Gastric acid and pancreatic polypeptide responses to modified sham feeding: indication of an increased basal vagal tone in a subgroup of duodenal ulcer patients

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SUMMARY The effect of sham feeding upon gastric acid secretion and pancreatic polypeptide release was investigated in 28 patients with duodenal ulcer in order to evaluate whether high basal vagal activity is the cause of basal acid hypersecretion in patients with duodenal ulcer and basal secretion higher than 30% of their peak acid output. The patients were divided into two groups based on the ratio of basal/pentagastrin stimulated peak acid output (BAO/PAO) was higher or lower than 0·30: group A n=19 (BAO/PAO ≤0·30) and group B n=9 (BAO/PAO >0·30). Gastric acid response to sham feeding (SAO) was significantly higher than basal level in group A (SAO: 11·4 mEq/h (2·5–20·1) vs BAO: 5·2 mEq/h (0·8–22·9), p<0·01, median (range)) while in group B the acid secretion did not increase with sham feeding (SAO: 9·6 mEq/h (4·5–13·6) vs BAO: 8·8 mEq/h (6·3–13·8) ns, median (range)). A negative correlation (r=-0·618226, p<0·01) was found between acid increase expressed as basal subtracted sham feeding response (SAO-BAO) and BAO/PAO ratio of the entire group of duodenal ulcer patients (n=28) suggesting that the greater is basal acid secretory capacity the smaller is acid increase in response to residual vagal activation. Pancreatic polypeptide response to sham feeding was higher in group A than in group B but no correlation (r=0·20, n=28) nor individual covariation was found between acid and pancreatic polypeptide secretions during vagal stimulation. Sham feeding did not change serum gastrin. It is concluded that an increased vagal stimulation seems to be the cause of basal hypersecretion in a subgroup of patients with duodenal ulcer. The lack of correlation between the pancreatic polypeptide and acid responses to vagal stimulation interferes with the reliability of pancreatic polypeptide as indicator of vagal tone on gastric parietal cells.

An increased vagal tone has been considered one of the possible mechanisms that could explain the increased basal acid secretion in some duodenal ulcer patients.1 2 Cephalic vagal excitation induced by sham feeding results in a potent gastric secretory stimulation.3 5 If high vagal activity in the basal state is the cause of increased basal acid secretion in some duodenal ulcer patients, these patients might be expected to secrete little or no additional acid in response to sham feeding.5 Feldman et al,6 have therefore considered as suggestive of increased vagal tone the failure to increase acid secretion after sham feeding of four duodenal ulcer patients, with markedly increased basal acid secretion, expressed by a BAO/PAO ratio higher than 0·30.

Pancreatic polypeptide is a 36 amino acid polypeptide, localised in a distinct population of pancreatic cells,7 8 released in part under vagal control.9 10 Sham feeding produces a rapid increase in plasma pancreatic polypeptide concentrations 11–13 which is abolished by vagotomy12 and/or by atropine administration.13,14 Pancreatic polypeptide secretion is regulated by tonic vagal cholinergic activity15 and its concentrations fluctuate synchronously with the spontaneous secretions of gastric acid.15 Pancreatic polypeptide and acid secretion,
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however, are not mutually dependent, but the coordination in their basal secretions has led to the suggestion that pancreatic polypeptide secretion might serve as an independent indicator of vagal activity on gastric parietal cells.

In order to assess the hypothesis that high basal vagal activity causes the basal acid hypersecretion in some duodenal ulcer patients (having basal secretion rates higher than 30% of their peak acid output), patients with duodenal ulcer were divided into two groups by means of their BAO/PAO ratio (lower or higher than 0.30) and the relationship between basal acid secretion and acid response to sham feeding was investigated. In addition the reliability of pancreatic polypeptide secretion as indicator of vagal activity on gastric parietal cells has been investigated studying the relationship between gastric acid and pancreatic polypeptide secretion under basal conditions and during vagal stimulation.

