Effect of inhibition of *Helicobacter pylori* urease activity by acethydroxamic acid on serum gastrin in duodenal ulcer subjects

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Abstract

The mechanism of the hypergastrinaemia associated with *Helicobacter pylori* infection is unknown. It may be an effect of the ammonia produced by the bacterium near the antral epithelial surface. We have examined the effect on serum gastrin of inhibiting *H pylori* urease activity with acethydroxamic acid in six duodenal ulcer patients. On day 1 the fasted patients received placebo tablets at 8 am, a peptide meal at 10 am, and a 14C urea breath test at 11.30 am. The next day 750 mg acethydroxamic acid was administered orally in place of the placebo. The median (range) 30 minute breath test value (dose/mmol CO2×kg body wt×100) was 152 (111–335) on day 1, but only 24 (14–49) the next day (p<0.03). Further studies performed in one subject confirmed that acethydroxamic acid lowered the ammonium concentration and raised the urea concentration in gastric juice. The inhibition of urease activity and ammonia production did not result in a fall in the basal gastrin concentration or in the median integrated gastrin response to the peptide meal, which was 78 ng/l.h (range 21–222) on day 1 and 79ng/l.h (33–207) the next day. Ten days after acethydroxamic acid, the urea breath test values were similar to those before treatment. This study shows that the raised gastrin concentration in patients with *H pylori* infection is not directly related to the organism’s urease activity. It also shows that temporary suppression of *H pylori* urease activity does not clear the infection.

Eradication of *Helicobacter pylori* infection of the gastric antrum results in a lowering of the circulating gastrin concentration.1,2 The fasting concentration falls by 27–33% and the integrated gastrin response to a meal by 30–58%. This stimulation of gastrin release by *H pylori* may be relevant to the organism’s role in duodenal ulcer disease. The mechanism by which chronic infection of the antral mucosa with *H pylori* results in increased gastrin release is not known.

*H pylori* is remarkable because of its high urease activity by which it hydrolyses urea to ammonia and carbon dioxide.3 As a result of this, patients with the infection have reduced concentrations of urea and increased concentrations of ammonium in their gastric juice.4 The production of ammonia by the bacterium at the antral epithelial surface could increase gastrin release by any of three theoretical mechanisms. Ammonia is a strong alkali and could therefore prevent the physiological inhibition of gastrin release by gastric acid.5 In addition, an increase in the antral surface pH by ammonia would facilitate the entry of dietary amines into the antral G cells and thereby their stimulation of gastrin release.6 Thirdly, ammonia could stimulate gastrin release directly, as has been shown to occur in the rat.7

In an attempt to elucidate the mechanism of the hypergastrinaemia associated with *H pylori* infection, we have examined the effect of inhibiting the bacterium’s urease activity and ammonia production on serum gastrin in duodenal ulcer patients.

Patients and methods

STUDIES IN PATIENTS WITH *H pylori* INFECTION

Six patients confirmed endoscopically to have duodenal ulceration within the previous year but currently in clinical remission were studied. Their median age was 39 years (range 26–52) and three were women. In each patient, an antral biopsy specimen obtained endoscopically within the preceding three months had shown gastritis associated with *H pylori* like organisms. The patients reported fasted and a venous blood sample was removed at 8 am for gastrin determination. Immediately after this they drank 50 ml water and further blood samples were taken at 30 minute intervals for two hours. At 10 am they took a standard meal consisting of two beef cubes (OXO Ltd, Croydon, England) dissolved in 200 ml water at 50°C. Further blood samples were taken at 10 minute intervals for 70 minutes and a final one at 90 minutes after the OXO drink. Immediately after this sample a 14C urea breath test was performed to measure *H pylori* urease activity. For this they drank 250 ml Ensure Plus (Abbott Laboratories, England) to delay gastric emptying, followed by 0.4 MBq 14C urea (Amersham International) in 25 ml water. Breath samples for 14C-CO2 analysis were obtained at 10 minute intervals for 90 minutes.

The next day, the study was repeated in an identical fashion except that the patients received 750 mg acethydroxamic acid (Lithostat, Mission Pharmacol, USA) with a 50 ml drink of water at 8 am.

Ten days later a third 14C urea breath test was performed to determine whether the temporary inhibition of urease activity had resulted in clearance of the infection.

