Liver histology in the arthrogryposis multiplex congenita, renal dysfunction, and cholestasis (ARC) syndrome: report of three new cases and review

S P Horslen, O W J Quarrell, M S Tanner

Abstract
We report three cases from two unrelated families of infants with arthrogryposis multiplex congenita, cholestatic jaundice, and renal Fanconi's syndrome. In both families the parents were consanguineous. All three children died by 7 months of age. This association was first reported in 1973 by Lutz-Richner and Landolt and again in another family by Nezelof et al in 1979. However, because of differing liver histology the two sibships were considered to have two separate conditions. Based on the histological findings in one of our cases we propose that all cases described so far represent variation within a single syndrome.

The association between arthrogryposis multiplex congenita, cholestatic liver disease, renal tubular acidosis, and death in infancy has been reported previously. Since the first report by Lutz-Richner and Landolt in 1973, four other similarly affected families have been reported. The cases described so far have been divided into two separate groups on the basis of hepatic histology. The first group has paucity of intrahepatic bile ducts and giant cell transformation of hepatocytes. The other condition has pigment deposition in liver cells and marked cholestasis. We describe three further cases from two families and show hepatic histological features in one of the cases compatible with both groups.

Case reports

CASE 1
Case 1, the first born child of healthy, consanguineous Pakistani parents, was born at 41 weeks' gestation after a normal pregnancy and delivery, weighing 2575 g. He was noted to have low set ears and arthrogryposis of his lower limbs which consisted of bilateral calcanoevalgus, fixed flexion deformities at the hips and knees, and dislocated hips (fig 1).

On the third day after birth he developed profuse diarrhoea and a metabolic acidosis which persisted even after the diarrhoea had resolved. At 7 days a conjugated hyperbilirubinaemia was noted.

Fanconi's syndrome was diagnosed based on the demonstration of renal tubular acidosis, glycosuria in the absence of hyperglycaemia, generalised amino aciduria, and gross phosphaturia with a fractional phosphate excretion of 0-63 (normal <0-15).

He was also noted to pass very large quantities of dilute urine in spite of plasma hypertonicity. Hypernatraemia was particularly troublesome when sodium bicarbonate was used to treat the renal tubular acidosis. Urine osmolality was greater than 310 mmol/kg. Nephrogenic diabetes insipidus was confirmed by administering a parenteral dose of desmopressin (400 ng) which failed to produce a rise in urine concentration. Treatment with chlorothiazide (10 mg/kg three times daily) did not produce a decrease in urine output, but it did increase the urinary sodium losses enabling adequate treatment of the metabolic acidosis with sodium bicarbonate. Total fluid requirements remained at 300 ml/kg/day.

Cholestasis persisted. Metabolic and infective causes of neonatal liver disease were excluded and there were no features to suggest Alagille's syndrome. Cytogenetic analysis was also normal. Abdominal ultrasound showed normal liver and bile ducts. However, there was no excretion of technetium labelled methylbromo-iminodiacetic acid into the duodenum, after intravenous injection, up to 24 hours later suggesting biliary obstruction. Percutaneous liver biopsy showed cholestasis, lipofuscin deposition, occasional multinucleate forms, and paucity of intrahepatic bile ducts confirmed with immunohistochemical staining for cytokeratin which specifically identifies bile duct epithelium (fig 2). Fat malabsorption continued despite feeding with a formula feed high in medium chain triglycerides and the addition of ursodeoxycholic acid (15 mg/kg).

With this supportive treatment and dietary supplementation, consisting of glucose polymer and medium chain triglyceride (MCT) oil, he began to gain weight and remained biochemically stable. However, at the age of 7 months he developed a chest infection to which he succumbed. Negative bacterial cultures, a lymphocytosis, and lack of response to broad spectrum antibiotics suggested a viral aetiology. Consent for necropsy was withheld.

CASE 2
Case 2 was a male born at 42 weeks' gestation to healthy, consanguineous Asian parents. The pregnancy was complicated by a raised α fetoprotein, but amniocentesis was refused. Birth

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Liver histology in the arthrogryposis multiplex congenita, renal dysfunction, and cholestasis (ARC) 63

Figure 1  Case 1 aged 4 months.

weight was 2750 g and Apgar scores were 8 and 8 at one and five minutes respectively. The placenta was grossly infarcted.

