Inhibition of nocturnal acidity is important but not essential for duodenal ulcer healing

G Bianchi Porro, F Parente, O Sangaletti

Methods

PATIENTS PART 1

One hundred and thirty patients with active duodenal ulcer (>5 mm in diameter) endoscopically confirmed within the previous four days (except for patients participating in Part 2) were admitted to the study, which was approved by the hospital ethics committee.

Exclusion criteria included: age less than 18 years, concomitant gastric ulcer, upper gastrointestinal haemorrhage, recent peptic ulcer perforation, pyloric stenosis, Zollinger-Ellison syndrome, antiulcer medications (except antacids for symptomatic relief) during the preceding three weeks, pregnancy or lactation, and any serious medical condition capable of influencing the outcome of therapy.

All patients gave their informed verbal consent before admission to the study and were randomly allocated, according to a double blind protocol, either to ranitidine 300 mg at 8 am or ranitidine 300 mg at 10 pm. Patients receiving ranitidine in the morning also took placebo at night, whereas those taking ranitidine at night received placebo in the morning. All tablets were identical in size and appearance and were dispensed in two coded bottles, marked ‘morning’ and ‘night’, each containing sufficient medication for 35 days’ therapy. Patients were also provided with a known number of antacid tablets (Maalox) to be taken for the relief of pain. They were also instructed to record daily on a diary card pain episodes occurring during the day or night and the consequent antacid consumption.

Treatment was for an initial period of four weeks; if the ulcer was endoscopically unhealed at the end of this first period, therapy was continued for a further four weeks and endoscopy performed again at eight weeks. Patients, however, were clinically reviewed every fortnight to assess drug compliance and the effectiveness of therapy.

At treatment randomisation, patients were asked about age, duration of dyspeptic symptoms and social habits. Basal acid output (BAO) and peak acid output (PAO), the latter after stimulation with pentagastrin 6 μg/kg subcutaneously (Pepavlon, ICI, Great Britain) were also determined before therapy in all patients, with the exception of those participating in Part 2 of the study. The χ² test was used to assess the statistical significance of differences in healing rates of the two groups. 95% confidence intervals (CI) for the differences in treatments were also evaluated. Other data were analysed by means of Student’s t test for unpaired samples.

PART 2

This part of the study was performed on 18...
patients (14 men, four women), mean age 36-7 (9-2) (SD) with active duodenal ulcer subsequently admitted to the clinical trial. Each patient was studied on two separate occasions with a free interval of 48 hours. Patients were divided into two groups, each consisting of nine subjects; those in the first group were given one tablet of placebo or ranitidine 300 mg at 8 am while patients in the second group received one tablet of placebo or ranitidine 300 mg at 10 pm according to a randomised single blind scheme. In no case did more than seven days elapse between initial endoscopy and clinical trial entry.

Each patient underwent two 24 h pH-metries on an ambulant basis, but the time of starting pH monitoring was quite different for the two groups: patients receiving the morning dose were admitted to our laboratory before 8 am after an overnight fast, whereas those taking the nocturnal dose were admitted before 5 pm after fasting from 12 am.

After a pharyngeal anaesthesia with 2% xylocaine solution, a miniaturised bipolar glass electrode with a combined reference electrode (model 440 M4, Ingold AG, Switzerland), connected to a portable recorder) Autronic CM 18, Autronic, FRG), was passed through the nasopharynx into the stomach. The measuring tip was positioned, under fluoroscopic control in the gastric corpus, approximately 10 cm below the cardia. The recorder was calibrated at room temperature using commercial buffer solutions with a pH of 7-00, 4-01, and 1-09 (Radiometer, Copenhagen, Denmark) at the beginning of each test. Acidity was measured every five seconds for a total period of 24 hours.

