Cimetidine pharmacodynamics and pharmacokinetics in healthy subjects: A comparison of tablets and suspension

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The objective of this study was to compare the effect of cimetidine 200 mg tablet with that of cimetidine suspension (200 mg/10 mL), a 20 mL suspension of 800 mg magnesium hydroxide and 912 mg aluminum hydroxide, and matching placebo on intragastric pH of healthy volunteers. There were 13 males and seven females, mean age 23 years (range 20 to 32) and mean weight 72 kg (range 55 to 89). The intragastric pH of each subject was measured over 6 h starting immediately before dosing on each of five study days. Cimetidine plasma levels were measured for 6 h after dosing on each of the cimetidine study days. Cimetidine tablet and suspension were superior to placebo tablet or suspension or to the magnesium hydroxide/aluminum hydroxide suspension in the area under the pH time curve from 0 to 6 h, percentage time pH of at least 3.5, change from pretreatment pH area under the pH time curve (0 to 6 h) and maximum increase in pH. Mean plasma cimetidine levels were significantly and positively correlated to mean intragastric pH for both cimetidine tablets and cimetidine suspension. Comparing cimetidine concentration (Cmax) and lower percentage time plasma cimetidine concentration was at least 0.5 µg/mL (the minimum therapeutic level). These pharmacokinetic variations between cimetidine suspension and tablets may have partially explained the pharmacodynamic differences of the lower area under the pH time curve (0 to 6 h). The results indicate that both...
cimetidine tablets and cimetidine suspension significantly increase intragastric pH relative to the magnesium hydroxide/aluminum hydroxide suspension, placebo tablets or placebo suspension. Based on the assumption that elevation of intragastric pH is an important factor for alleviation of the symptoms and for the healing of peptic disorders, the results of this study suggest that both cimetidine tablets and cimetidine suspension should be effective treatment.

Key Words: Antacid, Cimetidine suspension, Cimetidine tablets, Intragastric pH

Pharmacodynamie et pharmacocinetique de la cimetidine chez des sujets sains : comparaison entre comprimes et suspension

RÉSUMÉ : L'objectif de cette étude était de comparer l'effet de la cimetidine en comprimés de 200 mg, de la cimetidine en suspension (200 mg/10 mL), et d'une suspension de 20 mL de 800 mg d'hydroxyde de magnésium, de 912 mg d'hydroxyde d'aluminium, et de placebo correspondant, sur le pH intragastrique de volontaires sains. Le groupe comprenait 13 hommes et sept femmes dont la moyenne d'âge était de 23 ans (entre 20 et 32 ans) et le poids moyen de 72 kg (entre 55 et 89 kg). Le pH intragastrique de chaque sujet a été mesuré sur une période de six heures avant l'administration de la dose pour chacun des cinq jours de l'étude. Les taux plasmatiques de cimetidine ont été mesurés six heures après administration, chaque jour de cette étude sur la cimetidine. Le comprimé et la suspension de cimetidine se sont révélés supérieurs au comprimé ou à la suspension placebo, et à la suspension d'hydroxyde de magnésium/hydroxyde d'aluminium pour l'aire sous la courbe pH-temps de 0 à 6 heures, pour le pourcentage temps-pH d'au moins 3,5, pour tout changement par rapport à l'aire sous la courbe pH-temps comparé au pH en prétraitement (0 à 6 heures) et pour l'augmentation maximum du pH. Les taux plasmatiques moyens de cimetidine ont été en corrélation positive et significative avec le pH intragastrique moyen pour les comprimés de cimetidine et la suspension de cimetidine. La différence entre la concentration de cimetidine (Cmax) et le pourcentage minimum temps-concentration de cimetidine plasmatische a été d'au moins 0,5 g/mL (taux thérapeutique minimum). Ces variations pharmacocinétiques entre la suspension et les comprimés de cimetidine peuvent avoir expliqué partiellement les différences pharmacodynamiques de l'aire minimale sous la courbe pH-temps (0 à 6 heures). Les résultats indiquent que les comprimés et la suspension de cimetidine augmentent significativement le pH intragastrique comparativement aux suspensions d'hydroxyde de magnésium/hydroxyde d'aluminium, aux comprimés de placebo et à la suspension de placebo. Sur la base de l'hypothèse selon laquelle l'élévation du pH intragastrique est un facteur important pour le soulagement des symptômes et pour la cicatrisation des lésions duodénales, les résultats de cette étude suggèrent que les comprimés de cimetidine et la suspension de cimetidine constituent un traitement efficace.

SUBJECTS AND METHODS

The study was a placebo-controlled crossover trial design in 20 normal healthy volunteers. The intragastric pH of each subject was measured over 6 h starting immediately before dosing on each of the five study days. The cimetidine plasma level was measured for 6 h after dosing on each of the cimetidine study days.

