Opposite effects of bombesin on insulin and gastrin response to food in humans

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SUMMARY The effect of bombesin on insulin and gastrin response to a standard labelled meal was studied in eight healthy male volunteers. The gastric emptying of solids was simultaneously evaluated. During intravenous infusion of the peptide (5 ng/kg/min) the insulin release after eating was greatly reduced whereas food stimulated gastrin release was significantly enhanced. Both effects of bombesin are likely to be connected with the marked inhibition of gastric emptying induced by the peptide.

Bombesin is a frog skin tetradecapeptide which has been found to have similar immunologically characterised counterparts in mammalian gut. An heptacosapeptide has been isolated from porcine non-antral gastric tissue: the so called gastrin releasing peptide (GRP). The homology of its C-terminal decapptide with C-terminal decapptide of bombesin is impressive. As a consequence, both these peptides have the same spectrum of biological actions, although some qualitative differences have been reported. Many radioimmunological and immunocytochemical observations indicate that bombesin like peptides are confined to neurones and fibres of the intramural plexus of the gut. As a consequence, they have been put forward as putative neurotransmitters in mammalian gut.

Bombesin was shown to be a potent and polivalent releaser of several gastrointestinal hormones both in animals and humans. Its gastrin releasing effect, first demonstrated in dogs, was repeatedly confirmed in other animal species including man. On the contrary, results concerning its effect on insulin secretion have been contradictory. A stimulation of insulin release was observed in dogs, cats, and man whereas an inhibition was reported in rats. In vitro studies have provided conflicting results, as well. In fact, whereas bombesin stimulates insulin secretion in isolated and perfused pancreas from dogs and rats it inhibits the hormone release from isolated rat pancreatic islets.

The recent discovery of bombesin like immunoreactivity in the human pancreas suggests that this peptide may also affect human pancreatic secretion. Therefore we decided to study in man its effect on insulin response to food. Because bombesin was shown to affect gastric emptying, the effect of the peptide on emptying rate was simultaneously evaluated.

Preliminary results of the present investigation have been presented at the Italian Society of Gastroenterology (June 1981) and appeared in abstract form.

Methods

SUBJECTS

Eight male volunteers (average age 24 years) participated in the study after having given written informed consent. They were medical students without any gastrointestinal, endocrine, or metabolic disease. When compared with average weights based on height and age (Tables Geigy), all the subjects were within 10% of their predicted (ideal) value.

EXPERIMENTAL DESIGN

After an overnight fast, all the subjects reached the laboratory at 08.30 am. They remained in the supine position during the entire period of the study, except when eating when they were in a semirecumbent position. Two indwelling intravenous cannulae were inserted in the forearms: one for the infusions, the other for blood sampling. All the volunteers underwent the test twice, having an intravenous infusion of bombesin on one occasion and a control saline
infusion on the other, in a single blind randomised order. Infusion of bombesin (gift of Dr Chiara De Paolis, Farmitalia-Carlo Erba Research Labs, Milan, Italy) in saline (5 ng/kg/min corresponding to 185 pmol/kg/h) began 15 minutes before the meal and continued for 60 minutes. Blood samples for insulin, gastrin, and glucose assays were taken before and after the meal (see Figures).

During the infusion of the peptide, three subjects suffered slight discomfort in the form of nausea (one case) or abdominal discomfort (two cases), but this disappeared spontaneously after stopping the infusion.

**MEASUREMENT OF GASTRIC EMPTYING**

Gastric emptying was measured by a method previously described and validated. Briefly, the subjects ate a standard meal (containing 70 g carbohydrates, 24 g proteins, and 24 g fat) of hamburgers and potatoes purée. Its volume was 400 ml and its caloric value 600 calories. The purée was prepared instantly from commercially lyophilized potatoes, water, milk and a small quantity of butter. This was lightly homogenised using an electric blender after addition of 250 μCi of 90mTc sulphur colloid.

The subjects were asked to eat the meal in about eight minutes followed by a drink of 200 ml partially skimmed milk. Soon after finishing the meal, they returned to the supine position and radioactivity was continuously recorded for 90 minutes by means of a scintillation detector (Nuclear Accesseries, Castiglione O, VA) positioned over the stomach. All data recorded during the time of the study were corrected for radioisotope decay. From these corrected data the percentage of the volume remaining in the stomach (as percentage of radioactivity) was obtained at various time intervals (every 10 minutes).

The radiation absorbed dose, using this method, is very low. Dosimetric calculations done in our laboratory, however, suggested that male subjects are more suitable than female for this type of study. Indeed, the absorbed dose to the ovaries was higher than that absorbed to testes.

