Neonatal cholestasis and focal medullary dysplasia of the kidneys in a case of microcephalic osteodysplastic primordial dwarfism

A Berger, N Haschke, C Kohlhauser, G Amman, U Unterberger, M Weninger

Abstract
We report on a male infant who presented with intrauterine growth retardation, severe postnatal failure to thrive, microcephaly, facial dysmorphism, and skeletal dysplasia. The clinical and radiological findings are consistent with former descriptions of microcephalic osteodysplastic primordial dwarfism (MOPD) type I/III. In addition to previously published features, multiple fractures of the long bones, severe neonatal cholestasis, and histological dysplasia of the kidneys were found. The boy died at the age of 8 months. The new finding of focal renal medullary dysplasia further supports the hypothesis of a basic defect in tissue differentiation in the pathogenesis of this rare condition.

Keywords: microcephalic osteodysplastic dwarfism; bone fracture; neonatal cholestasis; focal medullary dysplasia of kidney

Based on the thorough review of “bird headed” dwarfism by Majewski and Goecke, a classification into four different types was established as follows: classical Seckel syndrome and MOPD types I, II, and III, the latter being distinguished from Seckel syndrome by distinct radiological features. According to recent clinical observations, there is now general agreement to consider types I and III as a single entity.

In this report, detailed clinical, radiological, and histological data of an affected infant with MOPD type I/III are presented. Some of our findings have not been described previously.

Case report
We report on a male infant, delivered at 31 weeks’ gestation by caesarean section because of preterm labour associated with abnormal CTG tracing. The mother of our patient was a 26 year old Turkish woman, married to her 23 year old first cousin. An older sister developed normally. Repeated sonographic examinations from 21 weeks of gestation onwards showed severe intrauterine growth retardation. Amniocentesis performed at 23 weeks showed a normal karyotype and normal amniotic alphafetoprotein.

Birth weight was 786 g (<3 SD), length 32 cm (<3.4 SD), and head circumference 21.7 cm (<6 SD). APGAR scores were 5 and 7 at one and five minutes, respectively. The child had mild respiratory distress syndrome requiring ventilatory support for only one hour.

The following abnormalities were noted (fig 1): microcephalic dwarfism with a sloping forehead, small anterior fontanelle, sparse eyebrows, large eyes, beaked nose, thick lips, moderate micrognathia, short neck, broad hands with marked tapering of fingers, clinodactyly of the 5th fingers, short and broad feet with prominent heels, bilateral cryptorchidism, and dry, slightly hyperkeratotic skin.

On cranial ultrasound, partial agenesis of the corpus callosum and minimal cortical gyral differentiation were noted. The renal tract and heart were normal on ultrasound. A normal male karyotype was confirmed.

Thorough investigations for intrauterine infections and metabolic and endocrine disorders were negative.

The clinical course was complicated by recurrent infections from the first week of life onwards. The differential count was extremely shifted to the left (immature to total neutrophil ratio up to 0.9), even in intervals without clinical signs of infection. No specific immunological disorder was found. Bone marrow aspiration of the tibia was unsuccessful. The procedure was not repeated because of generalised osteoporosis and subsequent multiple fractures.

Besides recurrent infections, the most striking clinical findings were hepatosplenomegaly and cholestasis. Repeated serological and metabolic investigations were negative and a biopsy of the liver showed non-specific pathology (giant cell hepatitis, fibrosis, cholestasis) without evidence of storage disease.

Growth and weight gain were poor and remained far below the 3rd centile; psychomotor development was severely retarded.

At the age of 7 months the boy developed persistent hypokalaemia, hypophosphataemia, and hypocalcaemia with urinary loss of protein, glucose, potassium, calcium, and phosphate. Despite repeated courses of antibiotic treatment the child died of pneumonia and sepsis at the age of 8 months.

**Radiological findings**
There was microcephaly and generalised osteoporosis. The spine showed mild platyspondyly. The pelvis was dysplastic with rounded, hypoplastic iliac wings and nearly horizontal, irregular acetabular roofs (fig 2A). Both femora showed enlarged proximal metaphyses (fig 2B). There was a total lack of epiphyseal ossification.
Figure 1  The proband aged 2 months (A, B) and 8 months (C). Note microcephaly, beaked nose, tapering of fingers, dorsal lymphoedema of the feet, and hypoplastic genitalia.

At the age of 7 months the first bone fracture was noted on routine x-ray. Following the initial proximal femoral location, numerous additional lesions involving the distal femoral, proximal tibial, and proximal humeral bones and the ribs were observed (fig 2C). Post-mortem studies confirmed these findings.