The diet was normal and standardised for all subjects on each study day. Three meals were given (breakfast at 8 30 am, lunch at 12 30 am and dinner at 8 30 pm) for a total daily energy intake of approximately 2000 kcal. Wine was allowed at lunch and dinner, but the patients was invited to drink the same quantity during the two days. Water was freely allowed. Patients returned to the laboratory the next day and the electrode was removed exactly 24 hours after placing. Two separate pH profiles for placebo and ranitidine were constructed for each group of patients by calculating median pH values every 30 minutes. The area under the curve (AUC) of the 24 h pH profiles for each test was calculated by the trapezoid rule, and statistical comparisons of the mean AUC values for placebo v ranitidine were made using Student’s t test for paired data.

**Results**

**PART 1**

Of the 130 patients initially admitted to the study 65 received ranitidine 300 mg in the morning and 65 ranitidine 300 mg at night. The demographic characteristics of these patients are reported in Table I. One hundred and twenty-four patients were evaluable for the determination of healing rates at four weeks and 122 for cumulative analysis of healing at eight weeks. Four patients on the morning regimen and two patients on the bedtime regimen defaulted at week 4; the reasons for withdrawal were: lost to follow up (three), lack of compliance (two), and late endoscopy (one). Two additional patients unhealed at week 4, one on morning therapy and the other one on the nocturnal regimen, refused endoscopy at week 8.

Endoscopic evaluation after four weeks of treatment showed that ulcer healed in 41/61 patients (67%) taking the morning dose and in 47/63 (75%) of those receiving the nocturnal dose (95% CI for the difference: -0-09 +0-25; p ns). After eight weeks the cumulative healing rates were 82% for the morning regimen and 85-5%.

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**Table I. Characteristics of patients entering the trial. No significant differences were reported for any variable between the two groups.**

<table>
<thead>
<tr>
<th></th>
<th>Morning dose (n=65)</th>
<th>Nocturnal dose (n=65)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex: male (n)</td>
<td>46</td>
<td>49</td>
</tr>
<tr>
<td>Mean age (yr) (SD)</td>
<td>41.1 (11.9)</td>
<td>41.8 (12.9)</td>
</tr>
<tr>
<td>Duration of dyspeptic symptoms (months) mean (SD)</td>
<td>94 (68)</td>
<td>100 (65)</td>
</tr>
<tr>
<td>Smokers (%)</td>
<td>72%</td>
<td>66%</td>
</tr>
<tr>
<td>Alcohol consumers (%) (&gt;50 g/d)</td>
<td>28%</td>
<td>25%</td>
</tr>
<tr>
<td>PAO (mmol H+/h) mean (SD)</td>
<td>40.4 (10.3)</td>
<td>40.7 (11.4)</td>
</tr>
</tbody>
</table>

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**Table II. Endoscopic evaluation of ulcer healing after four and eight weeks of treatment. In parentheses are reported the 95% confidence intervals for the differences.**

<table>
<thead>
<tr>
<th></th>
<th>Healing at four wk</th>
<th>Healing at eight wk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ranitidine at morning</td>
<td>41/61 (67%)*</td>
<td>49/60 (82%)*[t]</td>
</tr>
<tr>
<td></td>
<td>CI 95%</td>
<td>CI 95%</td>
</tr>
<tr>
<td></td>
<td>(-0-09 +0-25)</td>
<td>(-0-11 +0-17)</td>
</tr>
<tr>
<td>Ranitidine at night</td>
<td>47/63 (75%)*</td>
<td>53/62 (85-5%)*[t]</td>
</tr>
</tbody>
</table>

*p=0.47; †p=0.74.

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*Figure 1: Median half hourly intragastric pH in nine patients with duodenal ulcer receiving placebo or ranitidine 300 mg at 8 am.*
Inhibition of nocturnal acidity is important but not essential for duodenal ulcer healing

for the nocturnal regimen (95% CI for the difference −0.11 +0.17; p ns, Table II). Similar proportions of patients in the two treatment groups were pain free by week 4 and no significant difference was observed between the two regimens regarding additional antacid consumption.

