Fracture of the femur, fish odour, and copper deficiency in a preterm infant

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SUMMARY  A preterm baby boy with blood and bone changes of copper deficiency is described. Copper deficiency was suspected after fracture of the left femur during examination of the hip joint. A low serum copper concentration (2.7 μmol/l; 17.2 μg/100 ml) and caeruloplasmin (0.04 g/l; 0.004 g/100 ml) confirmed the diagnosis. Despite the introduction of solids at 18 weeks the copper concentration remained low, and treatment with copper sulphate (2.5 mg daily) was started at 6 months. Treatment was stopped at 9 months, when he was both physically and developmentally normal.

When given a choline-containing vitamin preparation (Ketovite) he developed a fish odour because of the accumulation of trimethylamine. Withdrawal of this preparation at 6 weeks and substitution with a choline-free preparation (Abidec) was soon followed by disappearance of the odour. It is speculated that prematurity rather than copper deficiency was responsible for the poor activity of liver enzyme, trimethylamine oxidase.

Copper deficiency is rare in man, but has been reported in infants with malnutrition\(^1\) and after parenteral nutrition.\(^2-5\) Only recently has pure copper deficiency in infancy been documented,\(^3-4\) although for some time it had been known that copper deficiency occurs in a syndrome characterised by hypopcoraemia, hypoferaemia, hypoproteinœmia, and oedema.\(^5\) In previous reports copper deficiency was found after the investigation of anaemia. This is the only report where a fracture sustained during examination of the hip joint led to the diagnosis of copper deficiency.

Case report

After a normal pregnancy a 1·015-kg baby boy was born at 29 weeks by spontaneous vaginal delivery to an Asian primigravida. After transient tachypnoea he developed recurrent apnoea necessitating ventilation and intravenous dextrose electrolyte solution from day 2 to day 5, formula feeds (Cow and Gate Premium) being fully established by day 8 (150 ml/kg). Despite treatment with theophylline, apnoea continued and on day 17 he again required ventilation and intravenous dextrose electrolyte solution, maintenance requirement (200 ml/kg) of formula only being achieved by day 23. After a fall of 20% his birthweight was regained by one month.

The haemoglobin at birth was 14.5 g/dl and by 2 weeks had fallen to 8.4 g/dl, when a blood transfusion was given. Two further transfusions were required at 4 and 6 weeks. Vitamins (Ketovite) were added at 16 days and iron (Sytron) at 3 weeks. When a strong smell of fish developed Abidec (choline-free) was substituted for Ketovite and the smell soon disappeared. A subsequent challenge with Ketovite at 8 months failed to induce the odour.

At 10 weeks, during routine examination of the hip joint, a click was felt and swelling of the left leg developed soon afterwards. X-ray showed a subtrochanteric fracture, osteoporotic bones with periosteal reaction, and metaphyseal irregularity (Figure). Suspicion of copper deficiency led to the diagnosis at 15 weeks when the serum copper was 2.7 μmol/l and serum caeruloplasmin 0.04 g/l (Table). Solids were introduced at 18 weeks but

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<th>Copper (μmol/l)</th>
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<th>Protein (g/l)</th>
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<tr>
<td></td>
<td>caeruloplasmin (mg/100 ml)</td>
<td>reticulocytes (%)</td>
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Conversion: SI to traditional units—copper: 1 μmol/l = 6.37 μg/100 ml. Traditional to SI units—caeruloplasmin: 1 mg/100 ml = 10 mg/l. Protein: 1 g/l = 0.1 g/100 ml.
copper concentration remained low and at 24 weeks treatment started (copper sulphate 2.5 mg daily). The effect of treatment on serum copper and other values is shown in the Table. When treatment was stopped at 9 months he was thriving and developmentally normal.

Discussion

Copper is a known constituent of many essential enzymes and deficiency in infancy is characteristically associated with haematological and skeletal changes. Unlike the blood changes, the bone changes in copper deficiency are similar to Menkes’s syndrome (copper absorption defect), while the mental and other changes in Menkes’s syndrome are not present.

