Enterooxyntin Release from Isolated Perfused Canine Jejunum

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A humoral factor may mediate the intestinal phase of gastric acid secretion. An ex vivo perfused segment of canine jejunum maintained by an oxygenated asanguinous physiologic perfusate was used to test for release of an enterooxyntin (EO) in response to balloon distention at 30 mm Hg for 15 min. Gastric acid secretion in guinea pig fundic mucosa was determined indirectly by a quantitative cytochemical bioassay (CBA) of oxyntic cell hydroxyl ion production (HIP). An increase in the optical density (OD) caused by the cytochemical stain in the oxyntic cells reflects HIP, an index of acid secretion. Basal OD for segments with distention was 16.6 ± 0.53 and for those without 15.5 ± 0.68 (NS). Results are expressed as mean change of OD from basal (mean ΔOD ± SEM).

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>10</th>
<th>30</th>
<th>60</th>
<th>120</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean ΔOD, distention</td>
<td>4.9 ± 1.6*</td>
<td>5.9 ± 1.1*</td>
<td>7.2 ± 1.2*</td>
<td>5.2 ± 0.9*</td>
</tr>
<tr>
<td>Mean ΔOD, no distention</td>
<td>-0.6 ± 1.0</td>
<td>-0.1 ± 0.8</td>
<td>0.58 ± 1.0</td>
<td>0.7 ± 1.2</td>
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* Significant stimulation of HIP (P < 0.02) by EO.

EO caused greater stimulation of HIP than gastrin or histamine. EO was heat stable. Trichloroacetic acid treatment decreased EO activity as did pronase digestion suggesting that EO is composed of one or more peptides. Conclusion: EO, an acid secretagogue, is a humoral agent probably composed of one or more peptides and is released by small bowel distention. Mechanical distention of the small bowel may be an important mechanism for the perpetuation of gastric acid secretion. The ex vivo perfused jejunal segment in conjunction with the CBA are ideal tools with which to study mechanisms of release of EO and the mechanism of action of EO on the oxyntic cell.

INTRODUCTION

The presence of an intestinal phase of gastric secretion has been recognized since the work of Le Conte [17] and Pavlov [26] in the early twentieth century. In 1953, Sircus demonstrated that simple mechanical distention of the jejunum in the dog resulted in increased acid secretion from a denervated gastric pouch and suggested that a humoral agent caused the intestinal phase of gastric acid secretion [29]. Perfusions of Thiry-Vella loops, small intestine and gastrojejunostomies with nutrient solutions in dogs with Heidenhain pouches, support the concept that a humoral mediator of the intestinal phase response exists [1, 2, 7, 14, 16, 31]. In both dogs and man, a humoral agent has been suggested as the agent causing the increase in gastric acid secretion following portacaval shunting [12, 22, 24]. Grossman proposed the name enterooxyntin (EO) for the putative humoral substance released by the gut which stimulates the gastric oxyntic cell to produce acid [9].

We have used distention of an ex vivo isolated perfused segment of canine jejunum to release an acid secretagogue which satisfies the definition of EO. Using a sensitive quantitative cytochemical bioassay (CBA) of oxyntic cell hydroxyl ion production (HIP) as an in-
dex of acid secretion [13], we measured the ability of EO to stimulate the oxyntic cell. In the CBA, EO was at least as efficacious as histamine or gastrin. Preliminary characterization of EO suggests that it is composed of one or more relatively heat stable peptides.

METHODS

Jejunal perfusion. Mongrel dogs of both sexes weighing between 15 and 25 kg were anesthetized with pentobarbital sodium (Nembutol), intubated, and ventilated. Through a midline incision, a 30 cm segment of jejunum beginning 5 cm distal to the ligament of Treitz was isolated. The jejunum was divided and the lumen was perfused with normal saline at a rate of 1.67 ml/min. The cephalad mesenteric artery and vein were then cannulated and pulsatile perfusion begun at a flow rate of 35 ± 5 ml/min. Any remaining attachments of the segment of jejunum were severed and the specimen placed on the perfusion apparatus. The jejunal segment was maintained on a sheet of polyethylene suspended in a water bath at 37°C in a closed chamber.

The perfusate consisted of a modified Krebs–Hensel solution containing: sodium 145 meq/liter, glucose 7 mmole/liter, bovine serum albumin 0.6 mmole/liter, phosphorus 3.8 meq/liter, and dextran (avg. M.W. 70,000) 8.5 mmole/liter.

