Low dose famotidine and cimetidine in single postprandial doses: a placebo controlled comparative study of overnight pH

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Abstract
To investigate the relative abilities of low doses of famotidine and cimetidine to raise intragastric pH after a single postprandial evening dose, 16 healthy volunteers were recruited to a four period crossover trial of famotidine 10 mg, cimetidine 100 mg and 200 mg compared with placebo. Intragastric pH was monitored between 1800 and 0730 with a nasogastric pH electrode. Median gastric pH rose from 1·35 (interquartile range 1·1-1·65) with placebo to 1·95 (1·6-5·35, p<0·001 Friedman rank) after dosing with famotidine 10 mg, to 1·46 (1·3-2·0, 0·05<p<0·1) after cimetidine 200 mg, and remained 1·35 (1·1-1·6, p>0·2) after cimetidine 100 mg. Intragastric pH was above 3 for 34% (p<0·005) of the time after dosing with famotidine, compared with 13·6% (p>0·2) after cimetidine 200 mg, 9·5% (p>0·2) after cimetidine 100 mg, and 4·7% after placebo. The rise of intragastric pH after famotidine 10 mg is significantly greater than that after either 200 mg or 100 mg cimetidine when the drugs are used postprandially.

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Low dose H$_2$ receptor antagonists have recently been marketed for 'over the counter' use in dyspepsia. They are recommended for postprandial and nocturnal dyspepsia and acidity. It is assumed that the symptomatic responses of dyspepsic patients will relate directly to the antisecretory effects of these agents and thus the pharmacodynamic activity of the drugs may be used as a surrogate measure of clinical efficacy. We therefore sought to compare the antisecretory properties of famotidine 10 mg (Pepcid AC, Centra Healthcare), cimetidine 100 mg and 200 mg (Tagamet, SKB) in a controlled study. The relative antisecretory effect of such doses has not previously been published.

Methods

Subjects
Sixteen healthy subjects (eight males, eight females) median age 21 years (range 19-25) and median weight 71 kg (range 53-80) were recruited into the study. The subjects included four smokers who were able to abstain from smoking on the days of the study. Median weekly ethanol consumption ascertained by questionnaire was 13 units (range 0-30). Each volunteer gave his or her written informed consent and the study protocol was approved by South Birmingham Health Authority Ethics Committee.

Study design
The investigation was designed as a four period crossover study with all subjects receiving the four different preparations in a random order. The study was partially blinded in that the drugs were recognisably different but the subjects were not aware of which was which. Of the two principal investigators one (RPW) was blinded to the drugs. There was a wash out period of at least six days between studies.

On each study day subjects arrived at 1700 on the investigation unit having fasted for six hours. Bipolar glass pH electrodes (Ingold M440) were calibrated in standard buffer solutions of pH 7·00 and 4·01 and calibration was verified at pH 1·69. These electrodes were passed by the nasogastric route. A drop in pH to less than 2 was taken as evidence of the tip of the probe having entered the stomach, and the electrode was advanced a further 8 cm from this point. At 1830 a standard meal was given, which consisted of a supermarket ready meal of cottage pie, peas, and carrots, followed by chocolate coated ice cream and two chocolate mints to give a total of 700 kcal, provided by 22 g protein, 70 g carbohydrate, and 37 g fat, accompanied by 250 ml mineral water.

The drugs were given as tablets (famotidine 10 mg and cimetidine 200 mg) or as elixir (cimetidine 100 mg) with 50 ml mineral water at 1930. The placebo was a tablet containing no active ingredient. Subjects had nothing further to eat before the end of the study period, but had a further 250 ml of mineral water at 2100. The subjects retired to bed at 2300 and the pH electrodes were removed at 0730 after an overnight stay on the investigation unit under the supervision of one of the investigators (TGR).

