Role of cholecystokinin in pancreatic adaptation to massive enterectomy

P Watanapa, M Egan, P H Deprez, J Calam, C E Sarraf, M R Alison, R C N Williamson

Abstract
Since pancreatic adaptation to massive proximal small bowel resection (PSBR) may be modulated through cholecystokinin (CCK) secretion, we tested the effect of the CCK antagonist CR-1409 on this response. Male Wistar rats (n=72) weighing 220–225 g were randomised to receive either PSBR or transection/resuture followed by saline or CR-1409 (12 mg/kg daily subcutaneously). Rats were killed one, two, and three weeks postoperatively, at which time blood was obtained for CCK assay and the pancreas was assessed for proliferative activity by three parameters: nucleic acid and protein content, bromodeoxyuridine (BrdUrd) labelling index, and proliferating cell nuclear antigen (PCNA) expression. PSBR increased plasma CCK concentration by 83–102% at 1–3 weeks, irrespective of CR-1409 administration. Total pancreatic DNA content per 100 g body weight increased by 34% at two weeks (p<0.05) and by 82% at three weeks (p<0.05), while RNA content increased by 60% and 175% (p<0.001) and protein content by 20% and 57% (p<0.05). PSBR increased the BrdUrd labelling index and the percentage of PCNA immunoreactive cells. CR-1409 completely abolished this proliferative response and also prevented the rise in nucleic acid and protein contents. Apart from growth stimulation, PSBR also enhanced pancreatic exocrine function, as shown by ultrastructural evidence of an appreciable decrease in zymogen granules; CR-1409 also inhibited this functional effect of hypercholecystokininæmia. The results confirm the trophic role of CCK after PSBR, and CR-1409 prevents this pancreatic adaptation.

The compensatory response to extensive proximal small bowel resection (PSBR) includes growth of the remaining small intestine (duodenum and ileum) and of the gastric parietal cells, causing acid hypersecretion. The pancreas shares in this adaptive response. Pancreatic hyperplasia can be detected as early as one week after 90% PSBR and persists for at least six months. Fifty and 75% PSBR also produce early and appreciable increases in basal pancreatic secretion of water and bicarbonate, but unlike the structural adaptation, these functional changes are transient.

Intestinal adaptation to PSBR is controlled by both luminal and humoral factors, but since the pancreas lies outside the alimentary tract, only systemic factors seem likely to mediate its adaptive response. There are several candidate gastrointestinal hormones for the role of 'pancreatotropin'. Although long term administration of pentagastrin can stimulate pancreatic growth and enterectomy causes hypergastrinæmia, studies in rats with antrectomy or vagotomy have excluded its involvement in the response to PSBR. Circulating levels of insulin, pancreatic glucagon, and neurotensin are unchanged after small bowel resection, whereas enteroglucagon is increased and concentrations correlate with indices of mucosal mass and cell turnover in the pancreas. Cholecystokinin is also increased 12 days after PSBR, and persistent hypercholecystokinæmia can certainly cause pancreatic hyperplasia. The present study was designed to test whether massive (90%) PSBR would remove sufficient enteroendocrine cells to prevent the hypercholecystokinæmia of lesser enterectomies and, if not, whether the CCK receptor antagonist CR-1409 would blunt the adaptive response of the pancreas. Daily administration of CR-1409 can inhibit CCK induced pancreatic growth after pancreatectobiliary diversion.

Pancreatic mass can be assessed by measuring wet weight, nucleic acid contents, and protein contents. We have attempted to measure cell proliferation, which should precede any increase in organ mass, by adopting two immunohisto logical methods to the pancreas: proliferating exocrine cells were labelled either with 5-bromo-2'-deoxyuridine (BrdUrd) or with a monoclonal antibody (PC10) against proliferating cell nuclear antigen (PCNA).

Methods

EXPERIMENTAL DESIGN
Male Wistar rats (n=72) weighing 200–225 g were housed in groups of five in animal quarters with a 12 hour day/night cycle. Standard pelleted rat food (Paterson and the Christopher Hill Group, Porton-diet PRD) and water were freely available. After 48 hours' acclimatisation, animals were randomised to receive either 90% proximal small bowel resection (as measured from the ligament of Treitz) or jejunal transection and resuture (controls). Operations were carried out under light ether anaesthesia. A continuous 6/0 silk suture was used for intestinal anastomoses.

