Discussion

Neurological complications in thalassaemia are rare. Extramedullary haematopoiesis is well recognised in thalassaemia at sites such as the liver and the spleen. Rarely such haematopoietic tissue may be found within the spinal canal, extradurally, causing compression of the spinal cord.1-3 Infrequently, infiltration of the middle ear by haematopoietic tissue may lead to hearing loss.4 These complications are more common on a low transfusion regime because it leads to bone marrow hyperplasia with widening of bones and extramedullary haematopoiesis.3 This patient was on a low transfusion regime due to poor compliance and had radiological evidence of widening of bones.

The commonest cause of an isolated unilateral lower motor neurone facial nerve palsy at this age is Bell’s palsy; however, in about 85% of cases, resolution occurs with time. Absence of evidence of middle ear infection, or a vesicular eruption around the external auditory meatus at the time of the facial palsy, excludes otitis media and herpes zoster as likely causes. Poliomyelitis or another enteroviral infection as a potential cause is unlikely in this patient as she had received appropriate doses of oral polio vaccine in the past and also because the two nerve lesions were spaced out in time. Therefore, the probable cause for the facial nerve palsy is pinching of the nerve at one of the narrow points along its course, such as within the facial canal, due to widening of diploic bone. The radiological evidence of widening of skull diploe supports this hypothesis, although an unresolved Bell’s palsy cannot be conclusively ruled out.

The reason for the phrenic nerve palsy is more difficult to explain. It may have been due to pressure from a small paravertebral mass of haematopoietic tissue. Although irradiation has been used successfully to treat paravertebral masses,2 it could not be used in this patient because exact localisation was not possible.

References


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Betaine for treatment of homocystinuria caused by methylenetetrahydrofolate reductase deficiency

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SUMMARY A 24 day old girl with homocystinuria and hypomethioninaemia caused by methylenetetrahydrofolate reductase deficiency presented with rapidly progressing encephalopathy and myopathy. An almost complete recovery was achieved by treatment with betaine.

5,10-Methylenetetrahydrofolate reductase (MTHFR) deficiency1 leads to deficient remethylation of homocysteine and is one of the causes of homocystinuria (figure). The other causes of homocystinuria are defects in cobalamin metabolism or deficient cystathionine β-synthase activity.

Methionine accumulates with homocysteine in cystathionine β-synthase deficiency,1 whereas in patients with disorders in cobalamin metabolism or with MTHFR deficiency methionine synthase is functionally deficient and the increased concentration of homocysteine occurs with normal or decreased methionine concentrations. In the latter conditions neurological impairment is the most important clinical finding. When symptoms occur in early infancy there is often rapid deterioration with respiratory failure.

Methylmalonic aciduria or macrocytic or megaloblastic changes of the bone marrow, or both, usually occur with homocystinuria in patients with disorders of cobalamin metabolism. These condi-

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tions usually respond to high doses of hydroxocobalamin. In MTHFR deficient patients, treatment with folate or cobalamin does not increase the methionine synthase activity as shown by the poor biochemical and clinical response.2

The alternative to the methyltetrahydrofolate dependent remethylation pathway of homocysteine is the betaine dependent pathway (figure). Treatment with betaine was efficient in lowering homocysteine concentrations and reduced methionine concentration to normal in a 2 year old child with MTHFR deficiency.3 The child was microcephalic and severely retarded when the betaine treatment was started, and the clinical response was limited. Two other patients treated with betaine have been reported (NJ Brandt, et al. Treatment of methylenetetrahydrofolate reductase deficiency from the neonatal period. Presented at the Society for the Study of Inborn Errors of Metabolism Annual Symposium, Amersfoort, The Netherlands, 1986). One was diagnosed prenatally. Betaine was started at the age of 2 weeks before any symptoms had appeared, and at the age of 15 months psychomotor development was normal. In the other child, treatment was started 27 days after the onset of symptoms at the age of 36 days. There was 'considerable developmental improvement' by 12 months of age. Two infants who died after 4 days and after 3 weeks of treatment have also been mentioned in a review.2 We now report an infant in whom successful treatment with betaine was started at the age of 7 weeks after a period of three weeks during which there was rapid progressive neurological deterioration.

Case report

A baby girl, the first child of healthy parents who are cousins with no family history of neurological, psychiatric, or thrombovascular disease, was born after a normal pregnancy. Vacuum extraction was used because of slow progression of the delivery. Apgar scores were 4, 5, and 8 at one, five, and 10 minutes. The birth weight was 3030 g and the head circumference 34 cm. Mild respiratory distress lead to two days treatment in an incubator. At 6 days of age the child was discharged from hospital. She was breast fed and remained healthy until 24 days of age, when she had an attack of cyanosis and unconsciousness. On admission she was pale, hypotonic, and lethargic. Blood glucose and serum electrolyte concentrations, acid base balance, and analysis of cerebrospinal fluid were normal. The haemoglobin concentration was 136 g/l. C reactive protein concentration was less than 10 mg/l.