Analysis of data
pH measurements were recorded using GastrograpH recorders (Medical Instruments Corporation, Solothurn, Switzerland). These instruments measure the potential difference between the electrode and the reference electrode in the tip of the tube four times per
second. Median pH values are stored giving data on pH 10 times per minute. The data were transferred to a computer for graphic and numerical analysis. Half hourly medians for each study period were extracted from the raw data and all subsequent data summaries and analyses were based on these half hourly medians. Friedman's rank test based on the $\chi^2$ distribution was used to compare different treatments.

The principal study periods analysed were as follows: the period from the drug being given to the end of the study (drug to end); and from the drug to seven hours (drug plus 7). Secondary descriptive analyses were performed in an attempt to define onset of action using the periods from drug administration to one hour, $1\frac{1}{2}$ hours, and two hours after dosing.

**Results**

Sixteen volunteers participated in all parts of the study. On two occasions in two different volunteers an error in programming the recording device necessitated a study being repeated. On both occasions the finishing time was incorrectly entered so that on the first occasion the recording ceased at midnight, and on the second it ceased at 0630. On both occasions the repeat study was the one used for analysis. All studies were well tolerated and no significant adverse events were reported during or after any study.

Figure 1 shows the group median half hourly pH over the whole study period for all studies and shows that pH rose substantially after famotidine and less after cimetidine 200 mg. The pH/time curve after cimetidine 100 mg was indistinguishable from that with placebo. The period when the difference between the $H_2$ receptor antagonists was most easily visualised was from seven hours after the drug administration.

**Primary analyses**

**Drug to end** – Figures 2–4 show the individual median pH values from drug administration to the end of the study for all dosings.
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Figure 4: Median pH placebo comparisons for 16 subjects after famotidine 10 mg: drug to end period.

for the different drugs. After placebo the group median pH (interquartile range) was 1.35 (1.1–1.65); after cimetidine 100 mg it was 1.35 (1.1–1.6; p>0.2 compared with placebo); after cimetidine 200 mg it was 1.46 (1.3–2.0; 0.05<p<0.1); and after famotidine 10 mg it was 1.95 (1.6–5.35; p<0.001). We examined the proportion of time that gastric pH was above 3 and 4 for each drug (Table). The proportion of time above both pH values was statistically significantly greater with famotidine 10 mg than placebo; differences with both doses of cimetidine did not reach statistical significance.

Drug to seven hours – As the half life of cimetidine is comparatively short (two hours) any effects of the drug may have worn off well before the end of the study. We therefore analysed the data from dosing up to seven hours. Compared with placebo, median pH was raised by famotidine 10 mg from 1.25 to 1.8 (p<0.005); after cimetidine 200 mg to 1.39 (0.05<p<0.1, not significant), and there was no significant difference between placebo and cimetidine 100 mg (1.25 v 1.25, p>0.2). The area under the pH/time curve from drug to seven hours after dose showed a significant difference from placebo for both famotidine 10 mg (p<0.0001) and cimetidine 200 mg (p>0.01).

Descriptive analyses
To assess the onset of significant activity we compared the effects of cimetidine 200 mg and famotidine 10 mg with placebo at earlier times after drug administration. Famotidine 10 mg showed a significant difference at two hours (p<0.02) but not before. After cimetidine 200 mg we could not show a significant difference from placebo at any time when the median pH was used for analysis.

Discussion
Low dose preparations of H2 receptor antagonists are of interest at present because they have recently become available to the public without prescription as over the counter drugs. This study sought to show whether there were differences in the pharmacodynamic activities of famotidine 10 mg, cimetidine 100 mg, and cimetidine 200 mg. In planning this investigation we designed a protocol that would closely mimic the use of these drugs for self treatment. We believe that postprandial heartburn is the symptom for which these drugs are likely to be used and therefore arranged to study the effects of the drugs taken soon after a realistically typical evening meal rather than at bedtime or in the fasted state. Healthy volunteers rather than patients were used for practical reasons and because in many cases self treatment would be by subjects otherwise recognised as normal. We did not include any snacks after the main meal as eating after the evening dose of an H2 receptor antagonist has been shown greatly to reduce the effectiveness of the drug.1–3 We timed the dose at one hour after the meal in a further attempt to mimic the symptomatic use of the drug postprandially. It is recognised that an early evening dose with potent H2 antagonists produces more acid inhibition than bedtime administration.4