Immediately after the operation, half the animals in each group were further randomised to receive either CR-1409 (12 mg/kg/day) or saline (3 ml/kg) by daily subcutaneous injection. CR-1409 was dissolved in distilled water and brought to pH 9 by 0.01 N NaOH to give a 0.4% solution. Food was reintroduced 12 hours postoperatively. After overnight fasting, representatives of each of the four groups were killed at 1, 2,
and 3 weeks after operation. Blood was obtained by direct cardiac puncture for cholecytokinin assay. The pancreas was excised, trimmed free of adherent fat and lymph nodes, and weighed. Small samples were removed for protein and nucleic acid content measurement, BrdUrd labelling index, and PCNA immunoreactive cell count.

NUCLEIC ACID AND PROTEIN CONTENTS
Pancreatic tissue was homogenised in phosphate buffered saline (PBS) (136.9 mmol/l NaCl, 2.7 mmol/l KCl, 8.1 mmol/l Na3HPO4, 1.5 mmol/l KH2PO4, 0.9 mmol/l CaCl2, 0.5 mmol/l MgCl2-6H2O, adjusted to pH 7.5 with NaOH), and the concentration of the homogenate was adjusted to 1 mg tissue per ml. Pancreatic contents of DNA and RNA were assayed by the fluorimetric method of Boer et al.11 but using an excitation wavelength of 536 nm and an emission wavelength of 602 nm.11 Protein content was determined by the method of Lowry et al.11 Since body weight has a profound effect on pancreatic weight,13 the results were expressed as mg per 100 g body weight to correct for this variable.

BrdUrd Labelling Index
BrdUrd is a thymidine analogue that is readily and specifically incorporated into DNA during the S phase of the cell cycle. Rats were given an intraperitoneal injection of BrdUrd (50 mg/kg body weight) one hour before sacrifice. A small sample of the excised pancreas was placed in Carnoy's solution (absolute alcohol:chloroform:glacial acetic acid=6:3:1 by volume) for two to four hours and transferred to 75% ethanol. Tissue was processed in chloroform, wax embedded, and 4 μm sections were cut. Sections were dewaxed and dehydrated in 100% alcohol. Endogenous peroxidase activity was removed by subsequent immersion in methanol/hydrogen peroxide (98.4%-1.6%). Dissociation of histones was achieved by using 1 mol/l HCl, which also partially denatured DNA. Each section was treated with a drop of primary antibody (mouse anti-human BrdUrd, Dako, High Wycombe; 1:20 dilution in PBS). After overnight incubation at 4°C, the section was incubated with a peroxidase conjugated second antibody (rabbit anti-mouse IgG, 1:50 dilution with PBS) for one hour at room temperature. The colour reagent diaminobenzidine plus H2O2 was used for staining, and each section was counterstained with haematoxylin. The number of labelled cells in each section was estimated from a count of 2000 exocrine pancreatic acinar cells, and the result was expressed as a percentage— the labelling index.

PCNA Labelling Index
PCNA is an acidic nuclear protein, expression of which is directly correlated with rates of cell proliferation and DNA synthesis. The monoclonal antibody PC10 will 'recognise' PCNA in conventionally fixed and processed histological material. The tissue sample of the excised pancreas was placed in buffered formulin for two to four hours and transferred to 75% ethanol. Tissue was processed in chloroform and embedded in wax before 4 μm sections were cut. Sections were dewaxed and taken down through graded alcohols; endogenous peroxidase activity was blocked by incubating the sections in 3% hydrogen peroxide and methanol for one hour. After washing in PBS, pH 7.4, each section was treated with a drop of primary antibody (1:20 dilution in PBS). After overnight incubation at 4°C, the sections were washed in PBS, 0-1% bovine serum albumin (BSA), and Tris-BSA. The second layer antibody, biotinylated goat anti-mouse IgG (Vector Laboratories, Burlingame, CA, USA) was applied at a dilution of 1:50 and incubated for two hours at room temperature. After washing in PBS, streptavidin-peroxidase (Jackson Immunonuclear Laboratories, Westgrove, PA, USA) was applied to the sections at a 1:5000 dilution in PBS with 1% BSA for 30 minutes at room temperature. Diamino-benzidine-hydrogen peroxide was employed as a chromogen, and a light haematoxylin counterstain was used. The PCNA labelling index was estimated from a count of 2000 exocrine acinar cells and expressed as a percentage.

