Absence of acidosis in the initial presentation of propionic acidaemia

J H Walter, J E Wraith, M A Cleary

Abstract
The clinical presentation and results of the initial biochemical and haematological investigations in 11 newborn term infants with propionic acidaemia are described. All patients had neurological symptoms. Only four had clinically important acidosis, but all had a raised blood ammonia. A diagnosis of propionic acidaemia should be considered in all newborn infants with unexplained neurological deterioration even in the absence of a metabolic acidosis.

(Arch Dis Child 1995; 72: F197–F199)

Keywords: propionic acidaemia, organic acidaemia, metabolic acidosis.

Propionic acidaemia (PA) is one of the most common inherited disorders of organic acid metabolism. The disease is caused by a deficiency in the mitochondrial enzyme propionyl CoA carboxylase and leads to an accumulation in propionic acid and other metabolites. The clinical phenotype is severe and most affected children present soon after birth. Those who do not die in the newborn period rarely survive beyond their first decade. The initial presentation in newborn infants is thought almost always to be characterised by a severe metabolic acidosis and it is this finding that usually suggests the diagnosis. We report our experience in which, in most patients presenting with PA, metabolic acidosis was either mild or absent.

Methods
Between 1986 and 1994, 11 newborn infants with PA, from nine families, were both diagnosed and had their clinical care provided by our unit. The diagnosis was based in 10 cases on finding characteristic metabolites in the urine on analysis by gas chromatography/mass spectrometry and subsequently confirmed by assay of propionyl CoA carboxylase activity in leucocytes or cultured fibroblasts. One child, whose previous sibling had died from PA, was diagnosed prenatally by assay of propionyl CoA carboxylase in cultured amniocytes. All patients were of Asian origin and in most cases the parents were consanguineous.

A summary of the clinical features at presentation is given in table 1. The median age at presentation was nine days, range one to 22 days. Neurological symptoms were prominent: all but one of the patients had poor feeding or vomiting and most were lethargic. Most were jittery or had frank convulsions. Axial hypotonia was often found on examination.

A summary of the biochemical and haematological abnormalities at initial presentation is given in table 2. Serial measurements of blood ammonia, arterial pH, and bicarbonate are shown in figures 1, 2, and 3, respectively. Hyperammonaemia was found in all children. The median value for the ammonium measurement at the time of presentation was 350 μmol/l (range 84–1300). Two children had values above 1000 μmol/l. Metabolic acidosis (bicarbonate of <18 mmol/l) was present in only four children at presentation. The median pH for the group as a whole was 7·39 (range 7·11 to 7·52). The median bicarbonate was 18·5 mmol/l (range 7·4 to 22·8 mmol/l). Two patients were alkalotic and these were the children with ammonium concentrations in excess of 1000 μmol/l. Hypocalcaemia, neutropenia, and thrombocytopenia were frequent, although these were often not present initially.

Treatment
All patients were treated with intravenous dextrose and electrolyte replacement. Five required assisted ventilation. Intravenous sodium benzoate (250–500 mg/kg/day) was used to treat hyperammonaemia in eight patients. Despite theoretical concerns regarding accumulation of benzoate within mitochondria we have found this to be a safe and effective treatment in patients with propionic acidaemia who have good renal function.
Two patients required peritoneal dialysis. Seven patients were treated with either oral or intravenous L-carnitine. All patients were initially treated with biotin, the co-factor for propionyl CoA carboxylase but none showed any biochemical response. Eight patients survived their initial illness and were subsequently discharged home.

**Discussion**

The early mortality for children with propionic acidaemia presenting in the newborn period is generally thought to be very high. However, with early diagnosis and intensive care support over two thirds of our patients survived. Although most of these children subsequently have had moderate to severe developmental delay, in keeping with other reported series, two children have shown normal development in the first year of life. Both had less than 5% enzyme activity in vitro, but were not severely ill. One of these children underwent a liver transplant at 15 months with encouraging results (P J McKiernan, et al, abstract presented at 32nd Annual Symposium of the Society for the Study of Inborn Errors of Metabolism, Edinburgh, 1994). The other, whose sibling died at four weeks with severe hyperammonaemia and who was diagnosed prenatally and treated from birth, is now being assessed for liver transplantation. In order to offer children with this disorder any hope of long term survival without severe psychomotor retardation, it is important that the diagnosis is made and treatment started before severe metabolic decompensation occurs. Metabolic acidosis was not present in most of our patients at presentation. By contrast, hyperammonaemia was found in all of them. The accumulation of propionate and other acidic metabolites seems to have been sufficient to inhibit the urea cycle but not to cause clinically relevant acidosis.

Hyperammonaemia is also known to cause an alkalosis and this may have further reduced or delayed the onset of acidosis. DNA analysis had not been undertaken in these patients but their similar clinical phenotype and racial origin may indicate that they have the same mutation responsible for their disease.

Thrombocytopenia and neutropenia are known to occur in propionic acidaemia and are thought to be due to a direct toxic effect of propionic acid on bone marrow proliferation and maturation. Although most of our patients had low neutrophil (seven patients) and platelet counts (six patients) during the course of their illness, only a minority were neutropenic or thrombocytopenic at presentation. Similarly, serum calcium concentrations were low initially in only four patients but subsequently in nine.

In conclusion, the diagnosis of an organic acidaemia should be considered in all newborn infants with unexplained neurological deterioration even in the absence of acidosis. A low white cell count, platelet count, and low calcium may support such a diagnosis but may be normal at presentation. The recognition and treatment of hyperammonaemia is likely to be crucial in improving the early outcome for these patients and of particular importance if liver transplantation and, perhaps in the future, gene therapy are found to be an effective treatment for organic acidemias.


Absence of acidosis in the initial presentation of propionic acidemia

Absence of acidosis in the initial presentation of propionic acidaemia.

J. H. Walter, J. E. Wraith and M. A. Cleary

Arch Dis Child Fetal Neonatal Ed 1995 72: F197-F199
doi: 10.1136/fn.72.3.F197

Updated information and services can be found at:
http://fn.bmj.com/content/72/3/F197

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/