Among patients completing the trial, compliance was good and similar in both groups. The percentage of drug consumed during the treatment period was never less than 80%. No clinically relevant side effects were observed with either regimen. No factor was shown to significantly affect healing of duodenal ulcers in the two treatment groups although cigarette smoking was associated with lower rates of healing at four weeks both with the morning and nocturnal regimen.

PART 2

The 24 h pH profiles for placebo and ranitidine given at 8 am and for placebo and ranitidine given at 10 pm are reported in Figures 1 and 2.

It clearly emerges from Figure 1 that the ranitidine pH curve is markedly different from the placebo curve most of the daytime. Ranitidine in the morning takes effect at about 10 am, when pH concentrations rise above 5; high pH values are then maintained until 3 pm after which pH returns to the basal values. During the remaining part of the day and the whole of the night the ranitidine curve resembles the placebo curve. As emerges from Figure 2, nocturnal ranitidine takes effect around midnight and pH values are maintained above 5 throughout the whole nocturnal period returning to basal values at about 10 am. During the remaining part of the day the pH profile for nocturnal ranitidine is essentially similar to that for placebo. Comparison between the two figures clearly reveals that the duration of action of nocturnal ranitidine on intragastric pH is longer than that of the morning regimen.

The 2-by-2 comparison between the mean AUC values of the 24 h pH profiles with the two regimens (Table III) shows that a significant difference exists between both placebo and ranitidine 300 mg in the morning (t=−5.78, p<0.001) and between placebo and ranitidine 300 mg at night (t=−9.24, p<0.001).

![Figure 2: Median half hourly intragastric pH in nine patients with duodenal ulcer receiving placebo or ranitidine 300 mg at 10 pm.](image)

**TABLE III** Individual values and means±SD of AUC of the 24 hour median pH profiles calculated for placebo and ranitidine (R) 300 mg at 10 pm (left side) and placebo and ranitidine (R) 300 mg at 8 am (right side)

<table>
<thead>
<tr>
<th>Patient</th>
<th>Placebo</th>
<th>Placebo</th>
<th>Placebo</th>
<th>Placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PI</strong></td>
<td>1.945</td>
<td>5.223</td>
<td>2.674</td>
<td>3.877</td>
</tr>
<tr>
<td>2</td>
<td>4.613</td>
<td>5.686</td>
<td>1.778</td>
<td>3.521</td>
</tr>
<tr>
<td>3</td>
<td>1.853</td>
<td>5.420</td>
<td>2.906</td>
<td>5.545</td>
</tr>
<tr>
<td>4</td>
<td>2.907</td>
<td>5.194</td>
<td>2.534</td>
<td>3.649</td>
</tr>
<tr>
<td>5</td>
<td>2.343</td>
<td>4.589</td>
<td>2.017</td>
<td>3.651</td>
</tr>
<tr>
<td>6</td>
<td>1.821</td>
<td>5.232</td>
<td>2.653</td>
<td>5.654</td>
</tr>
<tr>
<td>7</td>
<td>3.578</td>
<td>5.977</td>
<td>2.551</td>
<td>4.335</td>
</tr>
<tr>
<td>8</td>
<td>2.284</td>
<td>5.361</td>
<td>2.108</td>
<td>4.179</td>
</tr>
<tr>
<td>9</td>
<td>1.048</td>
<td>4.424</td>
<td>1.842</td>
<td>3.284</td>
</tr>
<tr>
<td><strong>Mean</strong></td>
<td>2.254</td>
<td>3.011</td>
<td>2.349</td>
<td>4.210</td>
</tr>
<tr>
<td><strong>(SD)</strong></td>
<td>0.628</td>
<td>0.673</td>
<td>0.422</td>
<td>0.868</td>
</tr>
</tbody>
</table>

**Discussion**

First, Dragstedt postulated that nocturnal acid secretion was the most important pathophysiological factor in duodenal ulcer and considered this the basis for its vagotomy operations. Subsequently, the importance of nocturnal acidinity was confirmed by the high degree of efficacy of bedtime H2-blockers at low doses in preventing ulcer recurrence. In more recent years, several clinical studies have shown that the inhibition of nocturnal acid secretion alone, as obtained by large single bedtime doses of H2-blockers, induces duodenal ulcer healing rates comparable to those achieved with regimens inhibiting gastric acidity throughout the whole 24 h period. Successful healing of duodenal ulcers, however, has also been obtained using meal time doses of H2-blockers and divided doses of antacids, regimens designed mainly to inhibit daytime intragastric acidity. This indicates that the exact extent and duration of suppression of gastric acid secretion necessary to promote healing of duodenal ulcers is still uncertain.

