Eptastigmine led to cognitive, clinical, and functional benefits in Alzheimer's disease


Question
In patients with moderate to moderately severe Alzheimer’s disease (AD), how effective and safe is eptastigmine?

Design
Randomised, double blind, placebo controlled trial with 24 weeks of follow up.

Setting
36 centres in the US and Europe.

Patients
491 patients (mean age 71.6, 63% women) who met the diagnostic criteria for AD established by the NINCDS-ADRDA and described by DSM-IV; and who had a cognitive deficit for ≥6 months, a Mini-Mental State Examination Scale score between 10 and 22, a Global Deterioration Scale rating of 4 or 5, a modified Hachinski ischemic score <3, and a Hamilton Depression Scale score <18. Exclusion criteria were other neurological or psychiatric disorders; renal, hepatic, or cardiovascular diseases; peptic ulcer; bronchial asthma; neutropenia; deficiencies in thyroxine, vitamin B12, or folate; or treatment with medication known to affect the central nervous system. 86% completed the study.

Intervention
164 patients were assigned to placebo, 166 to 15 mg 3 times daily of eptastigmine, and 161 to 20 mg 3 times daily of eptastigmine for a period of 24 weeks. Eptastigmine was given using a 4 week stepwise dose escalation starting from 5 mg 3 times daily.

Main outcome measures
Changes in cognition (Alzheimer’s Disease Assessment Cognitive Subscale [ADAS-Cog]), global function (Clinician’s Interview-Based Impression of Change Plus [CIBIC-Plus]), and activities of daily living (Instrumental Activities of Daily Living Scale [IADL]).

Main results
Data for 463 patients were included in the analysis with the last outcome carried forward for 15. The table shows the mean change from baseline to week 24 on the ADAS-Cog, CIBIC-Plus, and IADL. 48% of patients in the higher dose group had adverse effects compared with 54% in the lower dose group and 49% in the placebo group. There was a dose dependent transient and mild neutropenic effect associated with eptastigmine treatment.

Conclusion
20 mg 3 times daily of eptastigmine led to cognitive, clinical, and functional benefits in patients with moderate to moderately severe Alzheimer’s disease but with increased risk of adverse haematological effects.

Mean changes from baseline to week 24 in patients with AD assigned to placebo or 15 mg or 20 mg 3 times daily of eptastigmine

<table>
<thead>
<tr>
<th></th>
<th>Placebo</th>
<th>15 mg</th>
<th>20 mg</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADAS-Cog</td>
<td>Drug-placebo adjusted difference (95% CI)</td>
<td>2.5 (0.9 to 4.1)</td>
<td>1.9 (0.7 to 3.1)</td>
</tr>
<tr>
<td>CIBIC-Plus</td>
<td>Mean change</td>
<td>4.4</td>
<td>4.0</td>
</tr>
<tr>
<td>IADL</td>
<td>Drug-placebo adjusted difference (95% CI)</td>
<td>0.1 (0.1 to 0.7)</td>
<td>0.3 (0.1 to 0.6)</td>
</tr>
<tr>
<td></td>
<td>Mean change</td>
<td>1.2</td>
<td>0.8</td>
</tr>
<tr>
<td></td>
<td>Drug-placebo adjusted difference (95% CI)</td>
<td>0.4 (0.2 to 0.6)</td>
<td>0.7 (0.2 to 1.2)</td>
</tr>
</tbody>
</table>

*Not significant. AD = Alzheimer’s Disease; ADAS-Cog = Alzheimer’s Disease Assessment Cognitive Subscale; CIBIC-Plus = Clinician’s Interview-Based Impression of Change Plus; IADL = Instrumental Activities of Daily Living Scale.

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Commentary
The cholinergic hypothesis of memory loss has to contend with the glutamatergic hypothesis of amnesia and dementia.1 These studies by Imbimbo et al and Winblad and Porits are based on these alternative approaches.

Acetylcholinesterase inhibitors (AChEIs) have been the most widely investigated drugs in mild to moderate AD. Measures of their efficacy should tap 4 areas: cognition, functional activity, non-cognitive behaviours, and independent ratings of global change. Compared with placebo, the AChEIs tacrine, donepezil, metrifonate, and rivastigmine show a modest improvement of cognitive function of 2–4 ADAS-Cog points over 6 months, with a return then to baseline values or less.2 Ratings of global impression of change improve 1/3–1/2 point. Non-cognitive dysfunction has rarely been studied, and improvements in activities of daily living (ADL) have been hard to detect, except for rivastigmine.3 Overall, up to one quarter of patients do well on AChEIs, but the majority see little to no benefit. Which quarter improves, we cannot predict. The decrease in burden of care has been studied less.

