Case reports


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**Alpha-chain disease in an Englishman**

S. W. S. WANG*

M.B., Ch.B., M.R.C.P.

Liverpool Royal Infirmary

Summary

Eighty cases of the gastro-intestinal form of alpha-chain disease had been reported up to the time of writing this communication (September 1975). All patients were non-Caucasians. This report presents a case of this disorder in an Englishman who became ill whilst working in the tropics. His clinical features and response to therapy were entirely in line with the other cases so far reported.

Introduction

The first case of alpha-chain disease was described in 1968 (Ramboud et al., 1968; Seligman et al., 1968). By July 1975 a total of eighty cases had been reported (Doe et al., 1972; Manousos et al., 1974; Seligmann, 1975a, b).

Abdominal pain, weight-loss, steatorrhoea and marked finger clubbing are the main features (Ramboud and Matuchansky, 1973; Seligmann, 1975c).

* Present address: Department of Cardiology, Manchester Royal Infirmary, Oxford Road, Manchester M13 9WL.

A case is now reported where the presenting features were vomiting due to upper intestinal obstruction, weight-loss, steatorrhoea, anaemia and occult gastro-intestinal tract bleeding, and finally response to treatment with cyclophosphamide, prednisolone and tetracycline was achieved.

Case history

The patient, an Englishman, was first seen in 1966 (when he was 31 years old) at the School of Tropical Medicine, Liverpool, with a 7-month history of vomiting and diarrhoea, which had developed while he was working in Cameroon, West Africa. He had earlier worked for 3 years in Mauritius. Apart from having had 'stomach trouble' as an adolescent, there had been no significant illnesses in the past.

In December 1965 he developed vomiting and diarrhoea which lasted for about 1 week and then improved, but there were several recurrences in the course of the next few months. In July 1966 he developed severe and persistent vomiting and diarrhoea. He was sent back to Britain for medical
attention; he had lost 9.5 kg in weight in 7 months and now weighed only 62.5 kg.

When seen at the School of Tropical Medicine, no evidence was found to suggest that he might have contracted a tropical disease. However, steatorrhoea was discovered. He received a diet rich in protein and low in fat content. At review in January 1967, steatorrhoea had persisted and he had lost more weight (Fig. 1). There was no anaemia or oedema; blood urea and serum electrolytes, calcium, phosphate and alkaline phosphatase were all normal. The serum albumin was 2.85 g/100 ml and globulin 2.34 g/100 ml.

Jejunal biopsy was carried out using a Crosby capsule. Histology showed 'a mixture of partial and sub-total villous atrophy with marked infiltration by eosinophils, lymphocytes and plasma cells'. He was treated with a gluten-free diet and a temporary improvement was noted. Later, pancreatin, vitamin D and oral calcium therapy were added.

By February 1968 he had gained 7 kg in weight and the vomiting and diarrhoea had ceased. In November of the same year, however, his condition relapsed, with return of vomiting and diarrhoea. He was admitted to Liverpool Royal Infirmary where a barium meal revealed obstruction at the level of the distal duodenum and proximal jejunum. A laparotomy was carried out in January 1969. The whole of the small intestine was found to be oedematous. There was no enlargement of lymph nodes, and no hepato-splenomegaly. A posterior gastro-jejunostomy and a duodeno-jejunostomy were performed to relieve obstruction. Specimens of duodenum and lymph node were sent for histology. The duodenal mucosa showed marked infiltration with eosinophils, lymphocytes and plasma cells. The lymph node was reported as normal.

His condition fluctuated after the by-pass operation. In March 1970, he was found for the first time to be anaemic. The haemoglobin was 10.5 g/100 ml, the anaemia being microcytic and hypochromic. Repeated tests for occult blood in faeces were positive. He received oral iron therapy and vitamin B12 injection. In April 1971 vomiting and diarrhoea recurred and he was again admitted to hospital. He was wasted and pale. The haemoglobin was now 8.9 g/100 ml. The serum calcium was 9.3 g/100 ml, and both phosphate and alkaline phosphatase were normal. The serum albumin was 2.9 g/100 ml and globulin 2.8 g/100 ml. A barium meal suggested obstruction at the level of the proximal jejunum.

The possibility of Crohn's disease was considered and he was treated with prednisolone and sulphasalazine, and was transfused. During the next 3 years his condition remained static.

In April 1974 he was re-admitted to hospital owing to further deterioration. He had now lost 21 kg in weight altogether since 1966, and weighed only 49 kg. He was emaciated and pale. There was, however, no clubbing, oedema, or lymphadenopathy. The liver and spleen were not enlarged.

There was still evidence of blood-loss from the gastrointestinal tract. The haemoglobin was 7.4 g/100 ml. The serum iron was 30 µg/100 ml, and the total iron-binding capacity was only 7% saturated. The serum albumin was 2.3 g/100 ml and globulin 1.8 g/100 ml. A further barium meal showed obstruction at the previously constructed gastro-enterostomy.

A further laparotomy was performed to refashion
the by-pass. At operation, oedematous and atonic gut was observed. The post-operative course was stormy, with chest and wound infections occurring.

At this stage, alpha-chain disease was suspected. An operative specimen of jejunum was studied in the University sub-Department of Immunology (Director, Dr J. Bradley). The jejunal mucosa was infiltrated with lymphocytes and plasma cells at the villi and their bases. The cells were considered to be producing alpha-chain since they fluoresced with anti-alpha-chain antibodies. Unfortunately, owing to the shortage of anti-light-chain anti-sera, a confirmatory immuno-electrophoretic test of the patient’s serum for free alpha-chain (which is the heavy chain of IgA) was not carried out. Serum immunoglobulin studies showed: IgG, 680 mg/100 ml (normal, 220–1240 mg/100 ml); IgA, 190/100 ml (normal, 70–280 mg/100 ml); IgM 280 mg/100 ml (normal, 35–120 mg/100 ml).

The patient was treated with cyclophosphamide 100 mg daily, prednisolone 5 mg t.d.s. and tetracycline 500 mg q.d.s. Later, because of a fungal infection of the nails, he was given griseofulvin. Two months later, the vomiting and diarrhoea had disappeared and he had gained 7 kg in weight. In December 1974, his weight was 66-22 kg. The cyclophosphamide therapy was discontinued and the prednisolone dosage reduced to 5 mg b.d.

When last seen, in September 1975, the patient weighed 70-76 kg and had recovered his good health. Haemoglobin, plasma urea and electrolytes were all normal.

Discussion

Alpha-chain disease was initially regarded as geographically region-specific, and hence the term ‘Mediterranean’ was applied to it. Later, it was regarded as ethnic-related (Ramat and Hulu, 1975; Doe, 1975).

The present case is believed to be the first example of the intestinal form of this condition to be reported from a Caucasian.

There is a remarkable similarity between patients with this disease, and in the present case, the clinical picture, the course of the illness and the response to therapy were similar to those in the other reported cases. Unfortunately, the full confirmatory tests were not performed but it would be reasonable to make the diagnosis of alpha-chain disease on the basis of the features described.

It is interesting to note that the patient apparently responded initially to a gluten-free diet. It is hard to say whether such a response was merely a chance finding or whether he did initially have coeliac disease and later developed alpha-chain disease. Later, the apparent response to a regimen of prednisolone and sulphasalazine might well have been due to a partial response of the alpha-chain disease to the systemic steroid.

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Addendum

Since the submission of this paper, another case of alpha-chain disease has been reported (Shulman, Lai York and Grieve, 1975), bringing the total to eighty-two. But the present report remains the first from a Caucasian patient.

References


Alpha-chain disease in an Englishman.

S. W. Wang

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