Ethics: The study was conducted according to the provisions of the Declaration of Helsinki as amended in Venice in 1983. Ethical Review Committee approval was obtained. The nature of the study was explained to the subjects, and written informed consent was obtained from all subjects before entering the study. The subjects were also informed of their right to withdraw from the study at any time.

Subjects: Males and females between 18 and 50 years of age were eligible for inclusion. Pregnant or lactating women were ineligible for the study. Women of childbearing potential were included, but were cautioned against becoming pregnant during the study. Subjects receiving any medication other than oral contraceptives for two weeks before the study or who received any investigational drug within four weeks of the start of the study, were ineligible, as were subjects with a significant abnormality as determined by medical history and physical examination or with significant pretrial laboratory abnormalities. Any subject with a history of significant gastrointestinal surgery or any other surgery that might interfere with the absorption, metabolism or excretion of cimetidine was also ineligible. Any subject who was known to have been noncompliant or whom the investigators felt was unreliable or unsuitable to participate in a clinical trial was also ineligible.

Twenty subjects were evaluated for inclusion into the study. All subjects completed the study according to the protocol. There were 13 males and seven females; mean age was 23 years (range 20 to 32) and mean weight was 72 kg (range 55 to 89).

Materials: Cimetidine was supplied as a 200 mg tablet and 200 mg/10 mL suspension with matching placebo tablets and suspension. A 20 mL suspension of 800 mg magnesium hydroxide and 912 mg aluminum hydroxide was also used. Study medication was taken orally according to a randomization schedule. The dose was 200 mg cimetidine tablet, 200 mg/10 mL cimetidine suspension, 20 mL magnesium hydroxide/aluminum hydroxide suspension or matching placebo suspension or tablet.

Schedule of studies: Subjects began fasting at 22:00 the night before each study day. Water was taken ad libitum until the start of the gastric secretion study. No food was permitted during the 6 h study period. No fluids were permitted for up to 4 h after administration of the study medication. Thereafter, water was allowed, but was limited to a maximum of 200 mL/h. The volume and time of water consumption was recorded on the case report form.

Each subject was studied on five separate study days, separated by a washout period of at least five days. On each study day the subject reported to the Clinical Investigation Unit of the Walter C Mackenzie Health Science Centre (Edmonton, Alberta) at 07:00. The subject was weighed and exam-
TABLE 1
Intragastric pH from 0 to 6 h following treatment with cimetidine suspension, cimetidine tablet, placebo suspension, placebo tablet or magnesium hydroxide/aluminum hydroxide suspension

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Cimetidine tablet</th>
<th>Cimetidine suspension</th>
<th>Placebo tablet</th>
<th>Placebo suspension</th>
<th>Magnesium hydroxide/aluminum hydroxide suspension</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH AUC (0 to 6 h)</td>
<td>28.52±1.57</td>
<td>22.50±1.57</td>
<td>17.24±1.57</td>
<td>15.50±1.57</td>
<td>16.33±1.57</td>
</tr>
<tr>
<td>% time pH ≥ 3.5 (h)</td>
<td>65.28±5.22</td>
<td>45.7±5.22</td>
<td>27.15±5.22</td>
<td>22.80±5.22</td>
<td>22.36±5.22</td>
</tr>
<tr>
<td>Peak pH</td>
<td>7.03±0.32</td>
<td>6.02±0.32</td>
<td>5.31±0.32</td>
<td>4.63±0.32</td>
<td>5.39±0.32</td>
</tr>
<tr>
<td>Time to peak pH (h)</td>
<td>2.36±0.32</td>
<td>2.58±0.32</td>
<td>1.63±0.32</td>
<td>2.11±0.32</td>
<td>0.77±0.32</td>
</tr>
<tr>
<td>Change from pretreatment AUC (0 to 6 h)</td>
<td>13.75±1.86</td>
<td>8.21±1.86</td>
<td>-0.09±1.86</td>
<td>-2.07±1.86</td>
<td>-0.45±1.86</td>
</tr>
<tr>
<td>Maximum increase in pH</td>
<td>4.75±0.37</td>
<td>3.64±0.37</td>
<td>2.42±0.37</td>
<td>1.70±0.37</td>
<td>2.59±0.37</td>
</tr>
</tbody>
</table>

Mean ± SEM. AUC Area under the pH time curve. *Significant (P<0.05) versus cimetidine tablet; †Significant (P<0.05) versus cimetidine suspension