The bone changes in our patient were osteoporosis, periostitis, and metaphyseal spur formation, changes similar to scurvy and probably arising from the need for copper to convert ascorbic acid to its active form. The blood changes were anaemia, leucopenia, and neutropenia (<1.5 × 10⁹/l cells). Neutropenia is usually the first and sometimes the only manifestation of copper deficiency.¹

Previous reports of copper deficiency have affected small (under 1500 g) preterm infants, who presented at 3 months with anaemia.²⁻⁴ Three preterm infants with blood and skeletal changes of copper deficiency have been described in whom chest x-rays for respiratory problems at 3 months led to its detection. We suspect the infrequency of blood and x-ray investigations in preterm infants after 3 months partly accounts for the apparent rarity of this condition. Two-thirds of copper stores are deposited in the last trimester, making poor storage together with inadequate intake the most likely causes of deficiency. The range of copper requirement in infancy is between 200 and 500 µg/day and a reported failure⁵ to induce copper deficiency in preterm infants fed a low copper formula (15 µg/kg) is surprising, particularly as our infant received approximately 80 µg/kg (40 µg/100 ml). It should however, be noted that the infants of Wilson and Lahey⁶ received low copper for only 2 months, and that copper absorption is determined not only by copper content but also by formula composition; for example the quantity of zinc will influence the rate of copper absorption.

This is the first report of copper deficiency with fish odour. This followed the accumulation of trimethylamine after choline ingestion and the association suggested that copper might be necessary for trimethylamine oxidase, a liver enzyme preventing trimethylamine accumulation.⁷ We subsequently showed that prematurity was the probable cause of poor enzyme activity. Administration of Ketovite to 3 preterm infants of similar weights and ages induced fish odour in a single infant, who was not copper deficient.

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References

Late cerebral relapse of congenital toxoplasmosis

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SUMMARY Focal encephalitis occurred in a girl with activation of chorioretinitis which on clinical and serological grounds was taken to be caused by toxoplasma. The encephalitis and acute ocular inflammation resolved with treatment with pyrimethamine and sulphadiazine. This is presumably an example of reactivation of congenital cerebral toxoplasmosis.

Congenital ocular toxoplasmosis may be exacerbated years after the initial infection. It has been suggested that similar exacerbations of neurological toxoplasmosis occur, but these have been difficult to demonstrate. We describe an 8-year-old girl who had focal encephalitis associated with reactivation of ocular toxoplasmosis. We suggest this represents an example of reactivation of congenital central nervous system toxoplasmosis.

Case report

The patient had been born at term in eastern Kentucky after a pregnancy unmarred by maternal fever or lymphadenopathy. At 6 months strabismus had been noted, and at 3 years an optometrist diagnosed bilateral chorioretinitis. This was confirmed 2 years later by an ophthalmologist who found her bilateral visual acuity to be 20/40.

By the time she was aged 9, her school performance had deteriorated and her bilateral visual acuity was found to be 20/200. The left eye showed yellowish macular exudates with pigmentary irregularity and intraretinal oedema. A slight haze marred the overlying vitreous. Several small inactive scars lined the peripheral retina. The right eye had one large macular scar and several smaller peripheral ones, all appearing inactive. The optic discs seemed normal. The toxoplasma immunofluorescent antibody (IFA) titre was 1:64. Treatment with pyrimethamine and corticosteroids was offered but refused. She began to experience increasingly frequent headaches. Two months later she had five closely-spaced grand mal seizures. The first few were predominantly left-sided before becoming generalised. Immediately after the seizures she was afebrile, orientated, and co-operative. Her neck was supple and there were no focal neurological signs. The eye findings were similar to those described above. CSF obtained at lumbar puncture was under normal pressure but contained 100 nucleated cells (all lymphocytes) /mm³, with a glucose concentration of 50 mg/100 ml (2-8 mmol/l), protein concentration 19 mg/100 ml, and IgG concentration of 2-8 mg/100 ml. Bacterial and fungal cultures were sterile and cryptococcal antigen was absent. Her serum toxoplasma IFA titre was 1:256. An EEG 18 hours after the last seizure showed right posterior temporal 0-5 to 1 Hz slow wave activity, which persisted in a second EEG 4 days later. Skull and chest x-rays, and CAT of the head were normal. Serum VDRL and FTA were non-reactive, serum rubella titre was less than 1:10, serum herpes titre was less than 1:8, and cytomegalovirus titre was positive only to 1:8 dilution. The girl's mother had a serum toxoplasma IFA titre of 1:256.

The child was treated with folic acid, sulphamethazine, and pyrimethamine and sulphasalazine. She showed marked improvement. Computer-assisted tomography of the head was normal. She has been followed up for 5 years and her vision remains stable. The principal problem has been learning difficulties, which have been improved by refresher classes. Her neurological examination is normal.

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