CASE REPORTS

Defective jejunal brush border membrane sodium/proton exchange in association with lethal familial protracted diarrhoea

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Abstract

The spectrum of clinical disease associated with specific defects in jejunal brush border membrane sodium/proton exchange is poorly defined and only two patients have been described so far. Jejunal brush border membrane transport studies were performed in a boy who presented with lethal familial protracted diarrhoea in the first few days of life. Using jejunal brush border membrane vesicles prepared from conventional jejunal biopsy specimens, initial sodium uptake under H+ gradient conditions was found to be only 6% of the mean control value. In contrast, sodium stimulated glucose uptake was normal. Our data confirm the importance of a congenital defect in this exchanger as a cause of severe sodium-losing diarrhoea and extend the spectrum of disorders characterised by a specific defect in brush border membrane Na+/H+ exchange to include some forms of lethal familial protracted diarrhoea.

A wide variety of disorders is responsible for severe protracted diarrhoea in infancy.1 Investigations in familial protracted diarrhoea, however, usually fail to detect a clearly defined pathophysiological cause,2 although some cases may be associated with microvillus atrophy.3,4 Congenital brush border membrane defects are now established for sodium dependent glucose transport (glucose-galactose malabsorption)5 and for the Cl/HCO3 exchanger (congenital chloride diarrrhoea).6,7 Recently, a defect in brush border membrane sodium/proton exchange was also described8 and a second patient with this disorder has been reported.9 We present a further child with a specific defect in this exchanger, who presented with lethal sodium-losing familial protracted diarrhoea in the first few days of life.

Case report

The patient, a boy, was born after a normal pregnancy and weighed 2900 g. There was no history of hydramnios and the parents were unrelated. Two previous siblings, both girls, had died of severe protracted diarrhoea at 2 and 11 months of age; both had had very high stool sodium concentrations (>80 mmol/l).

The patient passed normal meconium within 24 hours, but watery stools began on the third day of life. He was formula fed and admitted to hospital on the sixth day of life with profuse watery diarrhoea, profound dehydration, and a severe metabolic acidosis (pH: 7-02; base excess: -27 mmol/l). Parenteral nutrition was started but profuse diarrhoea persisted. At the age of 3 weeks he developed severe necrotising enterocolitis and required an ileostomy. A high output secretory diarrhoea (60–90 ml/kg/day) persisted, despite receiving nil by mouth (ileostomy fluid (mmol/l): Na 128, K 10, Cl 96) and subsequent reanastomosis (stools (mmol/l): Na 90, K 38, Cl 72). His parenteral sodium requirement was high (6–10 mmol/kg/day) and remained so after reanastomosis. A persistent metabolic acidosis was present (base excess: -6 to -20 mmol/l), but there was no evidence of a renal tubular defect. Plasma aldosterone concentrations were raised, probably secondary to persisting marginal sodium depletion.

Extensive investigation of gastrointestinal function failed to show the cause of the diarrhoea. Tests of immune function (circulating immunoglobulins, E rosette formation, mitogen stimulation of lymphocyte, chemiluminescence and autoantibodies (including antienterocyte antibody), and gastrointestinal peptides (vaso-active intestinal peptide, pancreatic polypeptide, and gastrin) were all normal. Radiological findings in the gastrointestinal tract were unremarkable, and jejunal and ileal biopsy specimens showed mild partial villous atrophy but looked normal on electron microscopy. Mucosal disaccharidase activities were somewhat reduced (IU/g protein: lactase 14–9 (reference mean (SD) 65 (45)), sucrase 15–1 (reference mean (SD) 75 (30)) probably as a result of prolonged parenteral nutrition. Enzyme histochemistry of colonic biopsy specimens showed normal nerve fibres.

Brush border membrane vesicles prepared from a jejunal biopsy specimen performed when the boy was 4 months old showed a gross defect in sodium/proton exchange, but normal sodium-stimulated D glucose transport.