Methods

Patients

During a clinical trial to investigate the efficacy of H₂ antagonists on healing of duodenal ulcer, 28 outpatients with active duodenal ulcer, established by clinical and endoscopic examination, on their given informed consent, entered this clinical experimental study. These patients were divided into two groups according to their basal to pentagastrin stimulated peak acid output (BAO/PAO) ratio: group A) 19 patients with BAO/PAO ≤0.30 (five women, 14 men, mean age 41 years, range 29–80; mean weight 73 kg, range 53–93 kg); and group B) nine patients with BAO/PAO >0.30 (three women, six men; mean age 38 years, range 31–60 years; mean weight 67 kg, range 49–100 kg). All patients showed normal basal serum gastrin concentrations (Table 1).

The patients participated in three different studies (pentagastrin stimulation, modified sham feeding, standard meal), on three different days, at intervals of at least two days.

All tests were done after an overnight fast and any anticholinergic or antisecretory medication was withdrawn at least one week before the study.

Pentagastrin Stimulation

Firstly the peak acid output (PAO) as index of maximal secretory capacity was stated by injecting 6μg/kg of pentagastrin (Gastrodagnost, Merck) subcutaneously, and BAO/PAO ratio was calculated in order to allot patients to the above mentioned two groups.

Sham Feeding

Sham feeding was carried out using a modified sham feeding technique. Patients were served an appetising meal prepared in a separate building and consisting of 200 g fillet steak, 150 g French fried

Table 1 Values of BAO and PAO in group A and group B patients with duodenal ulcer

<table>
<thead>
<tr>
<th>Patients</th>
<th>BAO/PAO ≤0.30</th>
<th>BAO/PAO &gt;0.30</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients</td>
<td>BAO</td>
<td>PAO</td>
</tr>
<tr>
<td>1</td>
<td>7.5</td>
<td>26.8</td>
</tr>
<tr>
<td>2</td>
<td>15.1</td>
<td>71.9</td>
</tr>
<tr>
<td>3</td>
<td>7.1</td>
<td>28.6</td>
</tr>
<tr>
<td>4</td>
<td>10.0</td>
<td>52.6</td>
</tr>
<tr>
<td>5</td>
<td>5.2</td>
<td>35.6</td>
</tr>
<tr>
<td>6</td>
<td>22.9</td>
<td>94.0</td>
</tr>
<tr>
<td>7</td>
<td>4.4</td>
<td>27.0</td>
</tr>
<tr>
<td>8</td>
<td>5.8</td>
<td>101.8</td>
</tr>
<tr>
<td>9</td>
<td>2.5</td>
<td>18.3</td>
</tr>
<tr>
<td>10</td>
<td>2.0</td>
<td>28.8</td>
</tr>
<tr>
<td>11</td>
<td>7.0</td>
<td>25.9</td>
</tr>
<tr>
<td>12</td>
<td>4.8</td>
<td>25.6</td>
</tr>
<tr>
<td>13</td>
<td>0.8</td>
<td>41.8</td>
</tr>
<tr>
<td>14</td>
<td>0.9</td>
<td>20.0</td>
</tr>
<tr>
<td>15</td>
<td>5.3</td>
<td>18.9</td>
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<tr>
<td>16</td>
<td>6.5</td>
<td>40.7</td>
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<td>17</td>
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<tr>
<td>18</td>
<td>1.8</td>
<td>15.7</td>
</tr>
<tr>
<td>19</td>
<td>2.1</td>
<td>32.8</td>
</tr>
<tr>
<td>Median</td>
<td>5.2</td>
<td>28.8</td>
</tr>
</tbody>
</table>

BAO group A vs BAO group B: ns. PAO group A vs PAO group B: ns.
potatoes and 250 ml water. The food was tested, chewed, and spat out. The modified sham feeding procedure lasted 30 minutes. This length of time was chosen as it was shown that a modified sham feeding of 30 minutes is a powerful stimulus for vagal activation. Gastric secretion was collected during 60 minutes basal period and for two hours during and after sham feeding. Gastric aspirates were carefully checked for swallowed food particles and few or none were found.

