Lamotrigine and lithium are effective maintenance treatments in recently depressed people with bipolar I disorder


METHODS

What are the effects of lamotrigine and lithium for maintenance treatment in recently depressed people with bipolar I disorder?

METHODOLOGY

- Design: Randomised controlled trial.
- Allocation: Unclear.
- Blinding: Double blind.
- Follow up period: 18 months.

PATIENTS

- 463 adults with bipolar I disorder (mean age range 42 to 44 years), who were stabilised with lamotrigine during an 8–16 week open label phase (n = 966). Main inclusion criteria: current or recent (past 60 days) DSM-IV major depressive episode; >1 manic or hypomanic episodes, and >1 depressive episodes within 3 years of enrolment. Exclusions: >6 DSM-IV mood episodes in the past year; significant psychiatric or medical comorbidity; epilepsy or suicidal. All other psychotropic medicines were discontinued before randomisation.
- Intervention: Maintenance treatment with lamotrigine (50, 200, or 400 mg/day), lithium (serum levels 0.8–1.1 mEq/l) or placebo.
- Outcomes: Time to intervention for any mood episode; adverse events.
- Patient follow up: 95%.

MAIN RESULTS

Both lamotrigine (200 or 400 mg/day) and lithium significantly increased the time to intervention for any mood episode compared with placebo (200 days with lamotrigine v 170 days with lithium v 93 days with placebo; p = 0.029 for both comparisons v placebo). There was no significant difference in the time to intervention between lamotrigine and lithium (p = 0.915). 11% of participants withdrew due to adverse effects (see table for most common adverse events).

CONCLUSIONS

Lamotrigine and lithium are better than placebo for preventing mood episodes in people with bipolar I disorder.

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NOTE

There were initially three lamotrigine treatment groups; enrolment was stopped in the 50mg/day and 400 mg/day groups during the study because of poor enrolment. Efficacy data for lamotrigine was pooled for the 200 mg and 400 mg/day groups, a decision that had been made a priori. Authors point out that participants who were intolerant or unresponsive to lamotrigine would have been excluded during the prerandomisation stabilisation period, and this may have biased results.

Table

<table>
<thead>
<tr>
<th>Common adverse events occurring during treatment, n (%)</th>
<th>Lamotrigine (n = 169)</th>
<th>Lithium (n = 120)</th>
<th>Placebo (n = 121)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Headache</td>
<td>30 (18)</td>
<td>23 (19)</td>
<td>25 (21)</td>
</tr>
<tr>
<td>Nausea</td>
<td>28 (17)</td>
<td>24 (20)</td>
<td>15 (12)</td>
</tr>
<tr>
<td>Tremor</td>
<td>9 (5)</td>
<td>20 (17)†</td>
<td>6 (5)</td>
</tr>
<tr>
<td>Rash</td>
<td>12 (7)†</td>
<td>5 (4)</td>
<td>3 (2)</td>
</tr>
<tr>
<td>Dizziness</td>
<td>21 (12)</td>
<td>13 (11)</td>
<td>12 (10)</td>
</tr>
</tbody>
</table>

†p<0.05 versus lithium.

* p<0.05 versus placebo.

Commentary

This study investigates treatment with lamotrigine after an acute response to the drug in recently depressed subjects with bipolar disorder. This issue is clinically important, given the depressive burden of bipolar disorder. The design is that of a randomised double blind placebo controlled clinical trial comparing five groups: three doses regimens of lamotrigine (50, 200, 400 mg/day), lithium and placebo. Randomisation took place after 463 patients were stabilised on lamotrigine during an open label phase of treatment for acute major depression. Data analyses show lamotrigine and lithium being superior to placebo in preventing additional treatment interventions (the primary outcome) because of the occurrence of a mood episode.

This study is not a classic prophylaxis study; in that design, patients who have become euthymic in any manner (recently depressed, recently manic, or euthymic for a long time due to natural history) enter the study. This design is rather one of relapse prevention (or an “enriched” design), in which all patients who were randomised to maintenance treatment initially responded to lamotrigine (with or without other psychotropics) for an acute major depressive episode. In other words, the generalisability of these findings is limited to such patients, and not all patients who reach the maintenance phase in other ways.

Also, lithium is included as an active control—that is, as a test of assay sensitivity. As the study is not designed and powered to assess lithium efficacy, definitive conclusions about lithium’s efficacy cannot be made. Thus, it would not exactly follow from these data that lithium is not effective in prevention of depressive episodes (one of the secondary analyses). Rather, the primary conclusion one could draw is that lamotrigine is effective in delaying relapse of mood episodes, particularly of the depressive subtype, with the above constraints of generalisability.

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