Acid and gastrin responses during intragastric titration in normal subjects and duodenal ulcer patients with G-cell hyperfunction

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SUMMARY  Amino acid induced acid and gastrin responses during intragastric titration at pH 2.5 and 5.5 were compared in normal subjects and duodenal ulcer patients with G-cell hyperfunction. The latter were identified on the basis of raised basal or maximal acid outputs and increased gastrin responses to feeding. In normal subjects the mixed amino acid meal stimulated only modest increases in serum gastrin, and the highest observed increase was about 30% that after a standard meal. In contrast, in the G-cell hyperfunction group the highest gastrin concentrations were similar to those after a standard meal. In the G-cell hyperfunction group the increment in serum gastrin at pH 2.5 expressed as a proportion of that at pH 5.5 was 0.29 indicating that the capacity of acid to inhibit gastrin release was well established in these patients. Acid secretory rates were close to maximal at both pH 2.5 and 5.5 during intragastric titration in the ulcer patients, but in normal subjects acid output was about 50% maximal at 2.5 and close to maximal at 5.5. The results suggest that the enhanced gastrin response to feeding in G-cell hyperfunction patients is because of increased sensitivity to amino acid stimulation rather than to diminished acid-inhibitory mechanisms.

Duodenal ulcer is likely to be a common end result of a heterogeneous group of disorders. Subgroups of duodenal ulcer patients have been identified on the basis of genetic, epidemiological, pathophysiological, and immunological criteria.1-5 One such subgroup includes patients with hypergastrinaemia and acid hypersecretion.1-6 In at least a proportion of these patients, first degree relatives also show increased gastrin responses to feeding and raised serum pepsinogen (which is directly related to maximal acid output).6 We found that antral gastrin concentrations and G-cell numbers were normal in two patients belonging to this group, suggesting that the raised serum gastrin was because of G-cell hyperfunction, rather than hyperplasia.6 Acid normally inhibits gastrin release so that the combination of increased acid output and raised serum gastrin is inappropriate. There is evidence that in duodenal ulcer patients the mechanisms of acid-inhibition of gastrin release may be defective7 and this could obviously account for the increased serum gastrin in patients with G-cell hyperfunction. It is also possible, however, that the G-cells in the latter patients are more sensitive to stimulation. In the present series of experiments we have examined these possibilities by comparing the gastrin and acid secretory responses with intragastric instillation of a mixture of amino acids in normal and G-cell hyperfunction patients; the technique of intragastric titration8 was used to maintain a constant gastric pH at either 2.5 or 5.5, so that the effects of acid inhibition could be directly examined. The results do not support the idea of impaired acid inhibition of gastrin release in the G-cell hyperfunction group, but instead point to increased sensitivity of the G-cell to amino acid stimulation.

Methods

PATIENTS

Five patients (four men, one woman, Table) with G-cell hyperfunction were identified among a group of patients with endoscopically proven duodenal ulcer. The criteria for inclusion in the G-cell hyperfunction group were as follows: (a) basal
serum gastrin greater than 30 pmol/l and the peak increment in serum gastrin after a standard meal greater than 100 pmol/l; (b) basal acid output greater than 10 mmol/l and maximal acid output following pentagastrin (6 μg/kg) either greater than 35 mmol/l or the ratio of basal to maximal greater than 0.45. In our laboratory 90% of duodenal ulcer patients show an increment in gastrin after feeding of less than 90 pmol/l. The standard meal used for the gastrin studies consisted of two hard boiled eggs, one piece of toast and 120 ml beef bouillion (Oxo) eaten over a period of 10 minutes; serum was taken for gastrin at 20, 30, and 40 minutes after the meal. None of the patients in this study had recent complication by haemorrhage, perforation or pyloric stenosis, nor had they had previous gastric surgery or vagotomy; only one had active ulcer disease within two months of study. For this patient, antacid, and H2-receptor antagonist therapy was discontinued 72 hours before the studies. None of the other patients were receiving treatments at the time of study. The normal subjects (three men, two women, mean age 27 years) were laboratory or hospital workers and did not have a history of gastrointestinal disease (Table). All subjects gave informed consent, and the study was approved by the local ethical committee.

