Acid, motility, and ulcers: a comparison of cisapride with placebo in the prevention of duodenal ulcer relapse

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Abstract
In a single centre double blind study of 66 patients, the value of cisapride (10 mg twice daily) was compared with placebo in the prevention of duodenal ulcer relapse. Patients who remained ulcer free attended for clinical review every two months and had a mandatory endoscopy at 0, 4, 8, and 12 months or if symptoms suggestive of ulcer recurrence developed. The 12 month crude relapse rates (that underestimate the probability of ulcer recurrence) showed that cisapride was superior to placebo (34% (11/32) relapsed on cisapride v 68% (23/34) on placebo, p=0.007). This finding was confirmed using lifetable analysis, with a 35% reduction (95% confidence intervals 10-59%, p<0.05) in the proportion of ulcer relapses in patients who had received cisapride compared with those treated with placebo. These results are similar to those reported in maintenance trials of H$_2$ receptor antagonists analysed by the same method. Drug related adverse clinical events were mainly trivial, but led to three patients on cisapride and one on placebo withdrawing from the trial.

(Gut 1993; 34: 1042–1046)

Although duodenal ulcers rarely form in patients with achlorhydria, the widely held belief that all patients with duodenal ulcer disease secrete excessive amounts of acid is incorrect. About two thirds of duodenal ulcer patients have normal values of stimulated gastric secretion. Therefore follows that the therapeutic success of drugs that reduce gastric secretion might be explained if duodenal ulcers resulted from duodenal rather than gastric hyperacidity, possibly as a result of inadequate clearance of even normal amounts of acid from the bulb. We have suggested that properly coordinated motility might be of crucial importance in promoting acid clearance, and could therefore be an unrecognised factor in the pathogenesis of duodenal ulceration. If this hypothesis was correct, drugs that modulate upper gastrointestinal motility may have a role in the treatment of duodenal ulcer patients.

Nearly 20 years ago, Moshal presented data from a controlled trial that investigated the value of metoclopramide in the prevention of duodenal ulcer symptoms. Although maintenance treatment with metoclopramide significantly reduced the crude relapse rate of ulcer symptoms compared with placebo, Moshal’s work (which was published shortly before the first placebo controlled trial of an H$_2$ receptor antagonist), has largely been forgotten. The aim of our study was to find out if the more potent prokinetic drug cisapride could prevent duodenal ulcer relapse. Cisapride (Prepulsid, Janssen Pharmaceutica, Beerse, Belgium) stimulates coordinated upper gastrointestinal motility particularly in patients with disordered motor patterns. Cisapride does not affect gastric secretion either in animals or man.

Patients and methods
Patients aged between 18 and 75 years presenting to the Royal Hallamshire Hospital, Sheffield with duodenal ulceration were treated medically and had an endoscopy to confirm ulcer healing. They were then immediately entered into a single centre double blind trial in which they were randomised to receive either oral cisapride 10 mg twice daily or an identical placebo tablet for up to 55 weeks. Ulceration associated with the use of non-steroidal anti-inflammatory drugs or steroids was excluded from the study, as were patients with a high alcohol intake (>40 U/week), or a history of surgery or drug use which might interfere with gastric secretion or motility. Patients with severe cardiorespiratory, renal, hepatic, neurological, or malignant disease were also excluded, together with women of childbearing age not using a reliable means of contraception.

Patients were reviewed every two months and the duodenum inspected endoscopically at the beginning of the trial, at approximately 4, 8, and 12 months, and if the patient developed recurrent dyspepsia that persisted after three days of simple antacid treatment (Maalox, maximum 30 tablets per month). Treatment failure was defined as evidence by endoscopy of a recurrent duodenal ulcer crater. At each visit, any unused tablets were returned to check on compliance and a new prescription issued. Adequate compliance was defined as ingestion of at least 70% of the tablets prescribed.

This study received approval from our local Ethical Committee and all patients provided written, informed consent.

