Effect of taurine supplementation on fat and energy absorption in cystic fibrosis

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Abstract
In 10 children with cystic fibrosis and persisting steatorrhoea, supplementation with taurine (30-40 mg/kg/day) was given for two months as an adjunct to the usual pancreatic enzyme treatment. A three day fat and energy balance was performed in patients with cystic fibrosis, before and after the supplementation, and in seven healthy controls who did not receive taurine.

Faecal fat was measured by a gravimetric method and stool energy was determined using a bomb calorimeter. Patients with cystic fibrosis, before and after taurine, and healthy controls received the same fat and energy intake (calculated by a dietitian). In patients with cystic fibrosis taurine did not produce any improvement of steatorrhoea (mean (SD) faecal fat 8-7 (3-3) v 11-2 (7-0) g/day, respectively before and after the supplementation), of faecal energy loss (0-978 (0-468) v 1-133 (0-539) MJ/day), of faecal fat expressed as percent of fat intake (13-4 (5-6) v 15-1 (9-8)%), and of faecal energy expressed as percent of energy intake (9-9 (3-6) v 11-2 (5-7)%).

Healthy controls had significant lower fat (3-5 (2-3) g/day) and energy 0-576 (0-355) MJ/day faecal losses.

In conclusion, taurine failed to decrease significantly fat and energy losses. Our study does not support the use of taurine supplementation in the nutritional management of cystic fibrosis.

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The best known gastrointestinal complication of cystic fibrosis is pancreatic deficiency, which is present in 85% to 95% of patients.1 When untreated, pancreatic insufficiency may be responsible for a high degree of morbidity and may affect growth and nutrition. Adequate pancreatic enzyme replacement is required in order to correct nutrient malabsorption, but fat absorption often remains suboptimal despite the use of pancreatic extracts and of drugs which affect duodenal acidity.

As the predominance of glycine bile acid conjugates with almost complete disappearance of tauroconjugates could play a part in the fat malabsorption in cystic fibrosis,2 taurine supplementation has been proposed as an adjunct to standard treatment. Nevertheless, the results of published papers are conflicting. A significant improvement of fat absorption after taurine supplementation was demonstrated by two different groups,3,5 whereas the studies of Thompson et al concluded that taurine is not indicated in the nutritional management of cystic fibrosis.6 7 In addition to this, Fitzpatrick et al were unable to demonstrate a change in fat solubilisation in patients with cystic fibrosis with ileal resection after taurine supplementation.8

To determine whether taurine could play a part in fat malabsorption in cystic fibrosis, we studied the effect of taurine on fat and energy malabsorption in a group of patients with cystic fibrosis with various degrees of intestinal malabsorption that were only in part corrected by pancreatic extracts.

Patients and methods
PATIENTS
Ten patients with cystic fibrosis, seven boys and three girls, with a mean age of 7.7 years (range 2–13 years) were admitted to the study. The inclusion criteria were the persisting clinical symptoms due to an inadequate correction of the pancreatic insufficiency despite the enzyme replacement (mainly, persisting steatorrhoea and failure to gain weight while on an unrestricted diet). The patients had a Shwachman clinical score ranging between 45 and 96 (mean 71) and a degree of respiratory involvement judged from mild to severe. No ileal resection had previously been performed. No patient had clinical and biochemical signs of liver and biliary tract diseases and no one had a height ratio <5th centile. Vitamin supplementation, aerosol treatment, and physiotherapy were unchanged during the study.

All patients were supplemented with an enteric coated enzyme preparation of pancreatin (Pancrease, Cilag). The number of capsules, ranging from a minimum of six to a maximum of 20 a day in the whole study group, remained the same during the study.

DESIGN OF THE STUDY
The experiment was performed on an ambulatory basis and the patients, who were well at the time of the study, remained clinically stable for the whole period of supplementation. Taurine (O-Due, Nativelle) was administered for two months twice daily at a dose of 30–40 mg/kg/day as suggested by Darling et al.4

Before and two months after starting the supplementation a fat and energy balance study was performed in each patient. Weighed food intake was recorded for a period of six days by a daily diary and a dietitian controlled the records to check for errors. The weighed food intake was analysed using food composition data.9
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Gross energy intakes were estimated by multiplying the intakes of fat (39.3 kJ/g), protein (as nitrogen × 6.25; 23.6 kJ/g), and carbohydrate (17.6 kJ/g) by their respective heat of combustion values. All stools, collected during the three days of the record period, were kept in polyethylene bags and immediately frozen. Coal tablets were used as charcoal markers between the start and the end of the collection period.

All the patients were cooperative and no gross errors were found in the daily diary or the procedure for stool collection. During the collection periods the fat content of the diet was not different from the patients’ usual diet (about 3 g/kg/day).