PLASMA CCK ASSAY
Plasma CCK peptides were extracted from cardiac blood samples with C18 'SepPak' cartridges (Water, Harrow, UK),16 and eluates were dried by centrifugal evaporation (Savant, Farmingdale, NY, USA). CCK was measured by a specific radioimmunoassay based on antisera A3, raised by immunising a rabbit with a natural porcine CCK-33 (donated by Professors V Mutt and S R Bloom). Antiserum A3 (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 iodine-125 (1000 cpm, Amersham, UK) in 0-05 mol/l sodium phosphate buffer (pH 7.4) with 0-25% gelatin and 0-01% mol/l ethylenediamine tetra-acetic acid (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
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PEPTIDES (Fig 1)

Peptides that produced half maximal inhibition of binding of tracer to $A_2$ were 2-0 pmol/l for CCK-8, 2-4 pmol/l for CCK-33, and 2-2 nmol/l 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%.

ULTRASTRUCTURAL STUDIES

Samples of all categories of pancreas, not exceeding 1 mm$^3$ in volume, were fixed in 2% glutaraldehyde for two hours. After washing in phosphate buffer, tissues were osmicated and dehydrated in acidified DMP before routine embedding in Taab resin. Sections (1 μm) were cut and stained with toluidine blue for observation by light microscopy, followed by ultrathin sections of approximately 100 nm, collected on nickel grids and stained with uranyl acetate and lead citrate, for observation on a Philips CM-10 electron microscope.

STATISTICAL ANALYSIS

Student’s $t$ test for unpaired data was used for the group analysis. Nucleic acid and protein contents and plasma CCK values were expressed as mean (SEM), whereas those of BrdUrd labelling index and PCNA labelling index were expressed as medians. The Spearman correlation coefficient was used to assess the correlation between the results of the BrdUrd and PCNA labelling indices.

RESULTS

MORTALITY AND BODY WEIGHT (Fig 1)

There were two deaths from anastomotic leakage (one control and one PSBR rat). Rats undergoing massive small bowel resection weighed less than controls both at one and two weeks whether or not they received CR-1409 (8-14% less at one week, $p<0-001; 10-24% less at two weeks, $p<0-001$). By three weeks, resection rats weighed 15% less than controls, but there was no difference in body weight between the resection and transection animals receiving CR-1409.

PANCREATIC WEIGHT (Fig 2)

Proximal small bowel resection had a clearcut effect on both absolute and relative pancreatic weight. Absolute pancreatic weight was 4-9-9% less than controls at one week but was 18% greater than controls at two weeks and 27% greater at three weeks. Likewise, enterectomy caused a 32% increase in relative pancreatic weight over controls at two weeks and a 50% increase at three weeks. CR-1409 completely abolished this pancreatic growth, so that differences in absolute and relative pancreatic weights between resection and transection animals were no longer significant.
Plasma cholecystokinin concentrations (pmol/l) in rats with proximal small bowel resection (PSBR) or transection (control).

<table>
<thead>
<tr>
<th>Period post operation (weeks)</th>
<th>Group</th>
<th>Control - saline</th>
<th>PSBR - saline</th>
<th>Control - CR-1409</th>
<th>PSBR - CR-1409</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2.69 (0.29)</td>
<td>4.92 (0.66)*</td>
<td>2.90 (0.66)</td>
<td>5.52 (0.54)**</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>2.91 (0.46)</td>
<td>5.91 (1.13)*</td>
<td>3.74 (0.66)</td>
<td>7.45 (1.03)*</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>2.77 (0.30)</td>
<td>5.43 (0.78)*</td>
<td>2.76 (0.47)</td>
<td>7.45 (1.99)*</td>
<td></td>
</tr>
</tbody>
</table>

Significance *p<0.05, **p<0.005. Values mean (SEM).

**PLASMA CHOLECYSTOKININ** (Table)

Small bowel resection caused a sustained increase in circulating CCK concentrations – by 83% at one week, by 102% at two weeks, and by 96% at three weeks. Administration of CR-1409 had no appreciable effect on plasma hormone concentrations.

**NUCLEIC ACID AND PROTEIN CONTENTS** (Fig 3)

Pancreatic nucleic acid and protein contents increased after proximal small bowel resection, but the increase did not attain statistical significance until two weeks, when DNA was 34% higher, RNA 60% higher, and protein 20% higher than in controls. These changes were more obvious by three weeks, when increments were 82%, 178%, and 57%. With the exception of the DNA contents at two weeks, the CCK antagonist completely inhibited growth stimulation produced by PSBR.