At birth the baby had loose skin, a high arched palate, low set ears, pectus carinatum, and arthrogryposis consisting of adducted hips, dorsiflexion of both feet at the ankles, and abnormalities of the right arm and hand. There was loss of muscle bulk in the arms and legs and skeletal survey showed hip dysplasia.

Obstructive jaundice was noted on day five. Abdominal ultrasound showed a liver of normal echogenicity, the intrahepatic bile ducts were not dilated, and an IDA excretion scan showed no evidence of bile flow. On day seven a metabolic acidosis was noted and the infant had evidence of Fanconi's syndrome with generalised amino aciduria, glycosuria, and hyperchloraemia.

An echocardiogram showed a small secundum atrial septal defect. No abnormality was found on cerebral ultrasound or detailed ophthalmological examination. Cytogenetic analysis was normal. Metabolic and infective causes of liver disease were excluded.

This infant received supportive treatment but died at the age of 7 months. There was no necropsy.

CASE 3

A male sib of case 2 was born at term by elective caesarian section. Serial antenatal ultrasound scans had suggested abnormalities in the third trimester with evidence of oligohydramnios and arthrogryposis. Apgar scores were 6 and 9 at one and five minutes respectively and birth weight was 2950 g. At birth he had contractures of the hips and knees and talipes of the feet and ankles and muscular calcification was noted on radiographs. The ears were described as large but low set. He had a small chin with full lips.

By day four he had developed metabolic acidosis and jaundice. The acidosis was considered to be renal tubular in origin and was associated with generalised amino aciduria. IDA excretion did not occur, suggesting biliary obstruction. He also developed right parotitis. A decision was made to treat him symptomatically and he died at home aged 1 month. Again, necropsy was not undertaken.

Discussion

In 1973 Lutz-Richner and Landolt1 reported two sibs of consanguineous parents who had arthrogryposis, hepatobiliary disease, and renal dysfunction. Eight further cases have been reported to date (table). In the second report of a sibship with this association of features, Nezelof et al7 suggested X linked inheritance for the condition. However, the occurrence of affected male and female sibs in later case reports, together with parental consanguinity, suggests autosomal recessive inheritance as the most likely mechanism.5 4

All reports which looked at the neuromuscular pathology have concluded that the deformities were the result of neurogenic muscular atrophy with rarification of the motor neurones of the anterior horns of the spinal cord.4 4 Also, identical renal pathological findings of renal tubular degeneration with nephrocalcinosis have been described in previous case reports.14 However, Lutz-Richner and Landolt1 and Mikati et al2 describe paucity of intrahepatic bile ducts and multinucleate transformation of hepatocytes in their patients, whereas the findings of Nezelof et al7 and subsequently Di Rocco et al6 and Saraiva et al6 were of Dubin-Johnson-like pigmentary changes with cholestasis. In view of these differing histological findings, Di Rocco et al6 suggested that two separate conditions were
Major features of infants with arthrogryposis multiplex, cholestatic liver disease, and renal impairment described in published reports.

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Cases 1 and 2 Lutz-Richner and Landolt,3 4, 5, and 6 Nezelof et al,7 and 8 Mikati et al,9 Di Rocco et al,10 Saraiva et al,11 12, and 13 present report. + = present, − = absent, NK = not known.

being described. However, the hepatic pigment in Dubin-Johnson disease has the staining properties of lipofuscin and the findings of cholestasis, multinucleate cells, lipofuscin deposition, and intrahepatic biliary hypoplasia in our first patient suggest that there is overlap between all the cases described so far. Both bile duct paucity and lipofuscin deposition can be seen in a wide range of liver disease and both features probably represent non-specific changes resulting from a variety of insults to the liver. Similarly, a common insult may produce both lesions but the degree of each may vary from person to person and may depend on the site from which the liver biopsy is taken and the stage in the evolution of the pathological process at which the specimen was obtained. We propose that all cases represent the same condition. Although the genetic mechanism is clear, the underlying metabolic abnormality is unknown. Although the recent reports have spoken of this phenotype as ‘Nezelof’s syndrome’, we suggest that this term is inappropriate as this eponym is already used to describe a form of thymic immunodeficiency; instead we suggest the acronym ARC syndrome.

We would like to acknowledge Dr R Mueller for first making the diagnosis in case 1 and Mr Anderton and Dr Murphy for referring the patients to us.

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