An Infant with Both Cystic Fibrosis and Coeliac Disease

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Coeliac disease (gluten enteropathy) and cystic fibrosis are both diagnosed fairly often in paediatric practice. The incidence of coeliac disease in London was estimated in 1959 as 1 in 3000 (Carter, Sheldon, and Walker, 1959), but this is almost certainly an underestimate, for the data were collected before the routine use of jejunal biopsy enabled the condition to be diagnosed more accurately. A figure of 1 in 1778 was found in Glasgow (Black, 1964), and this estimate has recently been confirmed (McCræ, 1968). Recent figures for cystic fibrosis give an incidence of 1 in 2448 live births in Australia (Danks, Allan, and Anderson, 1965) and 1 in 1863 in the United States (Honeyman and Siker, 1965). The chances of both conditions occurring together in one patient is thus extremely remote, but the boy reported here satisfies the current criteria for making both diagnoses. We know of no other report in the literature of the disorders coexisting.

Case History

A male infant, born on 7 March 1967, after an uneventful pregnancy lasting 39 weeks was the first child of non-consanguineous parents. At birth he weighed 2·7 kg., and from the first day of life he developed loose stools from which Pseudomonas pyocyanea were cultured. He gained weight extremely slowly on a full-cream milk, and cereals were added to his diet at the age of 4 weeks. At the age of 12 weeks he was admitted to Bristol Royal Hospital for Sick Children for investigation of his failure to thrive.

On examination he was marasmic with a grossly distended abdomen (Fig. 1a), and was continually passing frequent bulky, pale stools. Relevant investigations showed Hb 10·6 g., with normal red cell morphology. His 5-day faecal fat excretion was 18·4 g. per day and the levels of stool tryptic activities were 6, 7, 21, 1, and 2 units (Azocasein; normal is above 10 units). Microscopy of the stools revealed no abnormality, culture was negative, the stool pH was between 6 and 7, and the sugar chromatogram was negative. The β-lipo-proteins in the blood were normal. Duodenal aspirate showed a tryptic activity of 2 units (Azocasein) and an amylase activity of 1600 Somogyi units per 100 ml. A chest x-ray showed some overdistension of the lung bases, with increased lung markings in these regions but no specific abnormality.

Clinical course. During the weeks after admission when the baby was being investigated, he developed at different times a coliform septicaemia, dysentery due to Shigella sonnei, and a chest infection from which no specific organism was grown.

Though usually excessively hungry he failed to gain weight (Fig. 2). This, together with a low stool tryptic activity and low enzyme activity in the duodenal juice, led to a clinical diagnosis of cystic fibrosis. At the age of 17 weeks, the first jejunal biopsy was per-

![Fig. 1.—Patient (a) before treatment, aged 4 months, and (b) after 3 months on a gluten-free diet, aged 7 months.](image-url)
Three kissed.

gluten-free diet infiltration of the lamina impressively continued, and progress was satisfactory, so that at the age of 1 year he was over the 50th centile for weight and height. He occasionally still had minor chest infections and wheezing.

During the second year of life he developed normally apart from a pot-belly, and at the age of 21 months was at the 75th centile for weight and the 50th for height. In order to confirm both diagnoses, two further sweat sodiums were estimated and were 72 and 86 mEq/l. He was challenged with gluten, and after 2 biscuits and a sausage sandwich he developed frequent foul-smelling stools, abdominal discomfort, and became irritable and miserable. His parents were unwilling to continue him on a gluten-containing diet.

**Discussion**

The diagnosis of cystic fibrosis seems incontrovertible. The steatorrhoea, liability to infection, low duodenal and faecal trypsin, and raised sweat sodium satisfy the usual diagnostic criteria. In our experience, the intestinal mucosa in this condition does not show the changes found in coeliac disease. In infants, we have in fact found these changes only in coeliac disease, though experimentally they have been noted to follow repeated applications of irritant substances to enterostomies (Townley, Cass, and Anderson, 1964). A similar appearance has also been reported after neomycin (Jacobson, Prior, and Falcon, 1960). The changes in the small gut mucosa in malnutrition and bowel infections have previously been investigated and do not show the changes found in coeliac disease or in this baby (Burman, 1965).

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Summary

An infant with failure to thrive was shown by jejunal biopsy and response to gluten-free diet to have coeliac disease and by examination of sweat electrolytes and duodenal juice to have cystic fibrosis. This association has not previously been reported. He responded well to treatment of both conditions.

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References


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