The study by Imbimbo et al is well designed and was done with a low dropout rate. Unfortunately, the magnitude of the cognitive benefit was in the lower half of the range reported for other AChEIs and the number with the change normally considered clinically significant (≥4 ADAS-Cog points) is not reported. But eptastigmine joins rivastigmine in showing substantial benefit on ADL. The dose response effect and the consistency of modest improvements in cognition and daily functioning supporting the clinical relevance of the results. Although the cholinergic tolerability of eptastigmine was favourable, 4.5% of patients developed a mild transient neutropenia, which was dose dependent and included 1 case of severe neutropenia in the eptastigmine 20 mg group (3 times daily). It is surprising that patients with neutropenia continued in the trial and that there is no comment on incident infections. Another patient on eptastigmine, 20 mg (3 times daily), however, developed aplastic anaemia, a risk of 0.3% (95% CI 0.01% to 1.7%), whereas the risk in the general population is 2 per million per year. The risk on eptastigmine, similar to that of clozapine, will probably limit the use of the (commentary continued on page 113)
Memantine led to functional improvement and reduced care dependence in severe dementia


Question
In patients with moderately severe to severe primary dementia, how effective and safe is memantine?

Design
Randomised, double blind, placebo controlled trial with 12 weeks follow up.

Setting
7 trial centres in Latvia.

Patients
167 care dependent inpatients with moderately severe to severe dementia defined by DSM-III-R criteria (49% Alzheimer’s disease, 51% vascular dementia) and with severity assessed by the Global Deterioration Scale (stages 5–7) and the Mini-Mental State Examination (<10 points). Exclusion criteria included duration of dementia <12 months; central nervous system active drugs within the previous 14 days; severe, chronic, or terminal diseases; impaired thyroid function, lowered B12 blood levels of severity, leave key issues unaddressed. The reported benefits and tolerability need to be established in a much longer duration trial, with health economic analysis. Adverse effects (not described) and serious adverse effects (mainly cardiac deaths) were equivalent in the two groups. Memantine seems useful not only in severe AD but also in vascular and mixed dementias, and in those without substantive behavioural problems (neuroleptics, hypnotics, and some antidepressants were excluded in trial patients), found in approximately 40% of patients with AD. Although glutamate antagonists may induce transient psychoses, memantine is thought not to do so. These 2 contrasting pharmacological approaches to AD, here studied at different levels of severity, leave key issues unanswered: who benefits? Is response genotype specific? How substitutable and valuable is a 2 point drop in carer dependence (memantine) in progressive dementia? Are there longer term side effects? Are skills regainable without rehabilitation? And how do the modest gains translate into quality adjusted life years and reduce burden of care and costs? In the future, non-cognitive symptoms should be assessed because they increase disease progression and disability, predict high family burden and costs, and are the primary cause of institutionalisation.1 Memantine’s possible advantage of neuroprotection suggests that treatment combinations with AChEIs might enhance the benefits of only 1 approach.

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Main results
At 12 weeks, more patients receiving memantine had a positive treatment response (defined as 3 categories of improvement on the CGI-C) than those receiving placebo (p<0.001) (table). There was a 3.1 point improvement on the BGP subscore “care dependence” for those in the memantine group compared with a 1.1 point improvement for those in the placebo group (p=0.016). A coincidental response of 3 categories of improvement on the CGI-C and an improvement of ≥15% on the BGP subscore was observed in more patients receiving memantine than those receiving placebo (n=151) (table).

Conclusion
Memantine led to functional improvement and reduced care dependence in patients with severe dementia.

Memantine v placebo at 12 weeks in severe dementia*

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Memantine</th>
<th>Placebo</th>
<th>RBI (95% CI)</th>
<th>NNT (CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment response on the CGI-C</td>
<td>73%</td>
<td>45%</td>
<td>62% (25 to 114)</td>
<td>4 (3 to 8)</td>
</tr>
<tr>
<td>Coincidental response on the CGI-C and BGP subscore</td>
<td>61%</td>
<td>32%</td>
<td>94% (55 to 186)</td>
<td>4 (3 to 8)</td>
</tr>
</tbody>
</table>

*CGI-C = Clinical Global Impression of Change; BGP = Behavioural Rating Scale for Geriatric Patients. Other abbreviations defined in glossary; RBI, NNT, and CI calculated from data in article.

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