**LABORATORY ANALYSIS**

Blood was collected in prechilled tubes containing 1.2 mg of EDTA per ml of blood. Specimens were centrifuged within one hour and plasma stored at −20°C until the time of assay.

Protein free filtrates for plasma glucose were made immediately after the blood samples had been obtained. Plasma glucose was then determined on the supernatant by using a semimicro glucose-oxidase method.

Plasma immunoreactive insulin (IRI) was assayed by the double antibody method of Hales and Randle, using a Liso-phase insulin system (Lepetit Diagnostic Products, Milan). In this assay the separation of insulin bound to antibody from unbound insulin was carried out by affinity chromatography (Sepharose gel to which the second antibody has been covalently linked). Human insulin was used as a standard and diluted in insulin-free human plasma. In our hands, the method has an intraassay coefficient of variation of 3%; the minimal sensitivity of this method is 2 μU/ml.

Plasma immunoreactive gastrin (IRG) was measured by radioimmunoassay using CIS reagents. The antiserum, produced in rabbits against a G-17 BSA-conjugate, crossreacted with G-17 I by 100%, with G-17 II by 73% and with G-34 by 38% on a weight basis. Synthetic human gastrin (ICI) was used as a standard and diluted in gastrin free human plasma. The method has an intraassay coefficient of variation of 10% and its sensitivity is sufficient to measure 20 pg/ml of IRG in a plasma sample.

Bombesin did not interfere in the insulin or gastrin radioimmunoassay at the concentrations used. In both assays all samples from a single subject were assayed in the same run in order to avoid interassay variations.

**EVALUATION OF DATA**

The fitting of the standard curves of radioimmunoassays was done by a LKB-WALLAC computer on line with the Rack-Gamma using the spline approximation. Quantitative evaluation of hormone production was made by an integration of areas under the curve of immunoreactivities in plasma, after subtraction of the basal values. From this an immunoreactive insulin production rate in μU/ml-min and an immunoreactive gastrin production rate in pg/ml-min were calculated by dividing the respective area by the time (min) of the duration of the test.

There are many approaches to analysing and reporting gastric emptying data. Plotting the emptying curves is the best way to examine and display the details of the emptying process. For comparison between different groups of subjects or different treatments, however, the whole emptying curve rather than specific time points should be taken into account. We used two ways to analyse gastric emptying data. First, we calculated an emptying half-time from the regression lines for the log gastric content against time. Second, we calculated from each curve an emptying index according to the formula reported below:

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EI = \left[ \frac{100-RR_{go}}{\text{Ago}} \right] \times 100
\]
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where RRgo is the residual radioactivity 90 minutes after the meal and Ago is the area under the radioactivity time curve. This index was also adopted because its use does not require a choice to be made with regard to the emptying pattern.

All values are presented as a mean±SEM. The two way analysis of variance was used for statistical evaluation of data. Student’s t test for paired data was used to check differences between the means of summarised responses (emptying half-times, emptying indexes, and integrated hormone responses).

Results

HORMONE RESPONSE TO FOOD

Bombesin, administered by intravenous infusion (5 ng/kg/min) to healthy volunteers blunted the expected postprandial increase in plasma glucose and concomitantly blocked the insulin response to meal (Fig. 1), an effect which lasts as long as the infusion. After stopping the peptide administration, plasma glucose increased slightly and an insulin response became evident so that plasma hormone levels 60 to 150 minutes after the meal were not significantly different from those observed during saline infusion (Fig. 1).

The background infusion of bombesin increased the gastrin response to food (Fig. 2) as already described in dogs and man.

Figure 3 depicts the immunoreactive gastrin secretory rates after the meal with or without bombesin infusion together with the immunoreactive insulin secretory rates. It is evident that bombesin exerts an opposite effect on gastrin and insulin response to meal; that is an increase of the former and a decrease of the latter.

GASTRIC EMPTYING

During the infusion of the peptide a considerable delay in gastric emptying of solids was observed (Fig. 4) Indeed, bombesin caused a significant increase in emptying half-times and a significant decrease in emptying indexes (both expression of a decrease in emptying rate, see Table).

Fig. 1 Plasma insulin and glucose after meal during saline (continuous line) or bombesin (broken line) infusion in eight male volunteers. Each point refers to the mean of the values obtained from eight subjects. Vertical bars are standard errors. Arrows represent the time of beginning and of stopping the infusion, respectively. B (basal) refers to the mean of two consecutive blood samples (-30 and -15 min) obtained before bombesin or saline infusion. The hatched horizontal bar represents the duration of meal. *p<0.05 **p<0.02
Fig. 2  Plasma gastrin response to food during saline (continuous line) or bombesin (broken line) infusion in eight male volunteers. Each point refers to the mean of the values obtained from eight subjects. Vertical bars are standard errors. Arrows represent the time of beginning and of stopping the infusion, respectively. B (basal) refers to the mean of two consecutive blood samples (−30 and −15 min) obtained before bombesin or saline infusion. The hatched horizontal bar represents the duration of meal. *p<0.05  **p<0.02.

