Exocrine secretory responses of the pancreas to insulin and to a meat meal in dogs

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SUMMARY  Studies in five dogs with chronic pancreatic and gastric fistulae have shown that insulin-induced vagal stimulation of the pancreas (gastric fistula open) resulted in protein and bicarbonate outputs very much smaller than those obtained with a 400-g meat meal. However, when the insulin-activated gastric acid secretion was allowed access to the duodenum (gastric fistula closed) peak outputs of both bicarbonate and protein were observed which were closely similar to the response to the meal. These findings suggest that insulin-induced hypoglycaemia results in stimulation of the pancreas within the physiological range when gastric acid is allowed access to the duodenum with consequent release of secretin.

Stimulation of the vagus nerves has been shown to cause a small increase in the volume of exocrine secretion of the pancreas and a considerable increase in output of enzymes (Lobassov, 1910). Cephalic, gastric, and intestinal phases of pancreatic secretion have now been identified (Grossman, 1971), but little is known of the relationships of these phases to the response to a meal. The aim of this paper is to compare the exocrine secretory response of the pancreas during the cephalic phase of pancreatic secretion as induced by insulin hypoglycaemia with that obtained by a meat meal of 400 g in dogs.

Methods

Observations were made on five mongrel dogs weighing between 15 and 22 kg. In each animal the stomach was drained at its most dependent part by means of a Thomas cannula and a chronic pancreatic fistula was formed by the technique described by Preshaw and Grossman (1965). Between experiments in this model pancreatic juice enters the duodenum and the animals do not develop steatorrhoea.

At least six weeks were allowed to elapse after operation before beginning observations. Following an 18-hour fast the animals were placed in slings and both duodenal and gastric fistulae were opened, allowed to drain, and then washed out with 100 ml of 0.9% sodium chloride. Pancreatic juice was collected continuously by gravity drainage and every 30 minutes the volume was measured in millilitres. Total protein concentration (mg/ml) was estimated by Lowry's method (Lowry, Rosebrough, Farr, and Randall, 1951), and protein output (mg/30 min) was calculated from the product of the volume multiplied by concentration.

Bicarbonate concentration in each 30-minute sample was measured by the addition of 1.0 ml of 0.1 NHCl to 0.5 ml of the sample, heating the mixture to boiling for five seconds, and then back titrating the residual acid with 0.1 N Na OH using phenol red as indicator (to pH 6.4-8.6). Bicarbonate output (m-equiv/30 min) was expressed as the product of volume multiplied by concentration.

After two basal collections the exocrine secretory response of the pancreas to insulin (0.2 units/kg iv) was measured for four hours, first with the gastric fistula open (allowing gastric contents to drain freely and thereby preventing access of acid to the duodenum and consequent release of endogenous secretin), and secondly with the gastric fistula closed (allowing acid access to the duodenum with release of secretin). On a separate day, the exocrine response of the pancreas to a 400 g meat meal was measured. On this occasion the gastric fistula remained closed. Throughout the course of the study the dogs were fed on a normal diet of meat and biscuits and maintained their weight. One dog developed an abscess close to the duodenal cannula and this required incision and drainage. Observations were not made on this animal at this time.

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Results

Figure 1 shows the effect of insulin (0·2 units/kg) on the bicarbonate output of the pancreas. Peak bicarbonate output occurred 60 minutes after injection of insulin. With the gastric fistula closed and acid entering the duodenum releasing endogenous secretin, bicarbonate output, as expected, was significantly greater than in experiments with the gastric fistula open ($p < 0.001$). In experiments with the gastric fistula open a significant increase in bicarbonate output after insulin was noted when compared with basal secretion ($p < 0.05$). The pH of fluid obtained from the duodenum via the duodenal cannula was always greater than 6 when the gastric fistula was open during insulin-induced gastric acid secretion.

Figure 2 shows the effect of insulin on protein output from the pancreas. Peak output of protein with the gastric fistula closed occurred 30 minutes after the injection of insulin, and was significantly greater than that obtained in experiments with the gastric fistula open ($p < 0.01$). However, compared to the basal secretion in experiments with the gastric fistula open, a marked increase in output of protein was noted, suggesting vagal stimulation of the pancreas.