STATISTICAL METHODS
Lifetable analysis - In a study of this kind, it is rarely possible to follow up every patient until they either relapse or successfully complete the trial. To overcome this difficulty, the estimated cumulative proportion of ulcers relapsing on each treatment was calculated using life tables and the log rank test as described by Peto et al., because this method uses the actual duration of the ulcer free period for each patient and allows
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Results

Of the 68 patients entered, two were retrospectively excluded from analysis: one patient had an incompletely healed ulcer at the time of entry, and the other had a substantial prepyloric ulcer associated with small duodenal erosions. Thirty two of the remaining patients were randomised to receive cisapride, and 34 to receive placebo.

PATIENT CHARACTERISTICS

Patients randomised to each treatment were well matched for age, sex, smoking habit, alcohol intake, the duration of their previous dyspeptic symptoms, and their previous responsiveness to medical treatment (Table I). Most acute ulcers had been healed by H2 receptor antagonists, and only one patient (randomised to placebo) had received bismuth substitute before entering the trial. Three patients (two randomised to cisapride) had originally presented with large or multiple chronic duodenal ulcers associated with small solitary areas of slough on either the incisura or in the prepyloric region. A check on the number of tablets returned at each visit showed that all patients took at least 70% of the tablets prescribed, and that the median (interquartile range) number of antacid tablets consumed per week was similar in both groups (cisapride 2.2 (0.1–5.7); placebo 1.9 (0.3–3.6); p=0.76, Mann-Whitney U test).

WITHDRAWALS

In addition to the two patients excluded from analysis, another 13 patients were withdrawn from the study (Table II). Three failed to comply with the protocol restrictions; one patient continued to use H2 receptor antagonists until she was withdrawn three days after entry and another was unable to keep his alcohol consumption below 40 U/week; a third (asymptomatic) patient withdrew her consent within a month of randomisation. In addition, 10 patients were withdrawn because they either defaulted from follow up (n=3, all had moved away from the Sheffield area), or because they developed an adverse clinical event (n=7) which led to a change in their treatment (Table III). The reported occurrence of minor drug related symptoms (~that is, not severe enough to warrant withdrawal from the study), was similar in each group (Table III). Ten of the 13 patients withdrawn were in the group receiving cisapride (Table II).

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**Table I**  Patient characteristics on entry

<table>
<thead>
<tr>
<th></th>
<th>Cisapride (n=32)</th>
<th>Placebo (n=34)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years), mean (SD)</td>
<td>46.8 (13)</td>
<td>47.9 (14)</td>
</tr>
<tr>
<td>Sex (% men)</td>
<td>72%</td>
<td>74%</td>
</tr>
<tr>
<td>Smokers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>16</td>
<td>16</td>
</tr>
<tr>
<td>&gt;10/day</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Alcohol (units/week)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>range*</td>
<td>3 (0-12)</td>
<td>6 (0-12)</td>
</tr>
<tr>
<td>Duration of ulcer symptoms before randomisation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;3 months</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>3-12 months</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>1-2 years</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>&gt;2 years</td>
<td>16</td>
<td>15</td>
</tr>
<tr>
<td>Healing drug</td>
<td></td>
<td></td>
</tr>
<tr>
<td>H2 receptor antagonist</td>
<td>28</td>
<td>24</td>
</tr>
<tr>
<td>Bismuth substitute</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Cisapride</td>
<td>4</td>
<td>9</td>
</tr>
<tr>
<td>Duration of healing treatment before randomisation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;8 weeks</td>
<td>24</td>
<td>28</td>
</tr>
<tr>
<td>8-12 weeks</td>
<td>7</td>
<td>3</td>
</tr>
<tr>
<td>3-6 months</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>&gt;6 months</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

*One unit of alcohol=one half pint beer/lager, one glass of wine, or a single measure of spirits. Cisapride v placebo, p=0.70 (Mann-Whitney U test).