The boy was maintained on home parenteral nutrition because of intolerance of all enteral feeds other than small amounts of 10% glucose or Ringer’s solution. He grew along the 3rd centiles for height and weight, but developed increasing cholestasis and died of severe intercurrent infection at 18 months.
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Jejunal brush border membrane vesicle studies

CONTROL SUBJECTS
Histologically normal jejunal biopsy specimens were obtained from two sources:
(a) Partial thickness specimens (epithelium plus submucosa), weighing 20–50 mg, were obtained per orally from five patients (aged 1 to 6 years) being investigated for gastrointestinal symptoms, but subsequently shown to have no organic disease.
(b) Full thickness specimens weighing 90–150 mg, were obtained surgically from 10 patients (aged 2 to 7 months), undergoing a portoenterostomy procedure for extrahepatic biliary atresia.

The specimens were rinsed rapidly in 0·15 M NaCl solution and stored in air tight containers at −70°C. There were no differences in the transport characteristics of vesicles prepared from the two sources.

All studies were carried out with the approval of the local research ethical committee and the written informed consent of the patient's parents.

Methods

ISOLATION OF VESICLES

Vesicles were prepared from biopsy specimens using a previously described technique.14 In the case of porceral jejunal biopsy specimens, 20–50 mg frozen mucosa were thawed in 200 μl manninitol buffer containing (in mmol/l): manninitol 300; ethylene glycol tetra-acetic acid 5, Tris/HCl 12, pH 7·1. The buffer plus mucosa was diluted sixfold by volume with ice cold deionised water and homogenised for one minute (Sorvall Mini-Omnimixer, speed 5).

When using full thickness jejunal specimens taken at operation, 100–150 mg gut were thawed in 200 μl manninitol buffer, diluted sixfold with ice cold, deionised water, and vibrated for two minutes (Vibro Mischer El, Chemap AG). After removal of any intestinal muscle with forceps, the material was homogenised as above.

Magnesium chloride was added to the homogenate to a final concentration of 10 mmol/l, and after standing on ice for 40 minutes, the specimen was centrifuged for 15 minutes at 3000 g at 4°C. The supernatant (I) was removed and stored on ice.

The pellet was resuspended in 1 ml buffer containing (in mmol/l): manninitol 60, ethylene glycol tetra-acetic acid 1, Tris/HCl 2·4, pH 7·1 and homogenised with a glass Teflon tissue homogeniser. After addition of MgCl₂ (10 mmol/l) and standing on ice for 15 minutes, the homogenate was centrifuged for 15 minutes at 3000 g at 4°C.

The supernatant (II) was added to supernatant I and the combined supernatants were centrifuged for 30 minutes at 27000 g at 4°C. The resulting pellet was resuspended in a small volume of membrane buffer (about 125 μl) by means of a syringe and 25 G needle.

The membrane suspension was equilibrated at room temperature and 4°C for 30 minutes each, before transport studies.

ENZYME ASSAYS AND PROTEIN DETERMINATIONS

Two microvillus marker enzymes were used to assess brush border membrane purification. Alkaline phosphatase (EC 3.1.3.1) activity was measured using a fluorimetric procedure15 with 0·1 mm methyl umbelliferyl dihydrogen phosphate as substrate in borate buffer, pH 8·8, containing 5 mmol/1 MgCl₂ and 0·1% Triton X-100; sucrase (EC 3.2.1.48) activity was determined fluorimetrically, using 0·028 M sucrose as substrate. The glucose product was measured by glucose oxidase-peroxidase with homovanillic acid as fluorogen.

Protein measurements were made with the Coomassie blue dye binding method,15 using the Bio-Rad protein assay kit.

Both brush border marker enzymes were measured in whole tissue homogenates and in the final vesicle preparations. Specific enzyme activities were enhanced 15 to 20-fold in the vesicle preparations.