Famotidine 10 mg raised pH significantly versus placebo over the whole postdose period while the less potent drug (cimetidine) did not achieve this. After dosing with famotidine 10 mg a significant difference in median pH compared with placebo was found at two hours but after cimetidine 200 mg we could not show a significant difference. Reference to the curve of median pH against time (Fig 1), however, shows that cimetidine 200 mg clearly does have an effect on acidity, and by using the area under the pH/time curve it is possible to show a significant difference from placebo for this drug over the seven hour period (p<0.01). There is debate as to which measure of central tendency should be used in studies such as this: we have used the median because, unlike the mean or the area under the curve, it...
permits direct comparison of pH and hydrogen ion concentration. As far as cimetidine 200 mg is concerned it is reasonable to suppose that a statistically significant difference shown by one method but not by another is unlikely to represent a difference of any clinical significance.

Cimetidine 100 mg had little effect on postprandial intragastric pH. A possible criticism of our method is that the liquid form used in the study has a different bioavailability from the tablet form available over the counter. While no published data exist for the 100 mg dose of cimetidine, a comparative study found that a 300 mg tablet of cimetidine was bioequivalent to a 300 mg dose of oral liquid, and it may be supposed that this relation holds good for the lower dose. These results suggest that in the doses tested famotidine acts more than cimetidine.

The greatest rise in pH after famotidine 10 mg was seen after seven hours and fits with previous data on the effects of H2 antagonists overnight. During this period, competition at H2 receptors between drug and histamine is likely to be minimal because by this time food stimulated acid secretion will have worn off and the diurnal pattern of acid secretion will have passed its peak. But substantial plasma drug concentrations will remain. It is also the period when in healthy stomachs spontaneous late night gastric alkalisation is encountered.

The surprising finding is the difference between the effect of cimetidine 200 mg and famotidine 10 mg. Famotidine is about 20 times as potent as cimetidine on a weight to weight basis, and it would therefore have been expected that the doses tested would be equipotent. In a previous study inhibition of meal stimulated acid secretion was seen within 90 minutes, and the range of mean pH values obtained was 2.0-2.3 between 1.7 and 3.2 hours postdose after oral famotidine 10 mg. These results are in accord with our findings, which additionally show for how long this drug is able to raise intragastric pH. Our study was not primarily designed to determine the onset of action, which would require further study and it is probable that the drugs were actively inhibiting acid secretion well before any detectable change in acidity. During the first two hours after a meal food in the stomach tends to buffer acid secretion hence obscuring drug action, so it is possible that both drugs were active earlier than our results would suggest.

Our results differ, however, from the only other study of low dose cimetidine in healthy volunteers. That earlier study showed that low doses of cimetidine (100 mg and 200 mg) taken at 2300 reduced nocturnal acid secretion over an eight hour period in nine healthy fasted subjects, achieving for both preparations intragastric pH values of above pH 3 for 100% of the time. Methodological differences can probably explain the apparent variance of results.

This study differed substantially from our own: the volunteers were fasted, the investigation only lasted eight hours and started late at night when the stomach should be resting, and continuous aspiration of gastric contents gave measurements of basal acid secretion rather than acidity. Other studies that have looked at low dose cimetidine have found less impressive inhibition of acid secretion, but have used duodenal ulcer patients and are hence not strictly comparable. One subject showed no response to any of the drugs and it is likely that he was a non-responder to H2 receptor antagonists. This finding re-emphasises the known wide variation of response within populations. Such variation in dose requirement is most noticeable with low doses and could influence symptomatic efficacy if the proportion of non-responders is great. It is expected that these over the counter drugs will be used in response to symptoms and it is likely that their effect will mirror their antisecretory activity. We conclude that famotidine 10 mg is more effective than either cimetidine 100 mg or 200 mg at raising intragastric pH with a single dose taken after a normal evening meal.

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