**BRDURD LABELLING INDEX** (Fig 4)

One week after PSBR, the labelling index was already higher than in controls (0.35 vs 0.05%). The increase in proliferative response became increasingly obvious at two weeks (1.43 vs 0.20%) and was still present at three weeks (0.88 vs 0.35%). CR-1409 abolished the effect of proximal enterectomy, so that there was no difference from control values at any time point. The antagonist had no effect in rats with small bowel transection.

**PCNA LABELLING INDEX** (Fig 5)

Broadly similar results were obtained for the cellular kinetic data derived by counting PCNA immunoreactive cells. PSBR caused an increase in the proliferation rate at one week (0.70 vs 0%). The differences were more marked both at two weeks (4.13 vs 0%) and at three weeks (2.10 vs 0.55). Again, CR-1409 largely abolished the growth stimulation produced by bowel resection but had no consistent effect within the control groups. There was a correlation between the BrdUrd labelling index and the PCNA labelling index, with a Spearman correlation coefficient of 0.68.

**ULTRASTRUCTURAL STUDIES**

Samples of pancreas from rats with transection and saline injection showed no differences from normal pancreas. Acinar cells were typically pyramidal in shape with closely packed rough endoplasmic reticulum, numerous mitochondria and abundant apical zymogen granules (Fig 6). By contrast, samples from animals with PSBR and saline showed incipient vacuole formation in the rough endoplasmic reticulum of many acinar cells, associated with marked degranulation (Fig 7). No degranulation was noted in either transection or resection animals that had been treated with CR-1409, but incipient vacuolation of the rough endoplasmic reticulum was a feature of all CR-1409 treated tissues as well as those of the PSBR group (Fig 8).

**Discussion**

In line with other work, the present data show that an extensive proximal small bowel resection stimulates pancreatic growth, as characterised by increased wet weight and increased DNA, RNA, and protein contents. The pattern of this adaptive growth is mainly one of hyperplasia, since the RNA/DNA ratio was little changed. Increased organ mass was accompanied by increases in two markers of cell proliferation, the BrdUrd and PCNA labelling indices.

Our results firmly implicate CCK as a key intermediary in the adaptive response of the pancreas to massive proximal small bowel resection. Not only were CCK levels increased two to
explanations: (1) loss of a negative feedback mechanism on CCK release by a depletion of the anticholecystokinin peptide normally released from the ileum, since our procedure removed more than 80% of the ileum; (2) hyperplasia of the endocrine cells within the remaining intestine, notably those in the duodenum (modest duodenal hyperplasia follows jejunectomy); and (3) impaired elimination of CCK after removal of jejunum.

Immunohistological methods of assessing cell proliferation have particular advantages over other techniques because of the maintenance of cellular and tissue architecture, the relative simplicity of the methodology, and the rapidity of results. Pancreatic acinar cell kinetics can be estimated by using the methods of BrdUrd labelling index and metaphase arrest after vincristine administration. Recently, PCNA, an auxiliary protein of DNA polymerase-delta, has been shown to play a critical role in the initiation of cell proliferation. The distribution of PCNA in the cell cycle makes it a useful marker for proliferating cells: increasing through G1, peaking at the G1/S phase interface, decreasing through G2, and reaching low levels (which are virtually undetectable by immunocytochemical methods) in M phase and quiescent (G0) cells. PCNA can be demonstrated by using monoclonal antibodies, usually requiring cryostat sections or specially prepared histological material. PC10 is a new antibody that has the advantage of recognizing PCNA in conventionally fixed and processed histological material. As expected, there were more PCNA staining cells than BrdUrd-incorporating cells giving the PCNA technique a potential advantage in organs like the pancreas where there is low level of proliferative activity.

Our preliminary studies indicate that raised circulating levels of CCK cause degranulation in pancreatic acinar cells. Animals that underwent proximal enterectomy alone demonstrated acinar cell degranulation typical of the response to raised CCK levels, but these changes were not seen after treatment with the antagonist. Thus, CR-1409 not only abrogates the tropic effect of CCK on pancreatic growth but also inhibits functional stimulation of the exocrine cell. The distension of the rough endoplasmic reticulum and the cell vacuolation caused by the antagonist probably reflect a direct effect on pancreatic acinar cells.

We thank the Royal Postgraduate Medical School and the Hammersmith and Queen Charlotte’s Special Health Authority for supporting this research and Dr W K Man and Mr S K Li for the nucleic acid and protein assay. The CCK antagonist CR-1409 was kindly provided by Dr L Rovati of Rotta Research Laboratories (Milan, Italy), and the PC10 antibody was donated by Professor David Lane, CRC Cell Transformation Research Group, Department of Biochemistry, University of Dundee.

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