The perfusate was heated and oxygenated with a humidified 95% O₂–5%CO₂ mixture using a modified rotating disc oxygenator (Sarnes Co., Inc.) suspended in the water bath. Organ perfusion was provided with a pulsatile pump (Harvard Apparatus, Model 1405). A single-pass perfusion technique was used so that fresh oxygenated perfusate entered via the arterial cannula, exited the segment via the venous cannula, and was retained for assay. This system is similar to that described by Eckhauser et al. for ex vivo perfusion of the pancreas and duodenum except that the perfusate is not recirculated [6].

Three milliliters of venous effluent were taken at 0, 10, 30, 60 and 120 min after a 10 min baseline period, placed in a polyethylene tube, and immediately frozen in a solid CO₂–ethanol bath at −70°C.

Perfusions without distension (control). After removal from the dog, the bowel was perfused for 130 min. Arterial flow was maintained at 35 ± 5 ml/min and the lumen was perfused at 1.67 ml/min with normal saline. No luminal distention was applied. Samples of effluent for testing in the CBA were taken 10 min after beginning the perfusion to determine basal activity, and labeled as time 0 min. Additional samples were taken at 10, 30, 60, and 120 min. Samples of jejunal tissue for histologic examination were obtained before and after the perfusion. Minimal differences were observed between these sections consisting of mild edema and loss of the tips of some villi, suggesting that the bowel remained viable.

Perfusions with distention (experimental). After removal from the dog, the bowel was perfused for 130 min. Arterial flow was maintained at 35 ± 5 ml/min. Samples of effluent for testing in the CBA were taken 10 min after beginning the perfusion to determine basal activity and labeled as time 0. Immediately after obtaining this sample, the lumen was intubated with a latex rubber balloon which was inflated to 30 mm Hg. After 15 min of bowel distention, the tube was removed and perfusion with normal saline at 1.67 ml/min was resumed. Additional samples of perfusate were obtained at 10, 30, 60, and 120 min after time 0 for testing in the CBA. Sections of jejunum before and after distention and perfusion were examined histologically and showed minimal changes between the sections consisting of tissue edema and loss of some tips of villi.

Quantitative cytochemistry of oxyntic cell function. After an overnight fast, female guinea pigs weighing 450 to 500 g were sacrificed by asphyxiation in a nitrogen atmosphere and their stomachs were rapidly removed. A 3–4 cm strip of gastric fundus was excised from a standard area just below the esophagogastric junction. The tissue was rinsed in 0.025 M HEPES (Sigma) buffer, (pH adjusted to 7.0
with 10 M KOH solution) and divided into 3–5 mm pieces. These were snap-frozen in a beaker immersed in an n-hexane, solid CO₂, absolute alcohol freezing mixture. The frozen portions of tissue were used within 72 hr of freezing.

Sections (18 μm) were cut from the mounted tissue in a −20°C cryostat. Sections were mounted on glass slides using a template to reliably position the sections in a predetermined area and stored for no longer than 6 hr in the cryostat chamber before use in the assay.

The slides were placed flat, section upwards, in a slide tray and allowed to equilibrate to room temperature for 10 min. The test solutions were diluted to desired concentrations in 0.025 M HEPES buffer at pH 7.0. One hundred microliters of each solution was delivered by a dispensing apparatus designed to deliver precise volumes of test solution directly and simultaneously to 24 sections and allowed to react with the sections at room temperature (20°C) for 90 sec. The reaction was terminated by the addition of the staining reagents. Within each experiment each test solution was reacted with tissue sections in triplicate.