**INTRAGASTRIC TITRATION**

All subjects fasted from midnight before the test. A three-lumen orogastric tube was positioned along the lesser curve as far as the antral region, and the position confirmed by radiography. One lumen opened in the fundus and was used to deliver the initial bolus of amino acids and thereafter to maintain a constant delivery of amino acids and sodium bicarbonate, and for return of the gastric contents removed for pH sampling. A second multi-fenestrated lumen was used for continuous sampling of the gastric contents by aspiration, and a third tube was used for equilibration of gastric and atmospheric pressure. A system of continuous circulation through the stomach was used with a separation of approximately 20 cm between inflow and outflow portals. Subjects sat upright at 45° and turned onto their right side, this ensured good mixing as indicated by steady rates of bicarbonate delivery and a prompt increase on changing from pH 2-5 to 5-5. The pH of the gastric contents was monitored continuously (Radiometer, Copenhagen) and sodium bicarbonate (8.4%) added to the delivery side at a rate sufficient to maintain pH at 2-5 during the first hour and 5-5 during the second hour. At the start of the experiment 200 ml of amino acid solution was instilled into the stomach and the same solution was then infused continuously at 4 ml/min. The solution consisted of 20 g of Aminutrin amino acid powder (Geislick) dissolved in 1 litre of distilled water and supplemented with 1% tryptophan. The osmolarity was adjusted to 325 mOsmol/kg by addition of NaCl. The pH of the solution used during the first hour was adjusted to 2-5. At the end of this period the solution was removed and replaced with 200 ml of the same amino acid solution adjusted to pH 5-5.

**GASTRIN RADIOIMMUNOASSAY**

Serum samples were taken every 10 minutes during the titration and stored at −20°C before radioimmunoassay. Gastrin was measured by radioimmunoassay using a C-terminal specific antiserum, 1296, according to published methods. Routinely serum was assayed at a final dilution of 1:40, but samples from the G-cell hyperfunction group that contained high concentrations of gastrin were frequently assayed at dilutions of 1:100 as well.

**STATISTICAL ANALYSIS**

Results are expressed as mean value ± standard errors. The differences between the two groups were compared by Student’s t test.

**Results**

Gastrin responses to the intragastric instillation of amino acids were significantly greater in the ulcer compared with the control group at all times (p<0.05). In the normal subjects, the increase in serum gastrin was significantly greater than basal (p<0.05) only during titration at pH 5-5 when it amounted to about 25% of the response to a standard meal (Fig. 1). In contrast, in the G-cell
Fig. 1  Serum gastrin concentrations measured by radioimmunoassay during the course of intragastric titration in normal subjects (●) and G-cell hyperfunction patients (○). The increase in gastrin concentrations over basal is shown as the mean ± SE (indicated by vertical bars) for the two groups. For the first hour intragastric pH of the amino acid solution was maintained at 2-5, and for the second hour it was maintained at 5-5 (see Methods for further details). To the right is shown the peak gastrin response to a standard test meal taken on a separate day. Basal gastrin concentration in the normal subjects was 26.4±4.8 pmol/l and in the G-cell hyperfunction patients was 53.8±10.2 pmol/l.

Hyperfunction group the increase in serum gastrin was significant at both pH 2-5 and 5-5. At the lower pH the increase was 30% of the response to a standard meal, and at pH 5-5 the increase in serum gastrin was 112% that in the same subjects after a standard meal. When the gastrin response at pH 5-5 was expressed as a proportion of that to the standard meal the difference between the two groups was statistically significant at all times (p<0.05). An index of the inhibitory effects of acid on gastrin release may be obtained from the ratio of responses at pH 2-5 and 5-5 (taking the response as the mean of the last three 10 minute periods at each pH). In the G-cell hyperfunction group the increment in serum gastrin at pH 2-5 was 0.29±0.076 that at pH 5-5.

Intragastric instillation of amino acids stimulated acid secretion in all subjects. The acid secretion in the G-cell hyperfunction group was significantly greater than in the control subjects at all times (p<0.05). In the normal group the highest rate of secretion at pH 2-5 (mean of the two consecutive highest 10 minute periods) was 62±12% of maximal acid output to pentagastrin; at 5-5 the highest rate of secretion was 135±36% of the maximal acid output (Fig. 2) and these values were significantly different (p<0.05). In the G-cell hyperfunction group the highest rate of acid secretion at pH 2-5 was 124.18% of maximal acid secretion following pentagastrin (Fig. 2), and at pH 5-5 it was 142±11%; these values were not significantly different.

