Hepatic dysfunction in association with pancreatic insufficiency and cyclical neutropenia

Shwachman-Diamond syndrome

The syndrome of pancreatic achylia and cyclical neutropenia (Shwachman-Diamond syndrome), has been recognized for 13 years, though it now encompasses a far wider range of clinical abnormalities than was initially realized. In 1964 Shwachman et al. first distinguished these patients from those suffering from cystic fibrosis, and commented on an association of pancreatic insufficiency with growth retardation, bone marrow dysfunction, raised Hb-F levels, and galactosuria. Skeletal abnormalities were next noted, and in 1967 Burke et al. observed metaphyseal dysostosis in 3 cases; subsequent reports by others have recently been reviewed by McLennan and Steinbach (1974). In addition to metaphyseal disturbances, abnormal tubulations of long bones, clinodactyly, phalangeal hypoplasia, and narrowing of the sacrosciatic notches have been noted. Dys-y-globulinaemia was next noted by Goldstein (1968) and has been described in 5 further patients (Doe, 1973), in each case involving a reduction in the serum concentrations of one or more of the immunoglobulins. As far as we can determine only one living patient with histological but not clinical evidence of liver dysfunction has previously been described (Bodian et al., 1964). In a retrospective review of 18 children whose necroses had shown exocrine hypoplasia and lipomatosis of the pancreas they noted that 11 had shown a fatty liver or cirrhosis. Cyclical neutropenia had not been shown in life in any of them. Higashi et al. (1967) observed inflammatory cell infiltration in the perportal tissue of the liver obtained at necropsy in a 2-year-old child with pancreatic insufficiency and cyclical neutropenia.

We presently have 4 children with this syndrome under our care as detailed in the Table. One of them (Case 1) has clinical evidence of hepatic dysfunction and is the subject of the case report.

Case report

A male, weighing 2.9 kg at birth, is the son of healthy unrelated parents. Frequent, loose, offensive stools were noted from birth, but appetite was good and there was no unusual frequency of respiratory or other infections. Height and weight remained below the 3rd centile, and he was referred (with his sister, Case 2) for investigation at the age of 9 months. On examination there were no abnormalities apart
from his stature and a mild degree of abdominal distension. Steatorrhoea was present (46.2 mmol fat/24 hours), and fat globules were present in the stool. After stimulation of the pancreas with pancreozymin no trypsin appeared in the duodenal juice, and the lipase level was very low (0.25 units/0.1 ml; normal >2 units/0.1 ml). Sweat electrolytes were normal on three occasions, as was xylose absorption. Hb was 11.7 g/dl and the range of white cell counts was 7.2–23.0×10⁹/l (neutrophils 2.6–13.3×10⁹/l). Platelet count was normal but has fallen to 11.7×10⁹/l under further observation. Hb electrophoresis showed an increase in the proportion of Hb-F (3.3%). Serum calcium, phosphorus, cholesterol, and α₁-antitrypsin levels were normal, as were urine and plasma amino acids. There were no reducing substances present in the urine.

Radiological examination showed normal lung fields and bone age, but there was expansion of the anterior ends of the ribs. At that time no other metaphyseal abnormalities were present; however by the age of 2 years dysostosis of the femoral metaphyses was noted, while the rib changes were less marked.

Immunological studies showed normal IgG levels (4.8 g/l) but reduced IgM (0.07 g/l) and only a trace of IgA. Salivary IgG was increased (0.22 g/l) but only trace levels of IgA were detected. Alanine aminotransferase levels (70 IU/l), and aspartate aminotransferase levels (135 and 179 IU/l) were high. Alkaline phosphatase was 32 KA units and serum albumin 42 g/l. At no stage over the next 4 years was he jaundiced, nor was hepatosplenomegaly observed. During this time he was given pancreatic with meals and vitamin supplements. He has remained well, his height increasing to the 3rd centile but his weight remaining below that level. Further investigation has confirmed the continuing presence of steatorrhoea (40.7 mmol fat/24 hours), cyclical neutropenia (0.18–9.96×10⁹/l) and a raised aspartate aminotransferase level (76 IU/l). A needle liver biopsy taken at the age of 2 years showed variation in size of the hepatocytes and expansion of the portal tracts due to the presence of reticulin fibres (Fig.)

### Discussion

We consider that this patient clearly falls within the so-called Shwachman-Diamond syndrome, and shows many of the features more recently described. The presence of a degree of portal fibrosis and continuing high serum transaminase levels, in the absence of any episode of acute liver disturbance, suggest that these findings represent chronic liver damage and should perhaps be regarded as forming part of the spectrum of the syndrome. The pathogenesis of this disorder remains obscure. It has been noted by others among sibs. It is therefore of interest that this patient’s sister (Table, Case 2) has pancreatic insufficiency and other features including metaphyseal dysostosis of the femora, low IgG, short stature, and increased concentration of Hb-F (3.5%). Neutropenia, however, has not been observed despite a careful search. Her sweat test and liver function tests are normal.

Flaring of the anterior ends of the ribs was noted in 3 of the children described by Burke et al. (1967), and in one of those in the review of McLennan and Steinbach (1974). It is notable that these patients were under 2 years of age, as was the boy we describe and one other patient under our care with a similar finding (Table, Case 2). By the age of 4 years the ribs of both our patients had returned to normal, but metaphyseal dysostosis had become apparent. All
our 4 patients have skeletal changes and are of short stature.

The dys-γ-globulinaemias described have involved a reduction in IgA levels in all but one case, while low levels of IgG or IgM were found in 4 cases (Hudson and Aldor, 1970; Doe, 1973; McLennan and Steinbach, 1974). The patient described here shows a more complete IgA deficiency than previously described, and the other patients in the Table show the wide variation in immunoglobulin abnormalities that can occur.

This report illustrates the broad spectrum of this curious condition, and supports the contention that hepatic involvement should be regarded as a feature of the syndrome.

Summary

A patient with pancreatic insufficiency and cyclical neutropenia is described who also has evidence of hepatic dysfunction. He and 3 other patients whose findings are given emphasize the wide range of abnormalities seen in this syndrome.

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References


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