A nasogastric tube was positioned in the stomach using the standard aspiration technique (5). Subjects remained ambulant during the study day. Gastric acidity was monitored by a method similar to that described by Pounder et al (5). At 08:00 the study medication was administered. Sampling began immediately before administration of study medication. Thereafter, a 5 mL sample of gastric juice was aspirated at 10, 20, 30, 45 and 60 mins, and then at 30 min intervals for a total of 6 h. The pH of the sample was measured to the nearest 0.1 unit using a combined glass reference pH electrode and meter. The sample was then returned to the subject's stomach. The pH electrode was calibrated with standard buffers of pH 2.0, 4.0 and 7.0. The calibration was checked immediately before each set of measurements. The time and result of each pH measurement was recorded. Immediately before the administration of tablet or suspension, a minimum 3 mL venous sample was drawn to establish a plasma baseline. Thereafter at 10, 20, 30, 45 and 60 mins, and then every 30 mins up to 6 h, blood was drawn to measure plasma cimetidine levels. The sample was drawn through a heparin lock or by venepuncture depending upon the subject's preference. The sample was centrifuged and the plasma was separated and stored frozen for later measurement of plasma cimetidine concentrations. The serum cimetidine concentration was measured by the high performance liquid chromatographic method of Sollin et al (6).

Safety monitoring: All adverse events that occurred during the study were recorded, together with their severity and relationship to study medication. Adverse events attributed to study medication, as well as those in which the relationship to the study medication was unknown or unspecified, were separately tabulated. The severity outcome, and dates of onset and clearance for each event were recorded. No adverse effects were recorded during the study. Routine blood tests for hematology, biochemistry and urinalysis were taken during screening and at follow-up. A pregnancy test was given during screening. Routine hematology included hemoglobin, white blood cell count and platelet count. Routine serum chemistries included measurements of aspartate aminotransferase, alanine aminotransferase, total bilirubin, creatinine, sodium, potassium, chloride and bicarbonate concentrations.

Data analysis: On each study day the subjects' individual intragastric pH values and the adjusted mean pH over 6 h were plotted. The following parameters were determined: area under the pH time curve from 0 to 6 h, peak pH (Cmax), time to peak pH (Tmax), percentage of time pH was at least 3.5, area under the change from pretreatment pH time curve and maximum increase in pH from pretreatment value. The individual plasma cimetidine level and adjusted mean values were plotted over the same period as the intragastric pH. The following parame-
TABLE 2
Cimetidine plasma levels from 0 to 6 h following treatment with cimetidine suspension or cimetidine tablet (adjusted mean)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Cimetidine tablet</th>
<th>Cimetidine suspension</th>
</tr>
</thead>
<tbody>
<tr>
<td>Area under the pH time curve (0 to 6 h)</td>
<td>3.33±0.10</td>
<td>2.58±0.10</td>
</tr>
<tr>
<td>Peak pH</td>
<td>1.04±0.05</td>
<td>0.81±0.05</td>
</tr>
<tr>
<td>Time to peak pH</td>
<td>1.75±0.26</td>
<td>1.78±0.26</td>
</tr>
<tr>
<td>Time to therapeutic level ≥0.5 µg/mL</td>
<td>56.94±3.42</td>
<td>41.24±3.42</td>
</tr>
</tbody>
</table>

Mean ± SEM from 20 subjects. *P<0.01

Figure 2) Mean plasma cimetidine levels. O—Cimetidine tablet; ●—Cimetidine suspension

Figure 3) Correlation of pH and plasma cimetidine concentrations for cimetidine tablets (O). ● Cimetidine suspension

RESULTS
A pH reading was missing at 2.5 h for subject 6 and a plasma cimetidine was missing at 2.5 h for subject 9. Linear interpolation was used to estimate the two missing values before summarizing the parameters. All regimens had subjects with some pretreatment intragastric pH values greater than 3.0; the group receiving cimetidine suspension had four, the placebo tablet and suspension groups have five each, the cimetidine tablet group had six and the magnesium hydroxide/aluminum hydroxide suspension group had seven. Analysis of the pretreatment pH levels showed no statistically significant differences among the groups.

The cimetidine tablet has a larger (P<0.05) area under the pH time curve and change in pH curves than the cimetidine suspension, the magnesium hydroxide/aluminum hydroxide suspension, placebo tablet or placebo suspension (Table 1). A representative result from one subject is shown in Figure 1. Cimetidine suspension has a greater (P<0.05) area under the pH time curve and change in pH curves than the magnesium hydroxide/aluminum hydroxide suspension, placebo tablet or placebo suspension. The percentage time the pH was at least 3.5 was significantly greater for the cimetidine tablet and suspension than...
for the magnesium hydroxide/aluminum hydroxide suspension, placebo tablet or placebo suspension. The cimetidine tablet resulted in a longer time at or above pH 3.5 than cimetidine suspension. The cimetidine tablet has a higher peak pH than the cimetidine suspension, magnesium hydroxide/ aluminum hydroxide suspension, placebo tablet or placebo suspension. The time to peak pH was not significantly different for the cimetidine tablet, cimetidine suspension, placebo tablet or placebo suspension. The time to peak pH was longer (P<0.05) for the cimetidine tablet than for the cimetidine suspension, magnesium hydroxide/ aluminum hydroxide suspension, placebo tablet or placebo suspension. The maximum increase in pH was significantly greater for the cimetidine tablet than for the magnesium hydroxide/aluminum hydroxide suspension, placebo tablet or placebo suspension, but not between the cimetidine tablet and suspension.