In the first part of this study, we have shown that the antiulcer efficacy of ranitidine 300 mg in the morning does not significantly differ from that of ranitidine 300 mg at night. The healing rate observed with the morning regimen was only slightly inferior to the nocturnal healing rate at four weeks (67% vs 75%), but rates were practically identical at eight weeks (82% vs 85-5%). The 75% healing rate at four weeks in the group taking 300 mg at night was comparable with the 73% rate obtained in a previous anti-
ulcer trial with ranitidine nocte recently conducted in our unit; it is also on a par with the 78% rate observed by Lee et al in a large multicentre study comparing ranitidine 300 mg at night with 150 mg bid in a population of 424 patients. On the other hand, the good antitulc efficacy of the morning dose shown in our study agrees with the preliminary results of a recent multicentre antitulc study comparing single morning versus single nocturnal doses of famo-
tidine. The authors found no significant differ-
ences between the healing rates obtained with the two regimens either at four or eight weeks of treatment. Our results, however, should still be considered preliminary ones, because of a differ-
ence of importance (25%) in favour of the noct-
urnal regimen which cannot be excluded be-
cause of the relatively small size of the trial. Therefore, it would be worthwhile confirming these findings in a larger series and also checking for healing after two weeks of treatment to exclude a real difference between the two treat-
ments in terms of rapidity of healing. In this regard Merki et al have recently shown that the antitulc effects of ranitidine 300 mg given at 6 pm or at 10 pm significantly differ at two weeks (healing rates of 74% vs 50%, respectively), but are comparable at four weeks. In the second part of the present work, we have shown that both ranitidine regimens are significantly effec-
tive in suppressing 24 h intragastric acidity as compared with placebo; however, the duration of the antisecretory action of the morning dose is shorter than that of the nocturnal dose, thus confirming the results observed by others with the twice daily dosage. The reason for the latter event is not very clear, but it may be seen as an ‘escape phenomenon’ of the gastric inhibitory effect of ranitidine induced by the meal taken at 12:30 am. In support of this hypothesis, Johnston and Wormsley have recently reported that the nocturnal antisecretory effect of rani-
tidine at 6 pm is virtually abolished by a meal taken later in the evening. As a large morning dose of ranitidine has virtually no effect on nocturnal intragastric acidity, its high antitulc activity strongly suggests that reducing daytime acid secretion may be as per-
missive as the suppression of night time acidity for the healing of duodenal ulcers. The patho-
physiological significance of this observation is not yet known; a possible explanation is that duurnal acidity, particularly postprandial, plays a more important role in ulcerogenesis than was hitherto believed. Increased or prolonged acid secretory response to meals is, in fact, a well recognised physiological abnormality in a significant proportion of duodenal ulcer patients; thus, a single large morning dose of H2-blocker V should be considered, the secretory response to breakfast and lunch, would result in high duodenal ulcer healing rates. In this regard, other authors have recently devoted their atten-
tion to the effect of food on the duodenal ulcer healing process showing that food induced gastric secretion may have a major negative effect, at least in ulcers which fail to respond to H2-blocker therapy. In conclusion, our findings suggest that the duodenal ulcer healing process may take place during the day as well as the night, provided that a sufficiently long period of gastric acid inhibition is assured. As a practical consequence, this means that it may not matter whether anti-
secretory drugs are administered at bedtime or during the morning if their dose is capable of reducing acid secretion for a long enough period.

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