STUDIES IN PATIENTS WITHOUT *H pylori* INFECTION

Two male patients (aged 25 and 52 years) with a
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Figure 1: Effect of acetohydroxamic acid on Helicobacter pylori urease activity assessed by the 14C urea breath test. The values are medians of six patients.

past history of duodenal ulcer disease, but in whom H pylori had been eradicated within the previous year, were studied in an identical fashion to that described above. This was performed in order to exclude the possibility that acetohydroxamic acid might have a direct effect on gastrin release.

STUDY IN A HEALTHY VOLUNTEER WITH H PYLORI INFECTION
The effect of acetohydroxamic acid on the concentrations of urea and ammonium in gastric juice was examined in a single healthy volunteer (AN, aged 32 years) with H pylori infection. Endoscopic antral biopsy specimens obtained three months earlier had shown gastritis and H pylori like organisms. The 14C urea breath test performed one month before the study gave a 30 minute value of 109% dose/mmol CO2 x kg body wt x 100 which is in the middle of our range for infected subjects (35–225).

He reported fasted at 8 am and a nasogastric tube was passed perorally. The resting gastric juice was aspirated and discarded and then constant suction applied. At 30 minutes the suction was temporarily stopped and a 5 ml sample of gastric juice was obtained by manual aspiration. A further such sample was obtained at 60 minutes. Immediately after this 750 mg acetohydroxamic acid was taken orally with 50 ml water and no suction was applied until one hour later when all the resting volume was aspirated and a 5 ml aliquot retained for analyses. Suction was then recommenced but temporarily stopped every 30 minutes over the following three and a half hours to allow manual aspiration of 5 ml samples of gastric juice.

ANALYSES
Serum gastrin concentrations were determined by a standard radioimmunoassay kit (CIS UK Ltd). Each patient’s day 1 and day 2 samples were assayed in the same batch. The concentration of urea in gastric juice was determined by a prospective analyser (American Monitor, West Sussex, UK) and the concentration of ammonium by an enzymatic method (Sigma, Dorset, UK). Preliminary studies confirmed that acetohydroxamic acid did not interfere with the analysis of urea or ammonium in gastric juice. The integrated gastrin response to the OXO meal was assessed by calculating the area under the serum gastrin concentration time curve using the trapezoid method.

Statistical analysis was performed by means of the Wilcoxon signed rank sum test. The study was approved by the Western Infirmary Ethical Committee and each patient gave written informed consent.

Results
The excretion of 14C-CO2 in the breath was considerably lower after the administration of acetohydroxamic acid (Fig 1). On the first study day, the median 30 minute breath test value was 152% dose/mmol CO2 x kg body wt x 100 (range 111–335) compared with a value of 22 (range 14–95) after acetohydroxamic acid (p<0.03). The median value 10 days after the administration of acetohydroxamic acid was 149 (range 126–257) which was similar to the pretreatment value (Fig 2).

In spite of the noticeable suppression of H pylori urease activity, there was no difference between the two study days with respect to the basal or meal stimulated serum gastrin concentrations (Fig 3). The median integrated gastrin response to the OXO meal was 78 ng/l h
of juice, collected one hour after the administration of the urease inhibitor, and persisted for the full three and a half hours studied. The ammonium concentrations in gastric juice before the inhibitor were 5.2 and 5.5 mmol/l and values after the inhibitor ranged from 1.9-3.6 mmol/l. The ratio of urea:ammonium in gastric juice before the urease inhibitor was 0.2 whereas after the inhibitor five of the six patients showed values more than 1.0.

None of the patients experienced any side effects after the administration of acetohydroxamic acid.

**Discussion**

Acetohydroxamic acid is a specific urease inhibitor that has been used to reduce bacterial ammonia production in patients with renal calculi secondary to chronic urinary tract infection. Reducing the ammonia production lowers the pH of the urine and thereby lessens the tendency for calculi formation. In these patients the drug has been prescribed in a dose of 250 mg four times a day. We administered the drug as a once off dose of 750 mg in order to achieve rapid and effective inhibition of urease activity.

