Liver, biliary, and pancreas

Acid resistant lipase as replacement therapy in chronic pancreatic exocrine insufficiency: a study in dogs

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SUMMARY Conventional treatment of pancreatic steatorrhoea in man has been unsatisfactory because 90% of the lipase content of therapy is inactivated by acid in the stomach and large doses of replacement treatment are needed to provide adequate supplementation. An acid stable agent (fungal lipase) was investigated in the treatment of pancreatic deficiency steatorrhoea in 11 pancreatetomised dogs maintained on a fixed dietary intake of fat and treated with pancreatin or fungal lipase. Ten grams (60 000 U lipase) of pancreatin was compared with 400mg (4800 U lipase) of fungal lipase administered with each meal against a no treatment group. There was no significant difference in stool bulk and faecal fat excretion between pancreatin and lipase treated animals. Both groups showed a significant reduction in stool bulk and faecal excretion when compared with the no treatment group (p<0.01). A markedly diminished treatment volume, in the form of fungal lipase, is as effective in controlling steatorrhoea as pancreatin and may prove to be a potentially valuable therapy for patients with pancreatic insufficiency.

The most common causes of pancreatic exocrine insufficiency are cystic fibrosis and chronic pancreatitis resulting in two major problems: poor nutrition and considerable social embarrassment. Steatorrhoea is mainly a consequence of failure of digestion and aqueous solubilisation of lipid leading to its malabsorption.

The mainstay of treatment in pancreatic steatorrhoea has been replacement of endogenous pancreatic enzymes by supplements from hog pancreas (pancreatin – Paynes and Byrne). This has often proved unsatisfactory in the long term as many patients require 20 or 30 tablets per meal and still experience the nutritional and social consequences of steatorrhoea. As much as 30 000 U lipase are needed with each meal to abolish steatorrhoea.1 Most commercial preparations contain about 5000 U of lipase per tablet and do not approach the replacement dosage. Unfortunately, 90% of the lipase content of pancreatin is inactivated by gastric acid2 and so larger numbers of tablets are required to provide adequate supplementation.

Pancreatin is available in powder, capsules or tablets. Powder, sprinkled on meals, is found to be unpalatable and is poorly tolerated except in very young children. Capsules (gelatin shells containing pancreatin) dissolve in the stomach releasing the powder. Enteric-coating of pancreatin, formulated to dissolve only above a pH of 6° (Nutrizym and Pancrex V Forte), is intended to protect pancreatin against intragastric inactivation. This preparation is seldom completely effective as the capsules can remain intact in the stomach until the meal has passed and when the capsules do enter the duodenum the coating may not dissolve because of duodenal hyperacidity3 – most patients with pancreatic insufficiency have reduced bicarbonate secretion. Preparations including the microencapsulated pancreatin, designed to release the enzyme in the jejunum, may be ineffective if jejunal acidification is significant. Adjunctive treatment with H2 receptor antagonists can lessen intragastric inactivation of pancreatin and therefore increases lipid digestion and fat solubilisation in the duodenum.4 Cimetidine (Smith Kline and French) and pancreatin together are more effective in abolishing steatorrhoea and reducing faecal fat excretion5 but in many patients this is unsuccessful.

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Expense, number of tablets required and associated poor patient compliance continue to encourage the search for more satisfactory forms of replacement therapy.

Digestion of fats by pancreatic lipase occurs mainly in the duodenum. Lingual lipase which is acid resistant contributes to this process.7 The advantages of acid resistant lipase are obvious but lingual lipase from animal sources is not available. Lipase AP (Paynes and Byrne) is a novel lipolytic enzyme preparation, manufactured by a fermentation process of aspergillus, which is water soluble and stable at a wide range of pH from 2–10. It is heat stable and non-toxic. The preparation has 12000 U lipase/g weight and its potential therapeutic advantage in reducing exocrine replacement therapy merits evaluation. This study was designed, therefore, to compare the effectiveness of conventional pancreatic enzyme therapy with acid stable fungal lipase (lipase AP) against no treatment in dogs with total pancreatic exocrine insufficiency.

