2  Efficacy outcomes after 3 months

<table>
<thead>
<tr>
<th>Assessment</th>
<th>Placebo</th>
<th>Galantamine 24–32 mg/day</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Classic ITT</td>
<td>ITT (LOCF)</td>
</tr>
<tr>
<td>ADAS-cog/11 (mean (SE) change from baseline)</td>
<td>+0.7 (0.47)</td>
<td>+0.6 (0.45)</td>
</tr>
<tr>
<td>ADAS-cog/13 (mean (SE) change from baseline)</td>
<td>+0.7 (0.52)</td>
<td>+0.7 (0.51)</td>
</tr>
<tr>
<td>No (%) ADAS-cog/11 responders</td>
<td>+4 points improvement</td>
<td>27 (22.0)</td>
</tr>
<tr>
<td>CIBIC-Plus (No (%)) patients in each category</td>
<td>&gt;Markedly improved</td>
<td>0 (0)</td>
</tr>
<tr>
<td></td>
<td>2=Moderately improved</td>
<td>1 (0.8)</td>
</tr>
<tr>
<td></td>
<td>3=Minimally improved</td>
<td>23 (18.5)</td>
</tr>
<tr>
<td></td>
<td>4=No change</td>
<td>54 (43.5)</td>
</tr>
<tr>
<td></td>
<td>5=Minimally worsened</td>
<td>36 (29.0)</td>
</tr>
<tr>
<td></td>
<td>6=Moderately worsened</td>
<td>9 (7.3)</td>
</tr>
<tr>
<td></td>
<td>7=Markedly worsened</td>
<td>1 (0.8)</td>
</tr>
<tr>
<td>NPI (mean (SE) change from baseline)</td>
<td>+0.5 (0.64)</td>
<td>+0.5 (0.65)</td>
</tr>
<tr>
<td>DAD (mean (SE) change from baseline)</td>
<td>−5.1 (1.17)</td>
<td>−5.2 (1.18)</td>
</tr>
</tbody>
</table>

*p <0.05; **p <0.01; ***p <0.001 vs placebo.

ITT = Intention to treat analysis; LOCF = Last observation carried forward analysis; OC = Observed cases analysis.

Of the patients who continued on, or reverted to, the lower dose during week 4, 72 (69%) completed the study. The baseline characteristics of the treatment groups were comparable (table 1). The proportions of patients taking concomitant medication (most commonly analgesics) during the double blind phase of the study were comparable between treatment groups (89%, placebo and 88%, galantamine). A slightly greater proportion of patients in the galantamine group (33% (85/261)) compared with those receiving placebo (25% (31/125)) took concomitant psychotropic medications.

Protocol deviations occurred in 38 (10%) of randomised patients. Use of prohibited medications (20 cases) was the most common problem.

**Efficacy**

Galantamine treated patients showed significantly superior cognitive function when compared with placebo treated patients, the mean treatment effect in favour of galantamine being 1.1 points (p <0.05) at 1 month and 1.9 points (p =0.002) at 3 months on the ADAS-cog/11 subscale (fig 2). These treatment differences were due to ADAS-cog/11 scores significantly improving in baseline galantamine treated patients at both time points (p <0.001 in both cases), while not changing significantly in the placebo group (table 2). At 3 months, there was no difference in those who received 32 mg/day or 24 mg/day galantamine in improvement from baseline in ADAS-cog/11 during the fixed dose treatment period (mean (SE) 1.4 (0.57), n =99, and 1.5 (0.54), n =71, ADAS points respectively). All of these findings were confirmed by both ITT analyses. Galantamine also produced a significantly better outcome than placebo on the ADAS-cog/13 subscale (p =0.004) and ADAS-cog/11 responder rates (p =0.02, table 2).