Inhibition of *H. pylori* urease activity was shown by the fivefold reduction in the 30 minute values of the $^{14}$C urea breath test. The concentrations of urea and ammonium in the gastric juice of the volunteer also confirmed inhibition of urease activity. We have previously noted that though there is overlap of *H. pylori* negative and positive subjects with respect to their gastric juice ammonium concentration, the urea:ammonium ratio in gastric juice provides clear separation of the two groups. The median urea:ammonium ratio before the urease inhibitor was 0.2, which is within our range for infected subjects (0.04-0.7), and that after administration of the inhibitor was 1.2 which is just within our range for patients in whom the infection has been eradicated (1.1-1.13). The finding of altered urea and ammonium concentrations in the first sample of juice examined and its persistence throughout the remainder of the three and a half hours observed is consistent with the fact that acetohydroxamic acid is rapidly absorbed from the upper gastrointestinal tract reaching peak plasma concentrations at 60 minutes and has a plasma half life of three and a half to five hours. It is not clear whether the inhibition of *H. pylori* urease activity by oral acetohydroxamic acid is the result of a topical or systemic effect, or both.

In spite of the suppression of *H. pylori* urease activity there was no accompanying fall in the basal gastrin concentration or in the gastrin response to the OXO meal. There was also no change in gastrin concentration in the *H. pylori* negative subjects, which excluded the possibility that a fall in gastrin in those with the infection had been masked by a direct gastrin stimulating effect of the drug.

It has been proposed that the hypergastrinæmia in patients with *H. pylori* infection is caused by the ammonia produced by the bacteria raising the antral surface pH. This would prevent the physiological suppression of gastrin.

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**Figure 3: Basal and meal stimulated serum gastrin concentrations on placebo and after 750 mg acetohydroxamic acid. The values are medians of six patients.**

**Figure 4: Subjects integrated gastrin response to the OXO meal on placebo and acetohydroxamic acid.**
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Figure 5: Effect of 750mg acetohydroxamic acid on gastric juice concentrations of ammonium and urea and their ratio in healthy volunteers with Helicobacter pylori infection.

Acetohydroxamic acid

Ammonium (mmol/l)

Urea (mmol/l)

Urea/ammonium

Time (min) in relation to administration of acetohydroxamic acid

-60 -30 +60 +90 +120 +150 +180 +210

0 1 2 3 4 5 6

Release by intragastric acid, resulting in inappropriate release of the hormone. If this were the mechanism of the hypergastrinaemia, the gastrin concentration should fall rapidly after inhibition of bacterial urease activity. In patients with hypergastrinaemia secondary to achlorhydria the gastrin value falls within five to 15 minutes of intragastric instillation of hydrochloric acid. Likewise, in healthy subjects the intragastric administration of acid at the same time as a meal noticeably suppresses or abolishes the gastrin response to the meal. The lack of a fall in serum gastrin even three and a half hours after suppression of urease activity indicates that the hypergastrinaemia is unlikely to be caused by the acid inhibitory feedback mechanism by bacterial ammonia. We cannot, however, exclude the possibility that the degree of inhibition of urease activity with the acetohydroxamic acid might have been insufficient to allow restoration of the acid inhibitory mechanism.

We have previously reported that increasing H pylori ammonia production threefold for four hours by intragastric infusion of urea does not raise serum gastrin. The present finding that inhibition of H pylori ammonia production does not lower serum gastrin provides further evidence against the hypergastrinaemia being directly linked with bacterial urease activity.

Other mechanisms to explain the hypergastrinaemia need to be considered. It may be related to the chronic inflammatory cell infiltrate which the infection induces in the underlying antral mucosa where the G cells are located. Studies using isolated perfused canine antrum have shown that the T lymphocyte products interleukin-2 and γ interferon stimulate gastrin release. Recent observations by Wyatt et al indicate that hypergastrinaemia correlates more closely with inflammation of the antral mucosa than with infection with H pylori.

This study also shows that normal urease activity is not essential for H pylori to survive within the human stomach. It has been suggested that the organism produces ammonia in order to create an alkaline microenvironment and thus shield itself from the acidic gastric juice. In spite of considerable suppression of urease activity the infection was not cleared in any of our patients, as shown by the repeat breath test 10 days later. Ammonia production may be important in protecting the bacterium from luminal acid when the organism is first ingested but less important when the infection becomes established in the less acidic environment of the deep mucus layer.

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