Enhancing effect of partial gastrectomy on pancreatic carcinogenesis

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Summary The controversial issue of enhanced pancreatic carcinogenesis following partial gastrectomy has been explored in male Wistar rats (n = 40) weighing 250–300 g. Animals were randomised to receive either 60% distal gastrectomy with Roux-en-Y reconstruction or gastrectomy and resuture (control). Immediately after operation each group was further divided into two subgroups, receiving i.p. injections of either saline or azaserine (30 mg kg⁻¹ wk⁻¹ for 3 weeks). At 15 months blood was obtained at 0, 5, 15 and 30 min after a fatty meal for cholecystokinin (CCK) assay: rats were then killed. Pancreatic wet weight was measured, and histological sections were examined for atypical acinar cell foci (AACF), the putative precursor lesion of carcinoma. There were no significant differences in body weight or pancreatic weight between controls and rats with gastrectomy. Only azaserine-treated rats had acidophilic AACF. Partial gastrectomy substantially increased the number of acidophilic AACF per pancreas (median 26.05 vs 2.09; P < 0.005), with a 9-fold increase in their volume (P < 0.005). Basal and postprandial plasma CCK concentrations were higher after gastrectomy than in controls (P < 0.05). Partial gastrectomy has an enhancing effect on azaserine-induced pancreatic carcinogenesis, probably by means of increased CCK release.

Since carcinoma of the pancreas is so difficult to cure and its aetiology remains obscure, it is important to investigate potential risk factors such as partial gastrectomy. Several reports have indicated an increased incidence of pancreatic cancer in patients undergoing gastric resection (Ross et al., 1982; Mack et al., 1986; Caygill et al., 1987; Offerhaus et al., 1987; Mills et al., 1988; Tersmette et al., 1990), but other work is contradictory (Maringhini et al., 1987; Tomaszewska & Stachura, 1988; Vecchia et al., 1990). The former popularity of partial gastrectomy for treating peptic ulcer disease, often in young patients, means that there are many patients alive today who could be at risk of cancer of the pancreas as well as cancer in the gastric stump (Schrumpf et al., 1977).

In the alimentary canal, carcinogenesis can be enhanced by luminal factors acting directly on the mucosa to produce hyperplasia (Rainey et al., 1984; Houghton et al., 1987), but in the pancreas any such influence seems to be exerted indirectly through humoral and neural mechanisms. Cholecystokinin (CCK) promotes pancreatic carcinogenesis in the rat-azaserine model, in which the population of atypical acinar cell foci (AACF) of acidophilic type reflects the ultimate number of malignant tumours. Long-term administration of exogenous CCK increases the number of these preneoplastic AACF (Douglas et al., 1989a), and the CCK antagonist CR-1409 blocks this effect (Douglas et al., 1989b). The promoting effect of pancreaticobiliary diversion on experimental pancreatic carcinogenesis may also be mediated through a sustained increase in circulating CCK (Stewart et al., 1991; Watanapa et al., 1991). Although partial gastrectomy does not increase fasting levels of CCK in man or the rat, the CCK response to ingested fat is markedly greater in both species (Hopman et al., 1984; Inoue et al., 1987; Malfertheiner et al., 1987).

We have tested the hypothesis that partial gastrectomy enhances experimental pancreatic carcinogenesis, using quantitative estimation of AACF to show early malignant change and measuring CCK secretion to determine its intermediary role. We avoided any independent effect of duodenogastric reflux, which may itself enhance pancreatic carcinogenesis (unpublished data), by using Roux-en-Y reconstruction after partial gastrectomy rather than a Polya procedure.

Methods

Experimental design

Male Wistar rats (n = 40) weighing 250–300 g were housed in groups of five in animal quarters with a 12 h day night cycle. Standard pelleted rat food (Patterson and the Christopher Hill Group, Porton – diet PRD) and water were freely available. After 1 week of acclimatisation, animals were randomised to receive either 60% distal gastrectomy or gastrectomy and resuture (controls). Immediately after the operation, half the animals in each group were further randomised to receive saline or azaserine (see below). Food was reintroduced 12 h postoperatively. After overnight fasting, all rats were killed at 15 months after operation. Immediately before death rats were anaesthetised and a catheter was inserted into the inferior vena cava. Blood samples (2 ml) for CCK assay were obtained at 0, 5, 15 and 30 min after a fatty test meal comprising 3 ml soya bean oil, 1 ml glucose solution (40%), 2 ml water and 0.4 g protein (Maxipro, Scientific Hospital Supplies, Liverpool, UK). The test meal was infused through a catheter placed just beyond the pylorus (in controls) or the gastric anastomosis (in rats with gastrectomy). The position of this catheter was chosen to avoid any effect of altered gastric emptying. Similar amounts of normal saline were infused intravenously to maintain the animals until the end of the test–meal study. After death, the pancreas was excised and trimmed free of adherent fat and lymph nodes. The wet weight of each gland was recorded before fixation in 10% buffered neutral formalin. Before immersion in the fixative solution, each pancreas was spread out on a piece of porous paper to ensure maximal transsectional sectioning.