The overall clinical response to galantamine at 3 months, as measured by CIBIC-plus, was significantly better than with placebo (p =0.003; significance confirmed by both ITT analyses, table 2). Only 21% of patients on galantamine (n =170) deteriorated compared with 37% (n =111) of those in the placebo group. Furthermore, galantamine produced significant benefits on activities of daily living as indicated by a drug–placebo difference in the mean change from baseline of 4.3 points in the total DAD score (p = 0.004; significance confirmed by both ITT analyses, table 2). Only 21% of patients on galantamine (n =170) deteriorated compared with 37% (n =111) of those in the placebo group. Functional performance was preserved in galantamine treated patients, as indicated by a DAD score that was not significantly different from baseline. This preservation of functional activity was seen regardless of whether patients completed the study on a dose of 32 mg/day or 24 mg/day (mean (SE) changes of 0.6 (1.21), n =99, and −0.5 (1.24), n =73, respectively). By contrast, the decline in total DAD score for the placebo group was statistically significant (p <0.001 for OC and ITT analyses). At 3 months, the decline in total DAD score for the placebo group was statistically significant (p <0.001 for OC and ITT analyses). At 3 months, the decline in total DAD score for the placebo group was statistically significant (p <0.001 for OC and ITT analyses).
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Vects of a flexible galantamine dose in Alzheimer’s disease

in severity in both treatment groups, and the reduced during the maintenance phase (fig 5).

ference between the placebo and galantamine greater than that in the placebo group. The dif-

of the study in the galantamine group was adverse events during the dose escalation phase

Figure 5 Incidence of adverse events by phase of study.

Table 3 Number (%) patients with adverse events occurring at least 5% more often with galantamine than with placebo

<table>
<thead>
<tr>
<th>Adverse event</th>
<th>Placebo (n=125)</th>
<th>Galantamine (n=261)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nausea</td>
<td>14 (11.2)</td>
<td>84 (32.2)</td>
</tr>
<tr>
<td>Dizziness</td>
<td>5 (4.0)</td>
<td>39 (14.9)</td>
</tr>
<tr>
<td>Vomiting</td>
<td>5 (4.0)</td>
<td>38 (14.6)</td>
</tr>
<tr>
<td>Anorexia</td>
<td>3 (2.4)</td>
<td>31 (11.9)</td>
</tr>
<tr>
<td>Somnolence</td>
<td>1 (0.8)</td>
<td>20 (7.7)</td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>2 (1.6)</td>
<td>18 (6.9)</td>
</tr>
<tr>
<td>Agitation</td>
<td>1 (0.8)</td>
<td>16 (6.1)</td>
</tr>
<tr>
<td>Any adverse event</td>
<td>79 (63.2)</td>
<td>229 (86.2)</td>
</tr>
</tbody>
</table>

Figure 4 Mean change from baseline in DAD cluster scores at 3 months.

months, significant differences between the galantamine and placebo groups were also seen for each DAD cluster (fig 4): initiation, planning, performance as well as both basic and instrumental activities of daily living (p<0.05 for all OC and LOCF-ITT analyses), except for DAD-leisure (p=0.06 in OC analysis). At 3 months, the mean NPI scores had not significantly changed from baseline for both the galantamine and placebo groups (table 2).

SAFETY

Galantamine was well tolerated. Adverse events occurring at least 5% more often in either galantamine group than the placebo group are listed in table 3. The incidence of adverse events during the dose escalation phase of the study in the galantamine group was greater than that in the placebo group. The difference between the placebo and galantamine groups in the frequency of adverse events was reduced during the maintenance phase (fig 5).

Most adverse events were mild to moderate in severity in both treatment groups, and the proportion of serious adverse events was comparable (6% with placebo, 8% with galantamine). There were two deaths during the study, both in the placebo group. Discontinuations due to adverse events were more common in galantamine treated patients than in those receiving placebo (fig 1). The events most commonly associated with the discontinuations during galantamine therapy were nausea (13% (33/261)), vomiting (6% (15/261)), dizziness (5% (13/261)), and anorexia (4% (11/261)). There was only one report of muscle weakness—in a galantamine treated patient—which was judged as probably unrelated to treatment.

Total PSQI scores did not change significantly in either treatment group during the study (a mean decrease of only 0.2 points in each case). There were no consistent trends or clinically important differences between treatment groups in blood chemistry, haematology, urinalysis, pulse rate, blood pressure, or ECG measurements during the study.

Discussion

This study showed that patients with mild to moderate Alzheimer’s disease receiving galantamine experienced statistically significant benefits in cognition and global function after 4 and 12 weeks treatment. The clinical relevance of these findings was emphasised by the improvements seen in both the ADL measure and CIBIC-plus in the galantamine group on both observed case and ITT analyses. No benefit was found on behaviour, likely owing to the short duration of the trial and the exclusion of patients with behavioural problems at baseline. Although it is not possible to draw direct comparisons with another trial, the duration of the study may be important as in a similar, but longer, 5 month study the placebo group showed a significant increase in behavioural symptoms, whereas the galantamine group remained stable.