A fat and energy balance study was also performed in seven healthy children, with a mean age of 10 years (range 6–12 years), who had no known gastrointestinal and metabolic diseases. They were studied as controls and did not receive taurine supplementation.

ANALYSIS

The pooled three day stools were weighed and dried in an oven at 99°C for 24 hours. The weight of the dried stool was recorded and expressed as a % of wet stool.

Fat analyses were performed on the dried stool samples. A modification of the gravimetric method of Zuckerman et al was used. Four weighed samples of dried stools (0.5–2 g) were subjected to acid hydrolysis by boiling in 4 M hydrochloric acid. The hydrolysate was filtered and the filter papers were washed with hot water until neutral pH was achieved. They were then dried and extracted with petroleum ether (60–80°C) into preweighed aluminium cups. The extraction was performed in a Soxtec Ht 1043 extraction unit (Tecator). The cups containing the extracted fat were dried and weighed again, and the difference in cup weight was defined as the sample’s fat content.

Energy was determined on dried stools using a ballistic bomb calorimeter (Gallenkamp) as reported elsewhere. Results are given as faecal fat (g/day), faecal energy (MJ/day), and % of fat intake and % of gross energy intake. All the values are expressed as mean (±SD).

The significance of the difference between two arithmetic means was evaluated with the paired and unpaired Student’s t test and differences between means were considered significant where p<0.05.

Results

Taurine supplementation was well tolerated and no side effects were noted. All patients complied with the treatment, as written records of the daily consumption of taurine and of pancreatic extracts were reported by the parents.

In patients with cystic fibrosis daily fat intake was not different in the two periods of stool collection (before taurine, 70±5 (22±5) g/day; after taurine, 76±3 (21±3) g/day). In healthy controls, fat intake was 79±4 (27±8) g/day.

In patients with cystic fibrosis the weight of wet stools was not different before and after taurine supplementation (179 (116) vs 168 (61) g/day). The dry weight expressed as a % of wet weight in both groups was 24±1 (4±4) vs 25±8 (5±6)%, respectively. In healthy controls the weight of wet stools was 141 (101) g/day and the dry weight 20±5 (5±1)% of faecal wet weight.

Data about faecal fat losses in patients with cystic fibrosis show that taurine administration did not improve faecal fat loss (before taurine, 8±7 (3±3) g/day; after taurine, 11±2 (7±0) g/day; p<0.006). Values of faecal fat loss in healthy controls were significantly lower than those found in patients with cystic fibrosis before taurine (3±5 (2±3) g/day vs 8±7 (3±3) g/day; p<0.001; fig 1).

As shown in fig 2, in patients with cystic fibrosis daily fat intake as a % of energy intake was not different before and after taurine supplementation (11±2 (3±3) vs 10±4 (2±7)%. In healthy controls, faecal fat intake was 10±3 (2±6)% of energy intake.

Figure 1 Daily stool fat expressed as g/day in cystic fibrosis patients before and after taurine supplementation and in the control subjects. The bars represent the mean values.

Figure 2 Daily stool fat expressed as % of fat intake in cystic fibrosis patients before and after taurine supplementation and in the control subjects. The bars represent the mean values.
fibrosis the faecal fat expressed as % of fat intake was 13.4 (5.6)% before taurine and 15.1 (9.8)% after taurine (the difference was not significant). In controls the faecal fat expressed as % of fat intake was lower than in patients with cystic fibrosis (4.3 (2.2)%; p<0.001).

Gross energy intake was 10.11 (2.69) MJ/day and 10.44 (2.57) MJ/day in patients with cystic fibrosis, respectively before and after taurine; in controls subjects it was 10.065 (2.215) MJ/day.

In patients with cystic fibrosis, faecal energy loss was not different in the two periods (before taurine, 0.978 (0.468) MJ/day; after taurine, 1.133 (0.539) MJ/day). Healthy controls had a mean faecal energy loss significantly lower than patients with cystic fibrosis (0.576 (0.355) MJ/day vs 0.978 (0.468) MJ/day; p=0.037; fig 3).

As illustrated in fig 4, in patients with cystic fibrosis faecal energy expressed as % of gross energy intake did not change before and after taurine administration (respectively, 9.9 (3.6)% v 11.2 (5.7)%; the difference was not significant). Healthy controls had a faecal energy expressed as % of gross energy intake significantly lower than patients with cystic fibrosis (5.5 (3.0); p=0.009).

Discussion
To our knowledge this represents the first longitudinal study that measures both faecal lipid and energy losses in patients with cystic fibrosis before and after taurine supplementation. The main consequence of pancreatic insufficiency in cystic fibrosis is the failure to digest and absorb some components of the diet. Fat and nitrogen malabsorption are common in these patients and the faecal energy losses reflect the nutrients' malabsorption.