The area under the cimetidine plasma concentration curve was larger (P<0.01) for the tablet than for the suspension (Table 2, Figure 2). The peak cimetidine plasma level (Cmax) was significantly greater for the tablet than for the suspension. There were no significant differences between the tablet and suspension for time to peak cimetidine plasma level (Tmax) or time to reach a cimetidine plasma level of 0.5 µg/mL (the minimum therapeutic level) or greater. The percentage time that the cimetidine plasma level was at least 0.5 µg/mL was longer for the tablet than for the suspension (P<0.01).

The mean 0 to 6 h intragastric pH for both cimetidine tablets and suspension was significantly correlated to plasma cimetidine levels (Figure 3) (Pearson correlation coefficient 0.888, P<0.001, with a y axis intercept of 2.38±0.29 and a slope of 3.82±0.36).

**DISCUSSION**

Antacids such as the magnesium hydroxide/aluminum hydroxide suspension are commonly used to relieve symptoms of heartburn and indigestion. Cimetidine has been proven to be effective in the treatment of patients with peptic ulcer disease and gastroesophageal reflux disease (7,8). Cimetidine has also been shown to ameliorate symptoms of these conditions. It is thought that the relief of such symptoms by cimetidine and antacids is related to the ability of these drugs to decrease gastric acidity. The purpose of this study was to compare the effect on intragastric pH of cimetidine tablets to cimetidine suspension, magnesium hydroxide/aluminum hydroxide suspension, and placebo tablet and placebo suspension. Pharmacodynamically, taking into account all the parameters measured, the cimetidine tablet was superior to all other treatment arms; cimetidine suspension was superior to the magnesium hydroxide/aluminum hydroxide suspension or placebo only in terms of maximum increase in pH (Table 1). Also, cimetidine tablets were superior to cimetidine suspension for the area under the pH time curve, peak pH, time pH was greater than 3.5 and the area under the change from pretreatment pH curve (Table 1, Figure 1).

Pharmacokinetically, the area under the pH time curve, Cmax and percentage time above the minimum therapeutic cimetidine level (0.5 µg/mL) were higher (P<0.01) following cimetidine tablets than cimetidine suspension (Table 2, Figure 2). This suggests that the cimetidine tablets were superior to the cimetidine suspension because the tablet maintained the therapeutic cimetidine level for a longer period than the cimetidine suspension, possibly as the result of different rates of dissolution or absorption. There was a significant correlation between mean pH and mean plasma cimetidine levels following both cimetidine regimens (Figure 3), including that the same pH effect was achieved from a given plasma cimetidine concentration achieved from the tablets and suspension. Whereas all of the cimetidine suspension-treated subjects showed plasma cimetidine concentrations less than 0.6 µg/mL, only half of the individuals treated with cimetidine tablets had values less than 0.6 µg/mL and the other half had higher values (Figure 3), suggesting that cimetidine tablets produced higher levels of plasma cimetidine and maintained higher levels of cimetidine in the plasma for a longer period of time. It is unclear what was the basis of the differences in the pharmacokinetics of the cimetidine suspension versus the tablets, but these variations in pharmacokinetics explained the differences in pharmacodynamics, ie, the pH measurements.

Although there were no significant differences between the baseline mean pH of the groups, all arms of the study had individuals with a baseline pH greater than 3. They were not necessarily the same individual in each arm of the study. The possibility that these high pretreatment pH subjects may not have been able to show their true response to the treatment or that they were false responders (elevated pH not due to treatment) cannot be discounted.

These results indicate that both cimetidine tablets and suspension are superior to the magnesium hydroxide/aluminum hydroxide suspension, placebo tablets or placebo suspension in increasing intragastric pH. Based on the assumption that elevation of intragastric pH is an important factor for alleviation of the symptoms of peptic disorders, the results of this study suggest that both cimetidine tablets and cimetidine suspension should be effective treatment. Although cimetidine tablets were statistically superior to cimetidine suspension for most pharmacodynamic and pharmacokinetic parameters, the magnitude of these differences was small, and the clinical relevance of the differences between the two formulations cannot be predicted from these data.

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**REFERENCES**