Methods

DOGS

Eleven dogs weighing 15–21 kg were used in the study. All animals underwent total pancreatectomy producing pancreatic exocrine insufficiency. All animals received intrasplenic islet autografts prepared by collagenase digestion of pancreatic tissue to preserve endocrine function.a These animals developed functioning endocrine grafts providing fasting euglycaemia within one month of pancreatectomy. Animals studied preoperatively acted as normal controls (group A). Pancreatectomised animals were assessed on no replacement therapy (group B), on fungal lipase (group C) and on pancreatin (group D). The animals were maintained on a fixed dietary intake which included 46 g fat per day. The study was designed in a randomised crossover fashion incorporating a different treatment regimen for three week periods. Animals were weighed before and after each period and three day collections of faeces were made and analysed for stool volume and fat content at the completion of each treatment. Animals on less than 10 g pancreatin had previously been observed to develop steatorrhoea, so 10 g (60000 U lipase) of pancreatin was compared with 400 mg (4800 U lipase) of fungal lipase given with each meal against no treatment. The stools were stored at −20°C until they were analysed for total fat content by the van der Kamer method.7 Statistical analysis was performed using the Wilcoxon’s rank-sum test for non-parametric paired data and signed rank test for unpaired data. Values indicated in the test are given as medians (range).

Results

Weight (Table)

Animals observed over a three week period pre-operatively showed no significant weight change within a range of −0.6 kg to +0.8 kg. Animals receiving no replacement therapy over a three week period showed significant weight losses of −0.9 kg (−1.7 to −0.2 kg) (p<0.01 v group A). When receiving fungal lipase or pancreatin the animals did not show significant weight loss compared with preoperative animals.

Faecal fat (Fig. 1)

A range of normal values was obtained from normal animals pre-operatively (9.9–12.3 mmol/24 h) with a median value of 10.8 mmol/24 h. Faecal fat excretion in group B was significantly raised when compared with normal animals demonstrating considerable fat malabsorption (p<0.01). There was no significant difference observed between animals of group A and

<table>
<thead>
<tr>
<th>Weight change (kg)</th>
<th>A normals</th>
<th>B placebo</th>
<th>C lipase</th>
<th>D Pancreatin</th>
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<tr>
<td>1</td>
<td>+0.5</td>
<td>-1.5</td>
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</tr>
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<tr>
<td>Median and range</td>
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<td>(-0.9 to -0.2)</td>
<td>(-0.2 to -0.3 to +0.4)</td>
<td>(-0.1 to -0.4 to +0.3)</td>
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<tr>
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<td>p&lt;0.01</td>
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</table>
group C. Similarly, no significant differences were found in group D dogs 14.4 mmol/24 h (9.0–35.6 mmol/24 h). No significant difference in fat excretion was observed between lipase treated and pancreatin treated animals.

**Stool volume** (Fig. 2)
Figure 2 shows that the pancreatectomised dog without exocrine therapy has steatorrhoea, but fungal lipase and pancreatin are equally effective in reducing stool volume to normal.

**Discussion**

Considerable and significant weight loss associated with fat malabsorption was confirmed in pancreatectomised animals receiving no exocrine replacement therapy over a three week period. The institution of either conventional pancreatin treatment or of acid stable lipase prevented this weight loss at the dosages used in this study. The lipase dosage of 400 mg provided 4800 U lipase and represented 8% of the lipase activity of the conventional therapy taking into account that over 90% of orally ingested conventional preparations of lipase are inactivated in the acid medium of the stomach.2

With the daily dosages of pancreatin and acid stable lipase, a statistically significant reduction of total faecal fat excretion/day of between 42 and 60% occurred when comparison was made over a similar period of no treatment. Considerable variations in faecal fat excretion in individual animals were noted, although weight changes were not so obvious. One animal with a mean faecal fat excretion before pancreatectomy of 11 mmol/24 h had concentrations of 40 and 36 mmol/24 h on replacement regimens. This observation may be because of the variation in acid output from the stomach and variable contributions of gastric and small intestinal lipase in different animals.9 This is probably a reflection of using a fixed dosage rather than tailoring the replacement therapy and the needs of the individual.

The most significant finding was the demonstration that a very small dose of acid stable lipase (400 mg) was as effective as pancreatin (10 g) in treating the steatorrhoea of pancreatic exocrine insufficiency in dogs. The study did not evaluate the effects of protein malabsorption, or examine the role of protein binding to bile acids as a cause of steatorrhoea. The results obtained may have considerable clinical implications in man. With the very large quantities of
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pancreatin necessary to replace missing enzymes in children with cystic fibrosis, problems of compliance as well as malabsorption are encountered. Furthermore, pancreatic extract contains high levels of purine and large doses of pancreatin can cause hyperuricaemia which may result in renal damage.\(^{11}\) Although the addition of H\(_2\) receptor antagonists\(^{12,13}\) and the development of microencapsulated pancreatin\(^{14}\) have improved the situation, the use of a high lipase content preparation which is acid stable may provide patients suffering from cystic fibrosis and chronic pancreatitis with a low dose, effective mode of therapy. Further studies in man would be worthwhile.

References

5 Regan PT, Malagelada TR, Dimagno EP, Glanzman SL, Go VLW. Comparative effects of antacids, cimeti-