Congenital glucose galactose malabsorption

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Introduction

Congenital glucose galactose malabsorption (CGGM) is a rare autosomal recessive disorder, which presents as a protracted diarrhoea in early neonatal life. It is due to a defect in sodium coupled transport of glucose and galactose in the enterocyte (1). Diarrhoea in CGGM is osmotic, caused by accumulation of unabsorbed glucose and galactose in the intestine (2), which results in severe malnutrition (3). When glucose and galactose are eliminated from the diet, infants with CGGM thrive and dietary intervention after early diagnosis will result in normal growth and development.

Case report

The first product of a consanguineous marriage weighing 2.82 kg at birth developed a profuse watery diarrhoea on the fifth post-natal day, while exclusively breast fed. He was admitted on the 11th day and treated for hypernatraemic dehydration with intravenous fluids, and a lactose free formula was started. On day 15 he weighed 2.4 kg and was severely dehydrated and emaciated. Investigations revealed a random blood glucose 3.5 mmol/l, serum Na⁺ 151.0 mmol/l, serum K⁺ 3.2 mmol/l, haemoglobin 14.6 g/dl, haematocrit 44%, stool smear- no pus or red blood cells, reducing substances +; stool culture negative for Salmonella, Shigella and Escherichia coli. Substitution of a lactose free formula for breast milk did not have any impact and reducing substances persisted in the stool. Congenital chloride diarrhoea was excluded (stool electrolytes Na⁺ 19.0 mmol/l, K⁺ 10.86 mmol/l and Cl⁻ 27.0 mmol/l). Normal serum zinc levels excluded acrodermatitis enteropathica. At 3 weeks, when exclusively maintained on intravenous fluids the diarrhoea settled. Introduction of 5% dextrose orally resulted in recurrence of watery stools loaded with reducing substances. Then it was evident that the child was having a malabsorption syndrome rather than a secretory diarrhoea. Thereafter he was managed with intravenous 5% dextrose and fresh frozen plasma. On the 34th day, parenteral nutrition with Aminoplasmin and Lipofundin (Braun) was started. These were also given orally, and well tolerated. Thereafter, the stools became semi-solid with disappearance of reducing substances, which favoured a specific carbohydrate absorption defect. A tentative diagnosis of CGGM was made and Galactomin 19 (Cow & Gate Nutricia), which is a glucose galactose free formula, was introduced. It was well tolerated and at 13 weeks the weight had increased to 4.8 kg. Boiled and blended chicken, Casilan 90 (Glaxo), pear and apple juice (which are low in glucose and galactose) were then introduced. MCT oil (Mead Johnson) was also given and well tolerated. At 18 weeks, soup containing mashed potato, dhal, beans, dried sprats, carrots and green leaves...
was given with rice-cunjee. He weighed 5.6 kg and passed two well formed stools daily. At nine months he weighed 7.8 kg (16.8% below reference range) and the development was appropriate for the age.

Discussion

Although a definitive diagnosis of CGGM is made by demonstrating a cotransporter defect after small intestinal biopsy, dietary manipulation may suffice in establishing it (4). In this patient, although the diagnosis could not be confirmed due to lack of facilities, the diagnosis of CGGM seems certain based on the clinical presentation and the response to dietary manipulations.

Once a presumptive diagnosis is made proper dietetic management should be initiated with a diet free of glucose and galactose, with fructose as the sole source of carbohydrate. Soya based products are not useful as they have glucose or glucose polymers as carbohydrate. All medications should be free of carbohydrate. By about the 4th month, complementary foods could be introduced (1). Home made formulae based on minced chicken, egg and Casilan can be made into a mixture and adjusted to suit individual requirements (5). The infant should be monitored closely for gastrointestinal tolerance of the food, based on stool frequency and consistency. Parents should be encouraged to introduce a variety of foods gradually one at a time. After the first year carbohydrate containing foods could be introduced gradually in small quantities and the tolerance monitored. Periodically the weight gain and linear growth should be monitored. Supplementation with vitamins and minerals is also required. Alternative sweeteners such as honey may be used after 12 months. With increasing age children begin to tolerate moderate amounts of glucose and galactose in the diet as the colonic flora adapt to chronic loading of glucose by increasing fermentation of the unabsorbed carbohydrate to short chain fatty acids (1).

References