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**Table II**  Patient progress

<table>
<thead>
<tr>
<th></th>
<th>Cisapride</th>
<th>Placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Randomised</td>
<td>34</td>
<td>34</td>
</tr>
<tr>
<td>Exclusions</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Valid entry</td>
<td>32</td>
<td>34</td>
</tr>
<tr>
<td>Withdrawn</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Protocol violation</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Defaulted*</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Drug related adverse events</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td>Other adverse events</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>

*Traced and interviewed by telephone >one year after leaving trial. All three patients had remained asymptomatic and none had received further treatment for dyspepsia.
ULCER RECURRENCE

Of the 34 recurrent duodenal ulcers detected, 23 (68%) occurred in patients taking placebo, most recurrences being picked up at the scheduled 4, 8, and 12 month endoscopies (15/23 placebo, 7/11 cisapride). Interestingly, although relapse on maintenance cisapride was almost invariably symptomatic (10/11), one third (8/23) of the placebo relapses were silent. The crude 12 month relapse rate was lower in patients treated with cisapride (11/32, 34%) compared with those who received placebo (23/34, 68% (χ²=7-306, p=0.007)). It should be emphasised, however, that the crude rate is probably an underestimate of the true proportion of recurrent ulcers, because it assumes that patients who were withdrawn (and therefore not followed endoscopically for the entire 12 months) did not develop an ulcer relapse.

A more accurate analysis of ulcer recurrence can be derived using lifetable analysis. Maintenance treatment with cisapride was associated with a significant increase in ulcer free survival at week 38 (Table IV), and a 35% increase in ulcer free survival at week 51 (estimated proportion relapsing on placebo=72% (95% CI 55–88%) v cisapride=37% (95% CI 19–55%), p<0.05)). However, the particularly wide 95% confidence interval for the difference between the percentage of cisapride and placebo treated patients estimated to remain ulcer free at 55 weeks (Table IV), reflects the inherent inaccuracy of this portion of the survival curve when the number of patients remaining was small.

The figure illustrates the cumulative ulcer free survival curves of patients randomised to each drug. As stated earlier, it is easy to misinterpret the unreliable later part of the curve (for example, the sharp decrease in the estimated ulcer free survival of patients treated with cisapride that occurred at week 52 resulted from a single ulcer recurrence). To avoid this kind of error, survival analysis was truncated at week 51, the point at which an adequate number (n>10) of patients were still present. Lifetable analysis of the truncated survival curve (which excludes one placebo and one cisapride recurrence) also indicates that patients treated with cisapride were less likely to develop a recurrent ulcer over the study period than those receiving placebo (χ²=3.673, p=0.055 log rank test).

Discussion

The results of this single centre pilot study show that maintenance treatment with cisapride 10 mg twice daily can reduce the likelihood of duodenal ulcer relapse. Lifetable analysis shows that 37% (95% CI 19–55%) of patients receiving cisapride are liable to develop a recurrent ulcer after 51 weeks of treatment, compared with 72% (95% CI 55–88%) of those randomised to placebo (complete separation of 95% confidence intervals). This estimated rate of ulcer recurrence is similar to that reported in lifetable analyses of other maintenance trials using both cimetidine 400 mg once daily (median annual recurrence reported=31%, range 21–55%)6,24–28 and ranitidine 150 mg once daily (median annual recurrence reported=35%, range 18–49%).21,24,27,29–31 Although other reports of maintenance treatment with H₂ receptor antagonist drugs have yielded superior results (15–39% annual recurrence),22,23,32–37 these studies only reported crude relapse rates (which underestimate the true rate of ulcer recurrence), or contained extraordinary inconsistencies in the presentation and interpretation of the trial data.33,37