Figure 3 shows the comparison of bicarbonate outputs to insulin and to a 400 g meat meal with the gastric fistula closed. It can be seen that the initial increase, and indeed peak responses, to each stimulant were very similar, but that the response to insulin decreased markedly after two and a half hours and was back at almost basal levels by four hours, whereas the response to a 400 g meal decreased more slowly and was still markedly elevated above basal levels four hours after ingestion of the meal.

Figure 4 compares the protein output of the pancreas to insulin hypoglycaemia with that to a meat meal with the gastric fistula closed. It can be seen that the initial rapid rise in protein output was
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Fig. 3 Comparison of bicarbonate outputs after insulin (0.2 units/kg iv) and after a meat meal (400 g): peak outputs were similar but the levels after food were significantly greater throughout the latter half of the experiments.

Fig. 4 Comparison of protein output after insulin and a meat meal (400 g).

gastric fistula open (p < 0.05) but was significantly less than that observed in response to a 400 g meat meal (p < 0.01).

Discussion

While it is possible that some degree of vagal denervation of the pancreas resulted from the construction of our experimental model, it is clear that some vagal innervation persisted, as was indicated by the sharp increase in protein output during insulin-induced hypoglycaemia with the gastric fistula open (Fig. 2).

If protein output to a 400 g meat meal is taken as a 100% response then the mean output of protein to insulin represented 41% (gastric fistula open) and 60% (gastric fistula closed) of the response to the meal. Similarly, if the mean four-hour output of bicarbonate to a 400 g meat meal under the conditions of the experiment is taken as 100%, then the mean four-hour bicarbonate response to insulin represented 10% (gastric fistula open) and 72% (gastric fistula closed) of the meal response. It is clear that when gastric acid is excluded from the duodenum the pancreatic outputs of protein and bicarbonate to vagal stimulation induced by insulin

Fig. 5 shows the mean four-hour bicarbonate output (milli-equivalents) for each of the three groups of experiments performed. It can be seen that the four-hour bicarbonate response to insulin with the gastric fistula closed was significantly greater (p < 0.001) than the bicarbonate output with the gastric fistula open, and significantly less than that observed in response to a 400 g meat meal (p < 0.05).

In Fig. 6 it can be seen that the mean four-hour output of protein to insulin with the gastric fistula closed was significantly greater than that with the
hypoglycaemia are considerably smaller than those occurring after a meat meal. However, in conjunction with endogenous secretin, ie, when the gastric fistula is closed, peak outputs of protein and bicarbonate were observed which did not differ significantly from those seen during the response to the meal. These findings suggest that insulin-induced hypoglycaemia results in stimulation of the pancreas within the physiological range when gastric acid is allowed access to the duodenum.

We further speculate that the rapid fall of both protein and bicarbonate outputs in experiments with the gastric fistula closed during insulin hypoglycaemia is probably due to a diminished release of secretin consequent upon the decrease in acid entering the duodenum from the stomach.

In the experiments with insulin, pancreatic protein output increased when the gastric fistula was closed compared to those in which it was open, suggesting that the presence of acid in the duodenum augmented the vagally stimulated protein output, possibly by release of pancreozymin. The stimulatory effect of acid alone (Preshaw, Cook, and Grossman, 1966) and of acid plus insulin hypoglycaemia noted by others (Eisenberg and Orahodd, 1971) on the pancreatic protein output is in keeping with such a mechanism.

Since both insulin and a meat meal are known to cause an increase in release of antral gastrin and of plasma gastrin concentration (Eisenberg, Emâs, and Grossman, 1966; Ganguli and Elder, 1971; Ganguli, 1970), the effects of gastrin in these experiments must be considered. Since gastrin is known to cause an increase in protein output from the pancreas (Emâs, Billings, and Grossman, 1968) it is likely that after both insulin and the 400 g meat meal, a part of the output of protein from the pancreas could be due to the stimulatory effects of antral gastrin directly on the pancreas. Direct evidence on this point from radioimmunoassay of plasma gastrin concentration during such experiments would be of great interest.

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