Fig. 3  Integrated responses of immunoreactive gastrin and immunoreactive insulin during saline (white columns) or bombesin (black columns) infusion in eight male volunteers. Each column represents the mean of the values obtained from eight subjects. Vertical bars are standard errors.

Fig. 4  Time course of residual radioactivity in stomach of eight male volunteers during saline (●) or bombesin (○) infusion. Each point represents the mean of the values obtained from eight subjects. Vertical bars are standard errors. Arrow represents time of stopping of the infusion.

Discussion

In 1971 Erspamer first reported that intravenous infusion of alytesin, a frog peptide of the bombesin family, causes a three-fold increase of immunoreactive insulin levels in dogs. Some years later Brown and coworkers showed that, like other peptides, bombesin can influence glucose homeostasis and affect insulin secretion. Afterwards, insulin release was found to be stimulated by the peptide in cats, dogs and man. In addition, preliminary experiments of our laboratory indicated that bombesin also enhances the insulin response to intravenous glucose.

Results obtained in the present investigation show that the insulin response to a standard mixed meal is almost completely suppressed when bombesin is infused intravenously. This unexpected finding, however, is only in apparent contrast with previous data. The lack of postprandial hyperglycaemia and the consequent inhibition of insulin release are likely to be connected with the strong inhibition of gastric emptying, the delay of which slows the entry of the chyme into the duodenum. In accordance with our results, Salera et al reported quite recently that during bombesin administration to healthy volunteers the oral glucose load was followed by a flattened glucose curve and a significant inhibition of insulin release. On the other hand, Jorde et al found that in Billroth II resected
patients an 80 minute ingestion of glucose was
followed by serum insulin concentrations signifi-
cantly lower than that observed after a two minute
ingestion of the same quantity of glucose. These
data suggest that the rate of glucose delivery into the
intestine is of importance in the insulin response to
oral stimuli (glucose or mixed meal). In line with
these conclusions, Thompson et al.\(^48\) showed that
gastric emptying is a determinant of oral glucose
tolerance test and actually suggested the possible
use of this latter for the assessment of emptying rate.

The flat plasma glucose pattern observed in our
volunteers during bombesin infusion raises the
possibility of an interference of the peptide with the
intestinal glucose absorption, either directly or
through an inhibition of intestinal motility.\(^39\) In a rat
model, however, bombesin failed to affect jejunal
glucose absorption.\(^40\) In addition, we\(^41\) recently
showed that there are no differences in plasma
glucose concentrations during bombesin or saline
infusion after an intraduodenal glucose load in
healthy volunteers. These data suggest that, on the
whole, the motor effect of bombesin on the intestine
is of less importance in comparison with the
enormous capacity of the upper intestine for glucose
absorption.\(^42\)

As regards gastrin secretion, our data show that
bombesin is not only a potent gastrin releaser\(^10\)
but also enhances the gastrin response to meal.
Various factors are thought to be involved in the
regulation of gastrin release. Among these, the meal
residence time in the stomach represents the most
often neglected one. Quite recently, Hirschowitz\(^43\)
demonstrated in dogs that the longer hypertonic
liquid meals remain in the stomach, a more sustained
gastrin response is seen than that with rapidly
emptying, hypotonic fluids. In our experimental
conditions gastric emptying of solids was strongly
inhibited by bombesin and consequently the time
food remained in the stomach was considerably
increased. This longer time of contact between
digesta and antral G cells may account, at least
partially, for the increased gastrin response to meal.

To summarise, bombesin administration strongly
reduced insulin response to meal and significantly
enhanced food induced gastrin release. Both these
effects are likely to be connected with the marked
inhibition of gastric emptying induced by the
peptide.

The exact physiological significance of these
observations remains to be established. According
to Grossman’s criteria,\(^44\) one of the conditions to be
satisfied before an action of a gastrointestinal hormone can be considered ‘physiological’ is the
demonstration that infusion of the hormone to
plasma concentrations comparable with those found
after a meal induces such an action. In dogs\(^45\) as well
as in man\(^10\) no increase in plasma bombesin like
immunoreactivity was detected after a meal. Be-
cause mammalian bombesin like peptides appear to
be confined to nerves of the gastrointestinal tract,\(^6\)\(^7\)
however, such peptide might have a neural rather
than hormonal role.\(^46\)

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