A direct comparison of the relative efficacy of ulcer maintenance drugs is hampered by discrepancies in the reported frequency of placebo related ulcer relapses, presumably because of geographical variation in the aggressiveness of duodenal ulcer disease. To overcome this problem, it is necessary to view the apparent success of an active drug in the context of the parallel placebo response. After 51 weeks of treatment, cisapride reduced the likelihood of duodenal ulcer relapse by 35% (95% CI 10–59%) compared with placebo (p<0.05). Comparable data from H₂ receptor antagonist maintenance trials are limited, but would seem to be similar for both cimetidine 400 mg once daily (estimated reduction in duodenal ulcer relapse rate compared with placebo ranges from 26%26 to 37%36), ranitidine 150 mg once daily (estimated reductions of 24%,30,39%30 and 48%30) and famotidine 20–40 mg once daily (estimated 32–34% reduction in relapse rate).35

Most ulcer maintenance trials, including this study, contain several potential methodological weaknesses that merit further discussion. Firstly, because patients who developed recurrent dyspepsia had another endoscopy before their next scheduled endoscopy, an unacceptable degree of bias could be introduced if the ulcer relapse rates associated with two drugs were identical, but relapses on one treatment resulted in more severe symptoms than those on the other. As some H₂ receptor antagonist maintenance trials have reported a greater proportion of

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**TABLE IV Relative efficacy of cisapride and placebo in preventing duodenal ulcer relapse after 6, 9, and 12 months of treatment**

<table>
<thead>
<tr>
<th>Duration (weeks)</th>
<th>Cisapride (%) CI</th>
<th>Placebo (%) CI</th>
<th>Difference (%) CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>26</td>
<td>+14% (−11 to +40%)</td>
<td>+27% (+2 to +52%)</td>
<td>+13% (+1 to +50%)</td>
</tr>
<tr>
<td>38</td>
<td>+19% (−7 to +54%)</td>
<td>+29% (+1 to +61%)</td>
<td>+10% (−7 to +35%)</td>
</tr>
<tr>
<td>51</td>
<td>+35% (+10 to +59%)</td>
<td>+42% (−8 to +96%)</td>
<td>+7% (+1 to +37%)</td>
</tr>
<tr>
<td>60</td>
<td>+26% (−9 to +60%)</td>
<td>+42% (−8 to +96%)</td>
<td>+16% (−7 to +41%)</td>
</tr>
</tbody>
</table>

Positive values indicate an increased estimated % ulcer free survival with cisapride treatment compared with placebo; negative values indicate decreased survival with cisapride.

*Denotes statistical significance at least at the 5% value.
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symptomatic ulcer recurrences in patients randomised to placebo, it would be easy to overestimate the placebo relapse rate. Secondly, the definition of treatment failure, which included both symptomatic and asymptomatic recurrences needs to be discussed. It has been argued that the detection rate of asymptomatic recurrences depends highly on the frequency of scheduled endoscopic examinations, and that infrequent endoscopy might allow some asymptomatic ulcers to heal without being detected. If, as intimated earlier, the proportion of asymptomatic recurrences was greater in the active treatment group, this could also lead to bias, and underestimation of the frequency of cisapride recurrences relative to placebo.

Although worthy of consideration, we feel that neither of these objections challenge the conclusions of this study. Endoscopic surveillance of treated and untreated asymptomatic ulcers has indicated that 75% persist unhealed over many months, so it is unlikely that an endoscopy every four months would have missed many. Moreover, the proportion of ulcer recurrences detected at unscheduled endoscopies was the same in cisapride and placebo treated patients (35–36%) and in contrast with the H2 receptor antagonist maintenance trials, symptomatic recurrence in this study was particularly common in the group randomised to cisapride, not placebo. If anything, this would suggest that we actually overestimated the frequency of cisapride relapses, and underestimated the placebo relapse rate. One group of investigators have argued that the inclusion of asymptomatic recurrences in the analysis of ulcer relapse trials should be discouraged, because these silent ulcers are clinically irrelevant; however, such optimism (based on findings made in a small group of patients with asymptomatic ulceration followed up for between 1 and 13 months) is probably misguided, because one third of duodenal ulcer patients presenting with perforation and side-effect profile in healthy man. DDig Di Sci 1987; 32: 1223–30.


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