TRANSPORT STUDIES

As a method control for confirming the integrity of brush border membrane vesicles, intact sodium stimulated D glucose uptake was confirmed in a fraction of each vesicle preparation. This requires showing that an inwardly directed sodium gradient (Na out−Na in) enhances glucose uptake, when compared with uptake without such a gradient (Na in=Na out). Brush border membrane vesicles were therefore preloaded with one of two buffers, containing either (in mmol/l): manninitol 300 and Hepes-Tris 50 (pH 7·4); or manninitol 100, Hepes-Tris 50 (pH 7·4), and NaCl 100, and incubated in a medium containing (mmol/l): manninitol 100, Hepes-Tris 50 (pH 7·4), NaCl 100, and D glucose 0·1 (+[H D glucose tracer).

To show the presence or absence of sodium/proton exchange requires proof that an outwardly directed proton gradient (pH in<pH out) enhances sodium uptake, when compared with uptake without such a gradient (pH in=pH out). Vesicles were therefore preloaded with a buffer containing (in mmol/l): manninitol 300, 2(N-morpholino)-ethane sulphonic acid (MES) Tris (pH 6·0) 50, K⁺ gluconate 20, α-1 mg/ml valinomycin, and incubated in a medium containing (in mmol/l): manninitol 300, NaCl 0·1 (+[2Na tracer], K⁺ gluconate 20, and either Hepes-Tris 50 (pH 7·4) or MES-Tris 50 (pH 6·0).

The uptake of [2Na or [H D glucose (Amersham International) by brush border membrane vesicles after 15 seconds and 90 minutes (glucose) or 15 seconds and 120 minutes (sodium) incubation, was measured using a rapid filtration technique, using 74 kBq [2Na or 185 kBq [H D glucose per time point.

Results

In brush border membrane vesicles from controls and the patient, an inwardly directed sodium gradient resulted in appreciable enhancement of D glucose uptake at 15 seconds; such that there was no significant difference in sodium stimulated D glucose uptake between controls and the patient (Figure).
**D glucose and Na\(^+\) uptake in brush border membrane vesicles from controls and in a patient with defective Na\(^+\)/H\(^+\) exchange.**

(a) D glucose uptakes in the presence and absence of an outwardly directed Na\(^+\) gradient. (b) Na\(^+\) uptakes in the presence (+) and absence (−) of an outwardly directed H\(^+\) gradient. The incubation media and membrane buffers used are given in the text. The control values are shown as means (SEM).

In contrast, brush border membrane vesicles from the patient showed no 15 seconds enhancement of sodium uptake under conditions of an outwardly directed proton gradient, when compared with no gradient; a sevenfold enhancement was seen in controls. As a result, 15 seconds sodium uptake under proton-gradient conditions was only 9 pmol sodium/mg protein in brush border membrane vesicles from the patient, compared with a mean (SD) uptake in controls of 150 (48) pmol sodium/mg protein.

Equilibrium values at 90 minutes (glucose) and 120 minutes (sodium) were not significantly different.

**Discussion**

The spectrum of clinical disease associated with the recently described specific defect in jejunal brush border membrane sodium/proton exchange is poorly defined. The clinical presentation and course of the two children published so far resembled congenital chloride diarrhoea, with a secretory diarrhoea beginning in utero with hydramnios, and a favourable prognosis. Both children grew and developed normally on appropriate electrolyte supplements.

Our patient had an equally severe and specific defect in jejunal brush border membrane sodium/proton exchange in vitro – again in association with intractable sodium-losing diarrhoea and a metabolic acidosis. A number of important clinical differences, however, were present. There was no evidence of true congenital diarrhoea, in that a history of hydramnios was absent, and the fatal outcome, strong family history, and prolonged dependence upon parenteral nutrition all suggest a different disease.

It is not clear to what extent the development of severe necrotising enterocolitis at 3 weeks of age may have enhanced any postnatal differences between our patient and those previously reported with a similar defect. The presence of normal D glucose transport in our patient’s brush border membrane vesicles does make it clear, however, that the observed changes in sodium transport did not result from unresolved non-specific mucosal damage after necrotising enterocolitis.

Our observations therefore confirm the importance of defective jejunal sodium/proton exchange as a cause of severe sodium-losing diarrhoea and extend the limited range of brush border membrane defects known to occur in association with lethal familial protracted diarrhoea.

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