Isolated lipase and colipase deficiency in two brothers

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Abstract

Two brothers of Arab origin, aged 15 and 10 years, with isolated congenital lipase and colipase deficiency are described. Both were normally developed with a history of passing greasy stools since early infancy. Both have remarkable steatorrhoea and low serum carotene and vitamin E concentrations. After exocrine pancreatic stimulation, lipase and colipase activities in the duodenal fluid were almost completely absent, while amylase tripsin, bile salt, and pH values were normal. No other aetiology for exocrine pancreatic insufficiency was found. This is the first report of congenital combined lipase and colipase deficiency in two brothers.

Results of pancreatic stimulation test (pancreozymin-secretin test) were consistent with isolated lipase and colipase deficiencies in both brothers (Table). Stool fat examined in three other siblings showed normal daily fat excretion. Vitamin E supplements together with pancreatic enzyme preparation were prescribed. Both patients, however, were lost to follow up.

Methods

PANCREOZYMIN SECRETIN STIMULATION TEST

A pancreatic stimulation test was performed as previously described. After an overnight fast, a nasoduodenal tube was placed under fluoroscopy into the third part of the duodenum. A basal sample of the duodenal fluid was collected into ice cold flasks during the first 10 minutes (basal). Pancreozymin (Kabi Diagnostica, Studsvik, Sweden) 2 U/kg was then slowly injected intravenously and three duodenal fluid samples at 10 minute intervals were collected as above. Subsequently, three more samples of duodenal aspirate were collected every 10 minutes following intravenous administration of secretin (Kabi 2 U/kg). The volume and pH value for each sample were determined immediately. Samples were then stored at –20°C until enzyme activity determinations.

ENZYME DETERMINATIONS

Trypsin was measured by a fluorimetric technique as described by Roth, using bovine trypsin (Sigma, Israel) as standard and expressed in μg/ml. Amylase was determined by the chromogenic enzymatic test adapted for duodenal juice (Phadebas amylase test, Pharmacia Diagnostics AB, Uppsala, Sweden) and expressed in U/ml. One unit of amylase activity is defined as the amount of enzyme catalysing the hydrolysis of 1 μmol glucosidic linkage per minute at 37°C. Bile salts were determined by an enzymatic spectrophotometric test for total bile acids (Sterognost -3a Pho, Nycomed AS, Oslo, Norway). Lipase activity in the duodenal samples was measured by turbidimetry according to Tietz et al., using absorbance measurements by spectrophotometry. Triolein 0.3 mM at pH 9.0 was used as substrate in the presence of 15 mM taurodeoxycholate and in excess of colipase (Sigma, Israel) 10–100 pmol. One unit of lipase activity is defined as the amount of enzyme that catalyses the transformation of 1 μmol triolein per minute at 37°C. Colipase activity was measured by its ability to reactivate bile salt inhibited lipase. Duodenal samples were heated to 80°C for 15 minutes for endogenous lipase inactivation, then added to a solution containing a standard amount of 100 pmol lipase (Sigma, Israel) derived from porcine
pancreas (MW 50000) and the restored lipase activity was determined by absorbance as described above. This activity was then transformed to moles of colipase (MW 11000) according to a standard curve (Figure).

Results

Amylase and trypsin activities showed normal responses to the pancreozymin secretin stimulation test, as well as the changes in pH and bile salt concentrations (Table). Lipase response in both patients showed only a trace – not more than 1 U/ml – (without exogenous colipase) which is less than 1.5% of the normal lipase activity (normal maximal response >80 U/ml). Colipase was unmeasurable in the duodenal fluid of the older brother (patient 1) and only a trace activity was detected in that of the younger (patient 2).

Discussion

Congenital, isolated exocrine pancreatic enzyme deficiencies are rare causes of malabsorption in infancy and childhood. Isolated lipase deficiency was reported in 12 patients in whom other more common causes of pancreatic disease were ruled out. In addition to its functional absence, no immunologically active lipase could be detected. The most characteristic manifestation was the passage of greasy-oily stools, often associated with steatorrhoea.

Colipase is a heat stable factor which was found to facilitate lipolysis in the presence of lipase. This is a glycoprotein secreted by the pancreas as procolipase which is cleaved by proteolytic enzymes, mainly trypsin, to result in the active moiety. It is the only factor that can activate lipase in the presence of bile acids. Colipase deficiency was implicated in the pathogenesis of some childhood onset pancreatic insufficiency. Congenital isolated colipase deficiency was reported only in two Assyrian brothers born in a consanguineous marriage. Their development was normal and no other pancreatic enzyme deficiencies were detected. Faye et al described the first case of combined isolated lipase and colipase deficiency. The measured activity of both enzymes was less than 2% of that found in normal control subjects. To the best of our knowledge the present report is the first to describe combined lipase and colipase deficiency in two brothers. In both, the residual lipase activity of the duodenal fluid was found to be less than 1.5% of that of the control, while the colipase activity was almost completely absent. The possibility of procolipase deficiency is valid since the pH and the trypsin activity which cleaves the procolipase were normal in the duodenal aspirate. Although steatorrhoea prevailed, probably since birth, growth and development were uneventful and no clinical signs of deficiency of any lipid soluble vitamin were noted. Serum carotene and vitamin E concentrations, however, were significantly decreased in both brothers. In patients with chronic severe fat malabsorption and vitamin E deficiency, a syndrome of progressive neuromuscular disorder characterised by ataxia, incoordination, areflexia, loss of vibratory sensation, and variable ophthalmoplegia develops. We believe that prolonged vitamin E deficiency would have eventually induced the neurological syndrome in both brothers. Therefore, aggressive vitamin E therapy as well as pancreatic extract is indicated in order to prevent any neurological degeneration. Although lipase and colipase activities in the pancreatic secretions of both brothers were almost completely absent, fat absorption was 70% and 79% of fat intake, respectively. This implies that in these patients extrapancreatic sources of lipase, such as lingual and gastric lipase, may have an important compensating role in fat digestion, thereby preventing the expected severe fat malabsorption and its morbidity sequelae.

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Gut 1990 31: 1416-1418
doi: 10.1136/gut.31.12.1416

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