Electrocardiogram and chest radiograph were normal. Two electroencephalograms obtained during the first week after admission were normal, although weak muscular twitching, tongue fasciculation, and seizures developed.

A positive urinary nitroprusside reaction was found two days after admission. Quantitative amino
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Acid analysis by ion exchange chromatography showed a high plasma homocysteine concentration and no detectable methionine. Increased urinary excretion of homocystine and cystathionine were also found. Urinary organic acid excretion analysed by gas chromatography and mass spectrometry was normal. The bone marrow morphology was normal. Plasma cobalamin and erythrocyte folate concentrations were also normal.

Treatment with pyridoxine (100–150 mg/day), hydroxocobalamin (1 mg/day), and folic acid (30–60 mg/day) was started. There was no response to the treatment, her clinical condition deteriorated rapidly, and respiratory failure necessitated assisted ventilation on the 21st day after admission. The electroencephalogram was then abnormal showing abundant spike and wave potentials. After three days of artificial ventilation betaine (3 g/day) and methionine (0-5 g/day) were given orally. After two days of treatment the muscular tone improved, and after two weeks there was no further need for assisted ventilation. The homocysteine concentrations in urine and plasma decreased below the detection limit and a normal plasma methionine concentration was obtained without methionine supplementation (table). The electroencephalogram returned to normal, and she caught up in growth and psychomotor development.

MTHFR deficiency was established by measurements in cultured skin fibroblasts as described by Mudd et al.4 The activity found was 0.34 nmol/(hour×mg protein). The mean activity in five controls was 15.5 nmol/(hour×mg protein) (range 9.0–32).

Total plasma homocysteine concentration—that is, the combined free and protein bound fractions, was 78 μmol/l (reference interval in adults 4.2–17.8 μmol/l) at the age of 9 months, when she was taking 3 g betaine daily. The daily intake of betaine was then increased to 6 g daily. The effect on total homocysteine concentration was limited; after two months total homocysteine concentration had only decreased to 70 μmol/l, but the plasma methionine concentration though still within the normal range, had increased.

Mainly because of persisting feeding difficulties the child was kept in hospital until the age of 5 months. When last seen at the age of 12 months the increments of length, weight, and head circumference were normal, following the 50th, 25th, and 10th percentiles, respectively. Her psychosocial behaviour was normal but her motor development was retarded by two to three months.

Discussion

Methylenetetrahydrofolate reduction is the first step in the biosynthesis of methyl groups, which—through homocysteine methylation and S-adenosylmethionine formation—participate in phospholipid biosynthesis and RNA and DNA modification, among other things.

As shown by the return to normal of plasma and urinary methionine and homocysteine concentrations in our case as in others, (NJ Brandt, et al. Treatment of methylenetetrahydrofolate reductase deficiency from the neonatal period. Presented at the Society for the Study of Inborn Errors of Metabolism Annual Symposium, Amersfoot, The Netherlands, 1986),2-3 exogenous betaine can supply methyl groups for homocysteine methylation. The accompanying clinical recovery in our case indicated that the metabolic derangement (presumably a shortage of methyl groups) responsible for the neurological symptoms was corrected by betaine treatment. Thus is imperative to recognise these patients early so as to start treatment before irreversible damage has occurred.

<table>
<thead>
<tr>
<th>Table</th>
<th>Clinical and biochemical responses to different treatments</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Age (weeks)</td>
</tr>
<tr>
<td></td>
<td>4</td>
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<tr>
<td></td>
<td>Deterioration</td>
</tr>
<tr>
<td>Treatment:</td>
<td></td>
</tr>
<tr>
<td>Pyridoxine, hydroxocobalamin</td>
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<tr>
<td>Folic acid</td>
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<tr>
<td>Methionine 500 mg/day</td>
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<tr>
<td>Betaine 3 g/day</td>
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</tr>
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<td>Plasma methionine concentration (μmol/l)</td>
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<td>Plasma homocysteine concentration (μmol/l)</td>
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<tr>
<td>Urinary homocystine concentration (mmol/mol creatinine)</td>
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</table>
To ensure long term health, betaine should if possible be given in doses high enough to return total plasma homocysteine concentration to normal, as even a moderate increase in a possible risk factor for thrombovascular disease.5

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References


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