Reversible intestinal mucosal abnormality in acrodermatitis enteropathica

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Acrodermatitis enteropathica (Margileth, 1963) is a rare hereditary disorder of infancy characterized by distinctive cutaneous lesions primarily localized to the periorificial areas and extremities, alopecia (usually total), gastrointestinal symptoms, dystrophy of nails, and growth failure.

Neldner et al. (1974), in a recent survey, stated that light microscope examinations of the gastrointestinal tract in acrodermatitis enteropathica had been essentially normal, though they referred to a few reports of necropsied patients who had shown focal areas of mucosal degeneration at various levels in the jejunum, ileum, and colon.

Ament and Broviac (1973) reported one case in which a mucosal lesion in the small and large intestine did not revert to normal on therapy which included di-iodohydroxyquinoline (DIH). Moynahan (1962) reported duodenal biopsy with normal histology in an 18-month-old female with acrodermatitis enteropathica treated with DIH, though there was no mention of whether biopsy was taken before or after DIH. Fry, McMinn, and Schuster (1966) referred to 2 cases of acrodermatitis enteropathica in whom biopsy was taken before treatment was started, but did not describe any structural abnormality of the intestinal mucosa. In contrast, all 3 cases described in this report showed similar intestinal mucosal abnormalities on duodenal biopsy at the outset of treatment and all reverted to normal with therapy.

Case reports

Case I. A male aged 9 months presented with a rash of 6 months’ duration which had appeared in the napkin area at 3 months and 2 weeks later around the mouth followed by ulceration in the mouth. It then spread around the eyes and nares and he developed blisters followed by areas of redness and scaling on the knees, hands, and feet. There was no diarrhoea. At birth black hair covered the scalp, but this was completely lost at 3 months of age. He was breast fed until 7½ months, and had received cereal with cow’s milk and other solids from the age of 2 months. A younger brother is normal.

A diagnosis of acrodermatitis enteropathica had been made before admission to hospital and DIH had been administered in dosage of 1300 mg daily for 9 weeks without significant improvement. After admission the dosage was doubled and expressed breast milk feed begun. The rash cleared and the dosage of DIH was progressively reduced with short temporary increases during mild recrudescences of rash which occurred with intercurrent infection. Expressed breast milk feeds were stopped after 2 months. This regimen was maintained for 5 years after which DIH was discontinued and zinc sulphate solution (50 mg three times daily) begun. Duodenal biopsy was performed 5 days after starting treatment with expressed breast milk and DIH and...
again after 4 weeks and 6 weeks of DIH therapy and before oral zinc. A further biopsy was obtained after 6 months' zinc therapy.

The rash cleared with DIH but after a cold he usually developed mild recurrences of rash which subsided with temporary increase in DIH dosage. On zinc sulphate, however, his skin has remained completely clear of rash at all times. Hair growth has remained normal. At the age of 6 years his height was 110 cm and weight 19·25 kg.

Case 2. A female aged 4 months presented with rash of 10 weeks' duration. One elder sister is normal. Ulcerated areas first appeared in the mouth at the age of 2 weeks, followed at 6 weeks by rash around the mouth and nares, on the cheeks, the back of the head, and the outer aspects of the ears. By 3 months the vulva, buttocks, inner thighs, knees, hands, and feet were affected and she developed diarrhoea with five or six liquid motions daily. Short sparse hair was present on the scalp and eyebrows and eyelashes were almost completely absent. She was fed on an artificial milk formula until the age of 4 months. A diagnosis of acrodermatitis enteropathica was made and treatment with expressed breast milk and DIH was begun. DIH dosage was increased from 900 mg to 1800 mg daily with progressive improvement of the rash and cessation of diarrhoea. Expressed breast milk was stopped and DIH continued alone, the dosage ultimately being reduced to 900 mg daily increasing up to 1800 mg daily during minor recurrences of rash and lethargy which occurred every 4 to 6 weeks. This was continued for 4 years when DIH was stopped and zinc sulphate solution (50 mg three times daily) begun. Duodenal biopsies were carried out before any treatment was given and again after 3 weeks and before oral zinc. A further duodenal biopsy was obtained after 6 months' zinc therapy. On DIH there was regrowth of scalp hair, eyebrows, and eyelashes. There have been no relapses on zinc sulphate treatment. At the age of 5 years she weighed 18·25 kg and height was 100 cm.