Necropsy macroscopic findings
The spongiosa of the long bones was hypoplastic and porotic. Massive hepatosplenomegaly was noted. The liver showed fibrosis and cholestasis. There was cardiac dilatation and marked gas distension of the gut. The kidneys were small with icteric nephropathy. The thymus appeared involuted.

The brain was small (118 g) and abnormally shaped with hypoplastic frontal lobes. Gyration was plump but symmetrical. Partial agenesis of the corpus callosum (fig 3A) and a synthalamus were found. The cerebellar vermis was hypoplastic, with distorted configuration.

Figure 2  Radiographs of patient at birth (A), 4 months (B), and 7 months (C). (A) Note dysplastic pelvis with nearly horizontal acetabular roofs and general lack of epiphyseal ossification. (B) Note enlarged proximal femoral metaphysis and proximally long fibula. (C) Note severe generalised osteoporosis and fractures of the left proximal humerus, both proximal and distal femoral bones, and proximal tibial bones.
A case of microcephalic osteodysplastic primordial dwarfism

blastic activity. The marrow cavity was enlarged and filled with haematopoietic cells. Irregular intracartilaginous ossification was seen with diminished height of cartilage cell columns and disturbed ossification.

**Brain**

The cerebral cortex was abnormally wide with a slightly disturbed lamination pattern. There were heterotopic neurones in the molecular layer of the cortex as well as irregular clusters of neurones in the white matter. Overlying the cortex was a layer of heterotopic gialle neuronal tissue (fig 3B), predominantly on the medial surface and dorsal convexity of the hemispheres. The hippocampus was malrotated. Cerebellar folia were dysplastic in some areas with multiple micropolygyric convolutions.

**Discussion**

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delineated three types of microcephalic osteodysplastic primordial dwarfism differing from Seckel syndrome on the basis of radiological findings. Types I and III are now considered to be a single entity with a possible age dependency of the x-ray findings. It was suggested that the term “type III” should be abandoned altogether, leaving only two types, type I, including all cases formerly reported as type I or type III plus the cases published as “cephaloskeletal dysplasia” and type II. Consequentially, the term “type III” became free. Confusingly, in 1992, Majewski again used this term to describe the syndrome of Caroline Crachami, a “new” entity of osteodysplastic primordial dwarfism.

Like previous cases, our patient died during an episode of respiratory infection. We observed extremely immature cells in the peripheral blood, even in intervals without clinical infection. Taybi and Linder reported a similar shift to the left in the differential of their second case. However, the reported blood count was taken shortly before death from pneumonia. Unfortunately, an attempt at bone marrow aspiration showed only peripheral blood cells. Based on reports of infections as the primary cause of death in most cases that survived for a longer time and supported by excessive extramedullary haematopoiesis involving the liver, spleen, kidneys, and testes (postmortem findings) in our case, we suggest that abnormalities in haematology and perhaps immune response, although not confirmed in our case, are part of this syndrome. In future, bone marrow examination at an earlier stage of disease could provide further important information.

A feature not formerly described was the marked cholestasis and hepatosplenomegaly of our patient. No other reported case of MOPD has shown this pathology. Taybi and Linder reported jaundice without hepatosplenomegaly. In our case, histological examination showed no specific alterations. In addition, repeated serological studies were not conclusive, leaving only parenteral nutrition and recurrent infections as possible explanations for this clinical finding. Only recently, Eason et al. reported the previously unrecognised feature of renal tubular leakage in a case of
Autosomal recessive inheritance is a common pattern in the study of bone proliferation and tissue differentiations. While necropsy was not performed in the case of Eason et al., Winter et al. described focal dilatation of the proximal renal tubules and fibrin deposits occluding the capillaries of a few glomeruli in their patient. With respect to these rather non-specific findings, the presence of focal medullary dysplasia in our case appears to be of particular interest, as it provides additional evidence that a basic defect of tissue differentiation might be primarily involved in the pathogenesis of this condition.

Multiple fractures of the long bones and ribs as observed in our case have not been described previously. We assume that they result from the renal problems and the primary disease. Eason et al. also described marked osteoporosis in their case. Histology of the bone was similar to previous descriptions. Our neuropathological findings showing a profound disturbance of cerebral development are similar to former reports. The clinical variability of this disorder seems to be broad; additional findings in our patient were neonatal cholestasis and multiple fractures of the long bones. The pathogenesis is unknown, but we propose a basic defect in cell proliferation and tissue differentiation. This hypothesis is further underlined by our finding of focal medullary dysplasia of the kidneys. Autosomal recessive inheritance is likely, based on reports of parental consanguinity (our case) and incidence among sibs of different sex. The gene responsible remains to be defined.

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