Although the study conforms to the usual requirements for valid studies, in our view, the short duration was the study’s main limitation. However, the duration of galantamine treatment was long enough to demonstrate a therapeutic effect, even with a fairly rapid dose escalation, and to test our primary hypothesis about the effects of flexible dosing on tolerability. We used the OC analysis at 3 months as the primary analysis so that we could capture the effect of treatment on completers who had been maintained on 24 or 32 mg/day galantamine, thus reflecting experience in clinical practice in patients who comply with treatment recommendations. Although the OC analysis may bias the results towards a larger treatment effect, especially as the drop out rate was higher in the galantamine group than in the placebo group, our findings are confirmed by more conservative ITT analyses.

Hitherto the lack of impact of cholinesterase inhibitors on function has been used to undermine the clinical importance of their cognitive effects, especially given the contribution of functional decline to caregiver burden and the cost of care. The general consensus view...
is changing as functional improvement has been reported in other studies of galantamine1, and at least for instrumental ADL, with donepezil.7 Whether the preservation of basic ADL seen in this study constitutes an advantage of galantamine over other treatments cannot be established without direct comparative studies, but evidence now exists that functional benefit can be demonstrated with cholinesterase inhibition.

The adverse events associated with galantamine in this study were generally those expected from cholinergic stimulation, and similar to those reported with other cholinesterase inhibitors.8 9 10 11 12

The adverse events were of mild to moderate severity, and generally resolved prima- ry during the dose escalation phase and may be reduced further by using a slower dose escalation.13 In the present study, there was no evidence of hepatotoxicity, muscle weakness, or sleep disturbance, as has been reported for other cholinergic treatments.8 9 10 11

Of the patients escalated to the higher dose of galantamine during week 4, most (76%) remained on this dose during that week, and most of these (82%) completed the study. Although flexible dose escalation seemed to allow investigators to select patients who would tolerate the higher dose, patients on 32 mg/day did seem to gain additional cognitive benefit over those who received 24 mg/day (mean ADAS-cog improvement over baseline 1.4 and 1.5 points respectively). This is consistent with a 6 month placebo controlled study that did not show any clinically or statistically significant differences on primary efficacy variables between the 24 and 32 mg/day doses of galantamine.14

This randomised, placebo controlled trial shows that galantamine produces cognitive and functional benefit in patients with mild to moderate Alzheimer’s disease. Galantamine was well tolerated with no evidence of hepatoxicity, muscle weakness, or sleep disturbance. These results suggest that galantamine will have a useful role in the treatment of Alzheimer’s disease.

Clinical investigators: Australia, Boundy K, Clarinette R; Canada, Accossato AR, Brown J, Caddy BH, Molloy D, Rockwood K, Wherrett J; New Zealand, Davidson G, Gilchrist N, South Africa, Browne G, Fritz V, Holmes R, Leong S, Laban P; France, Laborde G, Leonard Kingdom, Bonhala C, Ghosh M, Mahmood A, McMillan H, Pathy M, Bapuri Rao V, Roissier J, Tym E, Williams A, United States, Aronson S, Blan chette P, De Priest M, Ester L, Forrerd W, Frey J, Green R, Landlshoem R, Mintner J, Nakra R, Palj J, Pendlebury W, Rymer M, Shua-Hann J, Solomon P, Thien S, Thein J, Timon J. This study was supported by funding from Janssen Research Foundation, Beers, Belgium. We are grateful to Man Lagados of Oxford University who wrote an initial draft based on Clinical Trial Report N133910, which was prepared by Rolando Gutterres of Janssen Research Foundation. We have reviewed the clinical trial report and related analyses, and are guarantors for the results presented. This report includes data generated by protocol GAL-INT-2, sponsored by Janssen Research Foundation. KR has received honoraria for lectures, and three grants for clinical studies. JM has received compensation from Janssen for lectures, consulting and grants/studies. DW has received honoraria for conference presentations and for consulting activities, and DW and JM do not own any stock or options in Janssen Pharmaceutica. TW and LT are employees of Janssen Research Foundation.

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