Case 3. A male aged 5 months presented with a red, crusted rash of one-month duration appearing at first on the buttocks, groins, thighs, scrotum, and penis, with small patches elsewhere. It then appeared around the mouth and on the cheeks some days later. Scalp hair, eyebrows, and eyelashes were normal and there was no diarrhoea. He had been breast fed until the age of 3 months and was then changed to cow's milk. 3 elder brothers are normal. A diagnosis of acrodermatitis enteropathica was made and treatment begun with expressed breast milk alone. 10 days later DIH was added in dosage of 900 mg daily increasing to 1800 mg daily and then reducing to 900 mg daily after the rash cleared. DIH was discontinued after 20 months and zinc sulphate solution (50 mg twice daily) begun. Duodenal biopsies were performed before DIH therapy, and again before oral zinc. A third biopsy was obtained after 6 months' zinc therapy. At 2½ years of age his weight was 16·55 kg and height 93 cm.

Investigations. Haemoglobin estimations, skin and stool cultures, and serum immunoglobulins were investigated at initial presentation (see Table). Serum zinc levels were measured before, and several months after, starting oral zinc using the Varian Techtron AA3 spectrophotometer with an AA5 readout (Olson and Hamlin, 1968). Periodic eye examinations for optic neuritis were carried out in all patients when on DIH.

Duodenal biopsy. The biopsy procedure is described in detail elsewhere (Townley and Barnes, 1973). Specimens were taken from the fourth part of the duodenum.

Results

The initial biopsies taken before treatment or at the start of treatment showed loss of villous architecture with flattening of villi, which in some areas was as severe as that seen in coeliac disease. There was increased cellularity in the lamina propria due to infiltration with inflammatory cells. The intestinal epithelial cells were cuboidal in shape and the nuclei were enlarged, rounded, and less elongated with an open chromatin distribution in contrast to the normal compact chromatin arrangement. This same pattern of change was seen in all 3 patients (Figs 1a and 2a).

<table>
<thead>
<tr>
<th>Case no.</th>
<th>Haemoglobin (g/dl)</th>
<th>Skin swab culture</th>
<th>Stool culture</th>
<th>Stool and skin culture for Candida albicans</th>
<th>Serum immunoglobulins</th>
<th>Duodenal histology</th>
<th>Serum zinc (normal range 75–140 μg/100 ml)</th>
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<td>Normal</td>
<td>Normal</td>
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<td>NPI</td>
<td>Negative</td>
<td>Normal</td>
<td>Normal</td>
<td>65</td>
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</tbody>
</table>

NPI, no pathogen isolated.
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The biopsies taken after treatment with expressed breast milk and DIH showed partial re-establishment of villous architecture and reduction of the cellular infiltration of the lamina propria (Figs. 1b and 2b). The nuclei of the epithelial cells still showed the open distribution of the chromatin and were still more rounded than the normal elongated nucleus. Also the intestinal epithelial cell nuclei were located more centrally in the cell as compared with the normal basal location. After zinc sulphate therapy there was complete restoration of normal mucosa (Figs. 1c and 2c).

Discussion

This study clearly shows that in acrodermatitis enteropathica there is a small intestinal mucosal abnormality which is only partially corrected with DIH but which is completely restored to normal after oral zinc therapy. Since the abnormalities in the intestinal mucosa seem to improve at the same time as the skin changes during treatment with expressed breast milk and DIH, it may be that the intestinal mucosal changes are dependent on the same agency as the skin changes.

Remission of the clinical manifestations of the disease occurred in all 3 patients on DIH treatment.
but with some recrudescence of rash with intercurrent infection. All had low serum zinc levels which rose to normal after oral zinc sulphate (see Table). Thus oral zinc therapy maintains complete clinical remission, normal serum zinc levels, and normal histology of the small intestinal mucosa. However, the intestinal mucosa was still abnormal on DIH therapy. This may have been due specifically to zinc deficiency which disappeared with oral zinc and restored normal levels. The minor recrudescence of rash during intercurrent infections on treatment with DIH may have been due to an increased requirement of zinc at these times. These observations would agree with Moynahan’s postulate that all the manifestations of acrodermatitis enteropathica are due to zinc deficiency (Moynahan, 1974). The intestinal mucosa and the hair matrix have a high rate of cell multiplication and it would be likely that their morphology and function could be affected by zinc deficiency. Thus the appearance of the intestinal epithelial cells and the nuclei may reflect abnormal synthesis of nucleoprotein which is dependent on zinc-containing enzymes. Similar nuclear changes can be seen in folate deficiency (Hermos et al., 1972). Folate is essential for nucleoprotein synthesis and by analogy zinc deficiency may be the responsible factor in acrodermatitis enteropathica.

Since most previous reports of the small intestinal morphology in this disease did not show abnormality, or where found failed to revert with treatment (Ament and Broviac, 1973), further duodenal biopsy studies are indicated to show the true frequency of the intestinal mucosal abnormality in acrodermatitis enteropathica.

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REFERENCES


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