Production of hydroxyl ions (HIP) was quantified by a modification of Hansson's method [10] using 10.5 mM CoSO₄, 53 mM H₂SO₄, 157 mM NaHCO₃, and 1.17 mM KH₂PO₄ in 0.1 M HEPES buffer at pH 7.4 with 0.001% gum tragacanth. This solution was applied so as to just cover the sections. The staining reaction was allowed to proceed with gentle agitation for 2 min at 20°C and then the sections were washed twice with cold water after which they were exposed to a saturated solution of H₂S in water for 90 sec, again washed in water and mounted in Farrants Medium (G. D. Searle Diagnostics, High Wycombe, Bucks, England). The staining procedure led to the formation of visible brown precipitates of CoS within the sections: the most deeply stained cells were the oxyntic cells. The amount of precipitate formed is a reflection of the number of hydroxyl ions trapped [10] and was quantitated by means of a M85 scanning and integrating microdensitometer [5]. This staining is based on the reaction between Co³⁺ and OH⁻ ions produced during stimulation of the oxyntic cell. Whether these hydroxyl ions are produced by carbonic anhydrase (CA) or are produced by the proton pump is unknown. It has been suggested that CA catalyzes directly the reaction between the hydroxyl ion produced by the proton pump and CO₂ resulting in the formation of a bicarbonate ion [27]. This process requires fewer reactions than that proposed by Maren [20] and fits with observed events just as well. Thus, we believe the staining actually quantifies hydroxyl ions produced by the proton pump. This would provide a direct measurement of hydrogen ion secretion by the oxyntic cell and should parallel HIP mole for mole.

The cells selected for each reading were those which completely filled a mask which, when used with the 25X objective lens has a field diameter of 20 μm. Fifteen to twenty oxyntic cells in each of the randomized coded sections were read at 550 nm. The same number of measurements was made from the muscularis mucosa in each section. Readings from the muscularis reflect non-specific absorption of CoS to tissue and were therefore, subtracted from those of the oxyntic cells. To correct for possible instrument variation, the densitometer was calibrated by taking readings of a standard filter of known optical density before and after taking readings from each section. Values for the optical density of the CoS precipitate in each section were read as integrated extinction and were expressed as a percent using the following equation: (mean extinction of oxyntic cells) − (mean extinction of muscularis) × 100/D1, where D is the reading of a standard filter with an optical density of 1. The optical density thus calculated allowed comparisons between sections.

Characterization of EO. Preliminary characterization of the nature of EO was carried out on samples of perfusate shown to cause oxyntic cell HIP. HIP was measured before and after boiling for 10 min to test heat stability. To determine if EO was a peptide, a
sample containing EO was treated with 20% cold trichloracetic acid and the residual HIP measured in the supernatant. The importance of peptide bonds was determined by measuring HIP before and after a sample containing EO was treated with Pronase to destroy peptide bonds [11].

**Statistics.** The significance of the difference between the mean optical density (OD) of the oxyntic cells during the basal periods for the control and experimental groups was tested using Student's t test. Following the basal period, the change in OD between the basal value and subsequent values at 10, 30, 60, and 120 min was determined for the experimental and control groups and the significance of the difference between groups in the mean change in OD (meanΔOD) was determined for each time point using Student's t test. A P value less than 0.05 was accepted as significant.

**RESULTS**

During the 10 min basal period, treatment of oxyntic cell sections with venous effluent from the control perfusions (n = 3) resulted in a mean OD of 15.5 ± 0.68, while treatment with effluent from the experimental perfusions (n = 3) resulted in a mean OD of 16.6 ± 0.53. This difference was not statistically significant.

The mean OD in the CBA did not change from basal in the control group. In the experimental group, there was a significant rise in OD (P < 0.02) caused by the venous effluent (Fig. 1).

**FIG. 1.** Comparison of hydroxyl ion production (HIP) with and without distention of the segment. Mean Δ optical density reflects change of hydroxyl ion production from basal. ○ = Significant stimulation of HIP (P < 0.02) above basal. * = Significant difference between control and experimental.

**FIG. 2.** Comparison of EO, gastrin and histamine on oxyntic cell HIP. Mean optical density reflects HIP.
Comparison with gastrin and histamine. The peak value obtained from venous effluent of distended jejunum was 22.32 ± 1.29 seen at 60 min. This compares to a peak of 19.44 ± 0.22 for gastrin (10^{-12} M) and 17.97 ± 0.24 for histamine (10^{-14} M) (Fig. 2).

Physical characteristics of EO. After boiling venous effluent from distended segments, HIP in oxyntic cells was 59% of the value before boiling.

Treatment with 20% trichloroacetic acid and pronase completely prevented stimulation of HIP in the oxyntic cell by the perfusate samples. The results are shown in Fig. 3.