Discussion

The existence of a subpopulation of duodenal ulcer patients with raised acid output and exaggerated plasma gastrin responses to feeding has been recognised for some time. The circulating gastrin is primarily of antral origin as antrectomy reduces the gastrin response to feeding. The term G-cell hyperfunction offers a convenient operational definition for this group of patients; although it is possible that some of these patients may have G-cell hyperplasia, others have a normal density of G-cells. Acid normally inhibits gastrin release and a failure of this mechanism would obviously account for the hypergastrinaemia in G-cell hyperfunction. In the present study we examined this possibility by comparing gastrin responses in normal and G-cell hyperfunction patients during intragastric instillation of amino acids when pH was maintained constant at 2-5 or 5-5 by addition of sodium bicarbonate. The results do not support the idea that there is a defective acid inhibitory mechanism in G-cell hyperfunction patients, but instead they
suggest that gastrin cells might be particularly sensitive to certain modes of stimulation in these patients.

The results obtained with control subjects in the present study are broadly comparable with those of Walsh et al. who used an amino acid solution similar to that used by us but supplemented with cornstarch. In both studies there was little gastrin release at pH 2-5 in normal subjects, and at 5-5 the gastrin response was 30-50% that to a mixed meal (homogenised steak was instilled intragastrically by Walsh et al., compared with eggs and meat extract taken in the normal way in the present study). In an unselected group of duodenal ulcer patients studied by Walsh et al there was significant secretion of gastrin at pH 2-5, and in the G-cell hyperfunction patients studied here there was again significant secretion of gastrin at this pH. There were, however, differences in the degree of inhibition of gastrin release at pH 2-5 compared with 5-5, in the two groups of ulcer patients. Thus in the ulcer patients studied previously there was only about 30% inhibition of gastrin release at pH 2-5 compared with 5-5, whereas in the present study there was about 70% inhibition of gastrin release at 2-5 compared with 5-5. The latter is similar to that reported for normal subjects. Evidently, then, in the selected group of ulcer patients used in the present study the inhibitory effects of acid on gastrin release were much better developed than in the duodenal ulcer patients studied by Walsh et al. Moreover, the results suggest that the principal defect in gastrin release in the G-cell hyperfunction patients lies not in resistance to acid inhibition, but rather in sensitivity to stimulation by amino acids. In particular, when intragastric pH was held at 5-5 (thereby eliminating acid inhibition) the mixture of amino acids used by us evoked an increase in serum gastrin in the G-cell hyperfunction patients that was comparable with the response to a standard meal, while in normal subjects the response to amino acids was only 25% of that to the meal.

The meal used by us evoked about 50% of maximal acid output at pH 2-5 in normal subjects. As the rise in gastrin at this pH was not significant, it seems that non-gastrin mediated pathways are likely to be involved. In the ulcer group the acid response at pH 2-5 was near maximal. This might be because of enhanced activity of the non-gastrin mediated mechanisms of acid stimulation, but circulating gastrin could also make a significant contribution to this acid response.

Among the pathophysiological changes involving acid and gastrin in duodenal ulcer patients, previous studies have implicated enhanced gastrin-responses to feeding, decreased acid-inhibition of gastrin release, increased sensitivity to circulating gastrin and increased acid secretory capacity. Not all these abnormalities apply to all duodenal ulcer patients, and some progress has already been made toward the identification of distinct subgroups of ulcer patients on the basis of their acid or gastrin responses to stimulation. These studies, however, have not for the most part involved control of intragastric pH, and as a consequence the effects of acid inhibition could not be separated from those of increased sensitivity of G-cells to stimulation. The control of intragastric pH by titration in the present study has therefore allowed us, for the first time, to add increased sensitivity of gastrin to stimulation by amino acids to the list of abnormalities shown by a definable subgroup of duodenal ulcer patients. It seems likely that these patients produced exaggerated gastrin responses to modest levels of stimulation, which in turn cause exaggerated acid responses. The gastrin responses to luminal stimulation might involve direct action of luminal stimuli on the apical portions of G-cells, or indirect effects mediated by gastric neurones. The precise mechanisms involved in the control of gastrin release in G-cell hyperfunction patients now need to be established.

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References

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