**TEST MEAL**

A meal identical to that served during sham feeding was offered and completed within 30 minutes.

**DETERMINATION OF ACID SECRETION**

A nasogastric tube (modified Levine no 14) was positioned under fluoroscopic control with the radio-opaque tip in the gastric antrum 90 minutes before pentagastrin and sham feeding stimulations.

The gastric residue was completely aspirated and discarded. Collections were made at 15 minute intervals and acid content was titrated to pH=7.0 (PHM62 Radiometer, Copenhagen) with 0.1M NaOH.

Basal acid output (BAO) and sham feeding stimulated acid output (SAO) expressed as milliequivalent H⁺/hour were calculated as the sum of the four consecutive 15 minute outputs during the basal and sham feeding hours respectively. The sum of the two highest consecutive 15 minute outputs after pentagastrin multiplied by two represents the peak acid output by pentagastrin (PAO). Gastric acid secretion was also evaluated as ratio: basal/pentagastrin stimulated peak acid output (BAO/PAO) and basal/sham feeding stimulated acid output (BAO/SAO). In order to determine the residual vagal activation BAO value was subtracted from the acid increase in response to sham feeding (SAO-BAO mEq/h).

**PANCREATIC POLYPEPTIDE AND GASTRIN**

Venous blood samples were obtained from a peripheral vein every 15 minutes, three times before sham feeding and test meal and at 5, 10, 15, 30 minutes during and 45, 60, 75, 90, 120 minutes after. Blood samples were collected in EDTA plus aprotenin and plasma, separated by centrifugation, was stored at -20°C until assayed.

Pancreatic polypeptide radioimmunoassay was carried out by means of anti BPP serum 146-10 (kindly supplied by Dr R A Chance, Lilly Research Laboratories, Indianapolis) at a final dilution of 1:6,000,000. Highly purified bovine pancreatic polypeptide was iodinated using a modification of the chloramine-T method. Highly purified human pancreatic polypeptide was used as standard. Free antigen was separated from bound fraction by adding plasma-coated charcoal. The experimental detection limit was 3.75 pg/tube. The within assay coefficient of variation was less than 5% whereas the between assay coefficient of variation was 14%.

Plasma gastrin concentrations were determined by radioimmunoassay as previously described using antiserum 2604 (kindly supplied by Professor J F Rehfeld).

Serum pancreatic polypeptide and gastrin measured in each sample are expressed as pg/ml.

**STATISTICS**

All results are expressed as median with total range in brackets, because pancreatic polypeptide concentrations are not normally distributed.

Differences within a group were determined by Wilcoxon's matched pair signed rank test. Differences between groups were determined by Wilcoxon's sum rank test. Correlation was determined by Spearman rank correlation coefficient.

A p value of less than 0.05 was regarded as significant.

Pancreatic polypeptide response expressed as integrated pancreatic polypeptide response (IPPR) and calculated according to the following formula:

\[
\frac{\Delta PP_0 + \Delta PP_1 + \Delta PP_2 + \Delta PP_n}{2} \cdot (t_1 - t_0)
\]

\[
+ \frac{\Delta PP_1 + \Delta PP_2 + \Delta PP_n}{2} \cdot (t_2 - t_1)
\]

\[
+ \frac{\Delta PP_n-1 + \Delta PP_n}{2} \cdot (t_n - t_{n-1})
\]

\[\Delta PP\] is plasma pancreatic polypeptide concentrations minus basal pancreatic polypeptide concentrations; \(t\) is time in minutes, subscripts 0, 1, 2, . . . n refer to successive sampling periods (0 = basal sample, therefore PPₐ is always equal to zero).

The integrated gastrin response (IGR) was calculated according to the previous formula.