DISCUSSION

The cytochemical bioassay (CBA) used to measure hydroxyl ion production (HIP) by the oxyntic cell has been shown to reflect activation of guinea pig oxyntic cells by gastrin, histamine, and carbamylcholine, the three known acid secretagogues [13, 19, 28, 30]. Although this method does not directly measure hydrogen ion secretion, all current proposals to explain hydrogen ion secretion by the oxyntic cell require generation of intracellular hydroxyl ions [27]. Thus, measurement of hydroxyl ion production may well reflect acid secretion.

A number of studies indicated that the jejunum is the site of origin of an acid secretagogue tentatively termed enterooxyntin [1, 2, 4, 8, 9, 16, 25, 31]. However, it has not been clearly established that a hormone rather than the infused nutrient such as amino acids was the functional secretagogue [21]. For this reason, we chose to examine the effects of simple distention of the canine jejunum on the release of EO. The use of an ex vivo preparation offers several benefits. Removal of the jejunum severs all neural components and vascular components extrinsic to the segment of jejunum. Vascular and luminal flow are controlled and may be varied. Use of an artificial perfusate avoids contamination by other agents contained in plasma. A once through perfusion system avoids accumulation of waste products or active agents and allows an accurate evaluation of the time course of release. In addition, no major site of metabolic degradation such as the liver is present.

Earlier studies in our lab showed that the perfused bowel segment remained viable for about 2 hr. Histologic changes in the jejunum perfused for 2.5 hr or less showed minimal changes consisting of edema and loss of a few villi tips. Prolonged perfusion (>3 hr) or perfusion at reduced flow rates (<30 ml/min)
resulted in gross histologic evidence of cell death with loss of mucosa and necrosis of the muscularis.

Using this system, a definite and sustained release of a substance from the distended jejunum which stimulates HIP in guinea pig oxyntic cells was found. Since HIP was the same under basal conditions and there was no change during perfusion without distention, the release of EO is presumably due to distention of the jejunum and not the perfusion process itself. Because this substance is released into the venous effluent and stimulates the oxyntic cell, we have used the term enterooxyntin (EO) as first proposed by Grossman [9] to describe an agent released from the bowel causing stimulation of the oxyntic cell.

The mechanism of release of EO is unknown. Possible explanations include local neural pathways involving the enteric nervous system or direct action of stimuli on cells containing EO. Our model does not answer this question and either possibility may occur. Direct trauma to EO-containing cells by the perfusion process and balloon distention may cause release. Since no release was seen in control perfusions, the stimulus to secretion is likely to have been balloon distention.

After release of EO was established, a preliminary characterization was performed. EO was identified as a likely acid secretagogue. Comparison with histamine and gastrin indicated that EO was at least as efficacious a stimulus of HIP as gastrin and histamine. This may reflect a single more potent agent, the additive effects of several different agents released into the perfusate, or potentiation of local histamine within the gastric tissue. Treatment with trichloroacetic acid destroyed HIP stimulation indicating that the active substance was protein in nature. Pronase digestion destroyed the ability of EO to stimulate HIP indicating that peptide bonds are important in the action of EO. EO was relatively heat stable. These findings support the concept that the circulating agent or agents which comprise EO are made up of one or more peptides.

Since EO is believed to be at least partially responsible for the intestinal phase of acid secretion, it may be important in the development of peptic ulcer disease. Konturek et al. studied the effects of intestinal meals in patients with and without peptic ulcer disease. A distinct intestinal phase of gastric secretion was observed with a higher level of acid production present in patients with duodenal ulcer [15]. Bugat et al. have shown that a gastric secretagogue that is not gastrin exists in patients with peptic ulcer disease [3]. EO may contribute to this effect. Experiments by Orloff et al. have suggested that the increase in gastric acid secretion after portacaval shunting is mediated by a humoral agent which escapes hepatic degradation and is released from the jejunum [23, 25]. Our findings do not exclude the possibility that EO may be extracted by the liver and the stimulation of HIP we see may be due to an increased level of EO which might not be seen after circulation through the liver. Further study of the mechanisms of release of EO and its action on the oxyntic cell may provide a better understanding of the role of EO in acid secretion and peptic ulcer disease.

In summary, our study demonstrates the release into the venous system of an agent from the perfused, distended jejunum which was capable of stimulating the oxyntic cell, thus fulfilling the definition of enterooxyntin. Preliminary characterization suggests that EO is composed of one or more peptides. The ex vivo perfused jejunal segment and CBA are useful tools in the study of the release and action of EO.

REFERENCES