Operations (Figure 1)

Partial gastrectomy was performed by removal of the distal 60% of the stomach. Gastrojejunostomy was carried out in a Roux-en-Y fashion with a 15 cm Roux loop of jejunum (roughly corresponding to the 45 cm loop usually created in man). Control rats received a gastrostomy, consisting of a 1 cm incision in the greater curvature of the stomach, which was immediately resutured. Operations were carried out

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under light ether anaesthesia, and a continuous 6.0 silk suture was used for anastomoses.

Carcinogen
Azaserine (Sigma Chemical Company, UK) was dissolved in 0.9% NaCl to give a 2.5% solution and was administered by weekly i.p. injection into each rat, starting immediately after operation and continuing for the next 2 weeks. The dosage regime was 30 mg kg⁻¹ wk⁻¹, giving a total dose of 90 mg kg⁻¹. Controls received 1.2 ml kg⁻¹ of 0.9% NaCl by weekly i.p. injection.

CCK assay
Plasma CCK peptides were extracted from venous blood samples with C18 'SepPak' cartridges (Waters, Harrow, UK) (Eysellin et al., 1987), and eluates were dried by centrifugal evaporation (Savant, Famingdale, NY, USA).

CCK was measured by a specific radioimmunoassay based on antisera A₁, raised by immunising a rabbit with natural porcine CCK-33 (donated by Professors V. Mutt and S.R. Bloom). Antiserum A₁ (1:60 000) was incubated at 4°C for 3 days with standard CCK-8 or with plasma samples plus CCK-8 tracer-labelled with ¹²⁵I (1 000 c.p.m., Amer sham, UK) in 0.05 mol l⁻¹ sodium phosphate buffer (pH 7.4) containing 0.25% gelatin and 0.01 mol l⁻¹ EDTA. Free and bound tracer were separated by the addition of 6% (weight volume) charcoal (Norit PN5, BDH, Poole, UK) with 0.6% (weight volume) dextran. The concentrations of pure peptides that produced half-maximal inhibition of binding of tracer to A₁ were 2.0 pmol l⁻¹ for CCK-8, 2.4 pmol l⁻¹ for CCK-33, and 2.2 nmol l⁻¹ for gastrin 17. The coefficient of variation within assays was 8.2% and between assays 12.8%. The sensitivity of the assay (defined as minimal amount of CCK-8 that could be distinguished from zero with 95% confidence) was 0.2 pmol, and the recovery of CCK-8 and CCK-33 through the SepPak and assay procedure was 79%.

Quantitative estimation of AACF
The pancreas was spread out very thinly and then sectioned horizontally so that the whole gland could be examined. Histological sections (5 μm) of the whole pancreas were stained with haematoxylin and eosin, coded and scrutinised ‘blind’, i.e., the observers did not know what treatment each animal had received. The atypical acinar cell foci (AACF) were readily identified and classified as acidophilic or basophilic according to established criteria (Rao et al., 1982).

The total area of exocrine pancreatic tissue was measured directly in a single histological section from each pancreas by means of a VIDS III video image analyser (Analytical Measuring Systems, Cambridge). The same instrument was used to count acidophilic and basophilic AACF and to measure their transsectional area. Data were processed numerically by the Volugen computer package (InfoResearch Int., Bristol), using an algorithm based on that of Campbell et al. (Campbell et al., 1982) and modified by Pugh et al. (Pugh et al., 1983). Details of this analysis have been already described in our previous studies (Stewart et al., 1991).

Statistical analysis
Student's t-test for unpaired data was used for the group analysis of plasma CCK concentrations. The levels were expressed as means (SEM). Median values and ranges were quoted for body weight, pancreatic weight and quantitative estimation of AACF. Statistical analysis of these parameters were performed using Kruskal-Wallis one-way analysis of variance and the Mann-Whitney U-test.

Results
Mortality, body weight and pancreatic weight (Table I)
There were five premature deaths from anastomotic leakage with granuloma formation and intestinal obstruction (two in gastrectomy-azaserine rats and one in each other group). Yields of healthy survivors were as follows: control-saline, nine; control-azaserine, nine; gastrectomy-saline, nine and gastrectomy-azaserine, eight.

There were no differences in body weight between the four groups. Although both absolute and relative pancreatic weights of rats with gastrectomy were greater than those of controls, the differences did not reach a significant level. Macroscopic examination of the pancreas at autopsy revealed numerous small white elevated nodules on the surface of the glands of azaserine-treated animals, particularly those with gastric resection.

Plasma CCK (Figure 2)
Partial gastrectomy increased basal circulating CCK concentrations by 46%. Following the test meal, the plasma CCK response at 5 min in rats with gastrectomy was greater than in controls (52% vs 41% increments over basal). At 15 and 30 min, plasma CCK levels remained 19% greater than those of controls, but these differences did not show statistical significance.