**Results**

The range of distribution of basal acid output evaluated as a fraction of pentagastrin peak acid output (BAO/PAO) is shown in Figure 1. In the group of patients with BAO/PAO ≤0.30 (group A) the ratio ranged from 0.02 to 0.28 (median: 0.16) and in the group with BAO/PAO >0.30 (group B) the ratio ranged from 0.31 to 0.75 (median: 0.38). Individual variation of basal acid output evaluated in different days during pentagastrin stimulation and modified sham feeding test did not affect significantly BAO/PAO ratio, except for one case that was
Fig. 1  Ratio of basal acid output to peak acid output (BAO/IPAO) and of basal to sham feeding acid output (BAO/IPAO) and SAO-BAO in 19 patients with duodenal ulcer and BAO/PAO ≤0-30 (group A) (close circles) and in 9 patients with BAO/PAO >0-30 (group B) (open circles). Median values are shown as horizontal lines. p-values for differences between group A and group B are: <0-05 (BAO/IPAO); <0-01 (BAO/SAO); <0-01 (SAO-BAO).

discarded.

Figure 1 also shows that gastric acid response to sham feeding, when basal acid output (BAO) is expressed as fraction of sham feeding acid output (SAO) is statistically lower in group A than in group B (BAO/SAO: group A: 0·46 (0·15-1·84) vs group B: 0·94 (0·67-2·35); p<0·01; median (range)). When the sham feeding stimulated acid secretion is expressed as basal subtracted sham feeding response (SAO-BAO mmol/h) group A showed a statistically higher value than group B (group A: 5 (-2·1-14·2) mEq/h vs group B: 1 (-6·1-3·4) mEq/h; p<0·01; median (range)). (Fig. 1).

Figure 2 shows the time course of gastric acid secretion in response to sham feeding in both groups of patients. Gastric acid output in response to sham feeding (SAO) was significantly increased in group A (SAO: 11·4 (2·5-20·1) mEq/h vs BAO: 5·2 (0·8-22·9) mEq/h; p<0·01; median (range)) reaching the peak in the second 15 minute period of sham feeding stimulation, while group B did not show increased acid secretion during and after vagal stimulation (SAO: 9·6 (4·5-13·6) mEq/h vs BAO: 8·8 (6·8-13·8) mEq/h; ns; median (range)).

Figure 2  Time course of the median (range) gastric acid secretion in response to modified sham feeding in twenty eight patients with duodenal ulcer: 19 with BAO/PAO ≤0-30 (group A) (●●●●) range and 9 with BAO/PAO >0-30 (group B) (○○○○) range. The dark dash (---) indicates the length of sham feeding. The acid output in the group A patients (●●●●) is significantly increased (p<0·01) at 15 and 30 min during sham feeding.

Figure 3 shows that in the group of 28 duodenal ulcer patients (group A + group B) there is a negative correlation (r=-0·6118226, p<0·01) between BAO/PAO ratios plotted with SAO-BAO – that is, the greater the basal secretion expressed as percentage of peak pentagastrin-stimulated acid output (BAO/PAO), the smaller the acid increase (SAO-BAO) in response to residual vagal activation.

Modified sham feeding led to a significant and rapid increase in serum pancreatic polypeptide concentrations over the basal values in both groups of duodenal ulcer patients (data not shown).

No significant difference was found in the basal values of pancreatic polypeptide between the two groups of patients (group A: 70 (26-240) pg/ml vs group B: 75 (28-268) pg/ml; median (range)).

Table 2 shows the pancreatic polypeptide response to sham feeding evaluated as integrated response (IPPR) during the 30 minutes stimulation;
it was two times higher in group A than in group B (973 vs 445 pg/ml/30 min), but p value barely missed statistical difference (p=0.05). The pancreatic polypeptide response to standard meal was similar in the two groups of patients (Table 1).

Comparing gastric acid secretion and pancreatic polypeptide serum concentration measured at the start of each 15 minute collecting period in the individual patient (seven determinations), no positive covariation was found with a median correlation coefficient of 0.287, range (−0.006–0.942) (n=28). During the 30 minutes of sham feeding test no significant correlation was found between integrated pancreatic polypeptide response (IPPR<sub>sf</sub>) and the increase in gastric acid secretion (SAO-BAO, mEq/30 min) (Fig. 3).