As enzyme replacement diminishes but does not normalise the amount of fat and nitrogen lost in stools, the use of other drugs that can improve fat malabsorption has been suggested in cystic fibrosis. Nevertheless, a definitive resolution of the symptoms due to steatorrhoea is rarely achieved. In this study we were unable to demonstrate that taurine leads to an improvement in the absorption of fat and energy in 10 children with cystic fibrosis who had persisting fat malabsorption despite appropriate enzyme treatment. These findings contrast with most of the reports about the use of taurine in cystic fibrosis, but are similar to the conclusions of other studies, which did not support the therapeutic use of taurine in the nutritional management of cystic fibrosis.

The disagreement in the results of the studies on taurine supplementation in cystic fibrosis could be related to differences in the characteristics of the study population. People who suggest the use of taurine in cystic fibrosis recommend it particularly in patients with a significant degree of steatorrhoea, but our results, like those by Thompson et al, do not reveal any improvement in fat and energy absorption, even in cases with a low baseline high fat and energy losses.

The dose of taurine and the length of the supplementation cannot represent a bias: in our study taurine was given at the same dose and for the same period as in another study that concluded that taurine was effective in cystic fibrosis. However, Thompson et al showed that the glycine-taurine conjugates ratio is remarkably sensitive to even relatively small doses of taurine.

Recently, another group has published data regarding the energy and fat content of stools in patients with cystic fibrosis. Similar values of both faecal fat and faecal energy have been found in our patients.

In healthy patients we found a slightly higher fat and energy faecal excretion than those
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reported by Murphy et al (respectively 3·5 v 2·2 g/day and 0·577 v 0·339 MJ/day).14 Our values are similar to those seen in adults.15

Fecal energy does not represent only malabsorbed energy: in addition to maldigested and unabsorbed dietary residues, fecal energy losses can be due in part to colonic bacteria and to products from cellular lysis, which represents up to 30% of the fecal energy.14

In our study, fat absorption was estimated in terms of steatorrhea and of the percentage of dietary fat intake, which had been strictly checked by a dietitian. In the previous studies no detailed data about the dietary intake of fats are given: the variation of fat intake itself may account for changes in faecal fat loss unrelated to fat absorption. Furthermore, in one of these studies the authors give results in terms of 'steatorrhea', that is quite different from the percent of fat intake.6

In cystic fibrosis some evidence exists for impairment of fat absorption, which is distinct from the maldigestion of fat consequent on absence of pancreatic lipase. Some steatorrhea may persist with enteric coated enzyme preparations, even when optimal intraluminal pH for exogenous lipase is achieved by the use of cimetidine and sodium bicarbonate.13 A reduction in the bile acids pool resulting from decreased ileal reabsorption may be a partial explanation.4

The efficacy of taurine in cystic fibrosis is based on the assumption that this amino acid provides the necessary substrate for the synthesis of taurine-conjugated bile acids, which in turn, could improve the micellar solubilisation of lipolytic products.16 Our patients had no liver and biliary tract disease. Nevertheless, as ileal mucosal bile acids malabsorption has been postulated in cystic fibrosis,17 we cannot definitively state that our patients did not benefit from taurine supplementation because taurine-conjugated bile acids are less efficiently absorbed across the ileum.

In our study a variability in the response to supplementation appears in some individual patients. An explanation for this and for the considerable variability of the response to taurine supplementation among the various groups3-7 is the great heterogeneity of fat malabsorption in cystic fibrosis, which can be due both to lipase deficiency and to decreased solubility of bile acids micelles in the unstirred water layer. In this view the hypothesis of the association between taurine deficiency and biliary abnormalities in some patients with cystic fibrosis appears interesting. This finding supports the conclusion that taurine could be helpful only in selected patients, particularly if fat malabsorption is more related to the bile acid component than to the exocrine pancreatic insufficiency itself.

Taurine supplementation is also a controver-
sial issue in low birthweight infants, who usually present with a low fat absorption.18 The reasons for fat malabsorption in these babies are various and not fully defined: pancreatic lipase may be reduced, bile acid pool sizes, rate of synthesis and intraluminal concentrations are low, and absorption of bile salts may be impaired. No agreement has been reached about the effectiveness of supplementation of formulas with taurine on fat absorption in low birthweight infants.14-20

In conclusion, pending further studies on taurine administration in cystic fibrosis, we do not recommend an indiscriminate supplementation with taurine in every patient with cystic fibrosis suffering from persisting steatorrhea. More work is still necessary to confirm the exact role of taurine in cystic fibrosis.