Quantitative analysis of AACF
No pancreatic carcinomas were found. Acidophilic AACF, the putative precancerous lesions, were only seen in azaserine-treated rats (Table II), whereas a few basophilic foci appeared in controls as well (Table III). Among azaserine-treated groups, the observed transectional data (foci per cm²) revealed a marked increase in incidence of acidophilic lesions following partial gastrectomy compared with controls (2.29 vs 0.24). Quantitative stereological analysis of tissue sections confirmed the dramatic response of the pancreas to gastric resection with respect to acidophilic foci. Thus the number of lesions per cm² was substantially greater (15.40 vs 1.41), as was the total number of lesions per pancreas (26.05 vs 2.09). The median diameter of each lesion was increased by 69% and the volume by a factor of nine. Moreover, partial gastrectomy enhanced the percentage of the pancreatic volume occupied by acidophilic foci from 0.09% (control) to 3.38%. With regard to basophilic AACF, partial gastric resection increased the population of the lesions only with azaserine treatment, but the gastrectomised animals had fewer foci than the corresponding controls receiving saline.

Discussion
Partial gastrectomy clearly promoted experimental pancreatic carcinogenesis, as shown by the very considerable increase in the number and size of acidophilic AACF. Acidophilic
AACF are well established as the precursors of cancer in this model (Rao et al., 1982; Roebuck et al., 1984), and they were only found in rats receiving azaserine. Acidophilic AACF show considerable growth potential with a mitotic index (2.75) which greatly exceeds that of basophilic foci (0.125) or normal pancreas (zero) (Scarpelli et al., 1984). Their increased number after gastrectomy mirrored our subjective assessment that there were many more macroscopic nodules on the surface of the pancreas in these rats. We encountered fewer AACF in all groups compared with Longnecker’s reports (Longnecker et al., 1977; Roebuck et al., 1985) and our own previous experience (Stewart et al., 1991), probably because of the relatively large size of rat chosen to facilitate the gastric operation. Likewise, no actual carcinomas were found in the pancreas, although Longnecker and Curphey reported a few of these lesions at 1 year when much younger rats were given azaserine (Longnecker & Curphey, 1975). Although quantitative analysis showed that the population of basophilic foci was also increased in both number and size after partial gastrectomy, the relevance of this finding is doubtful since most modulators of the postinitiation phase of pancreatic carcinogenesis have little effect on basophilic foci (Roebuck et al., 1982; Roebuck et al., 1985).

Partial gastrectomy alters circulating levels of several gut peptides, notably gastrin, pancreatic polypeptide and CCK (Inoue et al., 1987; Malfertheiner et al., 1987; Rieu et al., 1990). Exogenous gastrin stimulates pancreatic growth (Johnson, 1976), whereas pancreatic polypeptide inhibits pancreatic secretion (Taylor et al., 1979); the effect of reducing their circulating levels has not been established in pancreatic carcinogenesis. The increase in basal CCK concentrations and the increased CCK response to a fatty test meal strongly implicate this peptide as an intermediary in the promoting effect of partial gastrectomy. The enhanced postprandial

![Figure 2](image-url)
CCK response is in line with other reports both in man and the rat (Homan et al., 1984; Inoue et al., 1987; Malfertheiner et al., 1987) but unlike other authors we also found a higher basal level. Previous studies were undertaken either 2 weeks after partial gastrectomy in rats or 1 month after partial gastrectomy in man; our data suggest that hypercholesterolaemia persists for up to 15 months and may even increase with time.

Since direct infusion of the fatty meal into the small bowel circumvented any variability in gastric emptying, the increased CCK release after partial gastrectomy may be due to an increased responsiveness of CCK-secreting cells. Diversion of peptidergic secretions from the jejunal limb of a Roux-en-Y anastomosis can cause mucosal hyperplasia (Miazzia et al., 1982), and this hyperplastic response might well involve the enteroendocrine cells and lead to increased cholecystokinin production. Lower concentrations of intraluminal protease have been shown in patients with subtotal gastric resection in the early phase after a fatty meal (MacGregor et al., 1977). Similar protease reduction in the gut might also contribute to an increased CCK response, since low levels of intraluminal trypsin are known to stimulate CCK release (Louve et al., 1986; Calam et al., 1987). Although CCK stimulates pancreatic growth, the fact that there is a loss of the normal pancreatic response to several tropic hormones (including CCK) with advancing age (Greenberg et al., 1986; Bouchier et al., 1991) might explain the non-significant increase in pancreatic weight 15 months after partial gastrectomy.

Data from an experimental rat model can only be of tentative relevance to man. Although our rats received a carcinogen, patients with previous partial gastrectomy have increased levels of nitrates and n-nitroso compounds in gastric juice, and these substances can act as pancreatic carcinogens (Schlag et al., 1980). Nitrosamines could be absorbed and subsequently secreted into the pancreatic juice or might reflux from the duodenum into the pancreatic duct, thereby inducing pancreatic cancer. The combination of increased post-prandial CCK release and greater exposure to pancreatic carcinogens might explain the increase in pancreatic cancer risk after gastrectomy. Our unpublished data showing only a few acidophilic AACC 6 months after a similar partial gastrectomy in rats underline the importance of a long-term experiment for a clear-cut effect to emerge. They could also explain why an increased risk of pancreatic cancer in man only appears to reach statistical significance 20 years after gastric resection (Caygill et al., 1987; Tersmette et al., 1990).

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