Basal serum gastrin concentrations were similar in both groups of patients (group A: 53 (28–95) pg/ml vs group B: 48 (30–100) pg/ml; ns; median (range)).

Sham feeding did not significantly affect serum gastrin concentrations in both groups of duodenal ulcer patients (Integrated gastrin response (IGR): group A: 37.5 (−227–610) pg/ml/30 min vs group B: 24.5 (−290–530) pg/ml/30 min; ns; median (range)).

**Discussion**

This study provides evidence that duodenal ulcer patients with high basal acid secretion, expressed by a BAO/PAO ratio higher than 0.30, and normal serum gastrin levels, do not show increased acid secretion in response to cephalic vagal activation – that is, modified sham feeding procedure.

This study based on a larger series of duodenal ulcer patients, confirms the previous observation reported by Feldman et al. Both our results and those of Feldman could be explained by the following hypothesis: if high vagal activity under basal conditions is present, little or no additional acid will be secreted in response to vagal stimulation by sham feeding. This hypothesis is further supported by our finding of a negative correlation between basal secretion, expressed as percentage of peak stimulated-acid output (BAO/PAO), and basal subtracted sham feeding response (SAO-BAO) in all 28 patients with duodenal ulcer (Fig. 4). Thus, in duodenal ulcer patients, either with or without basal hypersecretion, the higher their BAO/PAO ratio, the lower the acid response to vagal stimulation.

The amount of ‘vagal tone’ could hence be expressed by the degree of BAO/PAO ratio.

The BAO/PAO ratio, however, could vary in the same subject, owing to wide variation in basal acid secretion which is affected by such factors as environmental conditions, and emotional state of the individual, with cyclic variation during the day and great differences from day to day.

In our population of patients with active duodenal ulcer only in one case did variation of basal secretion affect BAO/PAO ratio. This subject, who was discarded, had been examined on two different occasions; pentagastrin stimulation during clinic and endoscopic ulcer recurrence (BAO/PAO=0.32) and

Table 2. Pancreatic polypeptide response to sham feeding (IPPR<sub>sf</sub>) and to meal (IPPR<sub>mr</sub>) expressed as 30 min integrated response (pg/ml/30 min) in the group A and in the group B patients with duodenal ulcer

<table>
<thead>
<tr>
<th>Group</th>
<th>BAO/PAO</th>
<th>IPPR&lt;sub&gt;sf&lt;/sub&gt;</th>
<th>IPPR&lt;sub&gt;mr&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>A (n=19)</td>
<td>≤0.30</td>
<td>973 (−199–6075)</td>
<td>8887.5 (3245–26500)</td>
</tr>
<tr>
<td>B (n=9)</td>
<td>&gt;0.30</td>
<td>445 (−75–1212)</td>
<td>9431.3 (3362.5–20250)</td>
</tr>
<tr>
<td>p=0.05</td>
<td></td>
<td>ns</td>
<td></td>
</tr>
</tbody>
</table>

**Fig. 3 Correlation (r=0.20) between integrated pancreatic polypeptide response (IPPR<sub>sf</sub>) and gastric acid increase (SAO-BAO) during the 30 min sham feeding in 28 patients with duodenal ulcer. The open circles beneath basal values, shown by broken lines, indicate three patients who have a gastric acid increase without any pancreatic polypeptide response to sham feeding and three other patients with good pancreatic polypeptide response and no acid increase.
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Fig. 4 Correlation (r = -0.6118226) between acid increase in response to modified sham feeding (SAO-BAO) and basal secretion expressed as a percentage of peak pentagastrin-stimulated acid output (BAO/PAO) in 28 patients with duodenal ulcer.

...pentagastrin-stimulated acid with polypeptide data should be contradictory by found feeding polypeptide pancreatic feeding. sham patients resembles secretion basal in group 1 minute response rule Gastroenterology 1983; 84:1211.

A preliminary report of this work has been published in abstract form (Gastroenterology 1983; 84:1211).

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