A family with spondyloepimeta physeal dwarfism: a ‘new’ dysplasia or Kniest disease with autosomal recessive inheritance?

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SUMMARY We present an Arab family with some features of Kniest disease. The proband was a six year old boy with rhizomelic short limbed dwarfism, ‘dish-like’ facies, cleft palate, deafness, and camptodactyly. Most radiological changes were compatible with Kniest disease. Two younger sibs, similarly affected, had died at a few months old, and the pedigree shows strong evidence of autosomal recessive inheritance, unlike previously reported cases of Kniest disease which have shown autosomal dominant inheritance.

There are many forms of dwarfism associated with spinal as well as epiphyseal and metaphyseal changes seen in radiographs. However, the most striking feature in this family was the enormous enlargement of the metaphyseal region of the long bones in infancy, which is one of the predominant radiological features of Kniest disease, first described by Kniest in 1952, and subsequently discussed in relation to similar disorders by other authors.3–5

Case reports

CASE 1
The proband (fig 1, III.5) was an intelligent boy born at term after a normal pregnancy and delivery. He had four normal sibs and two younger affected ones. His Syrian parents were phenotypically normal. He was referred to the Genetics Clinic because of disproportionate short stature diagnosed as either atypical achondroplasia, Crouzon syndrome, or Kniest disease. His height and weight were below the 3rd centile, with marked rhizomelic shortening in both upper and lower limbs (fig 2). He had a

![Family pedigree](image1)

![The proband (III.5) showing short limbs, particularly the proximal portion, and flattened face.](image2)
FIG 3 The dumb-bell shaped long bones of (a) patient III.9 and (b) patient III.10 in the first few months of life.

'dish-like' facies with frontal bossing, hypertelorism, depressed nasal bridge, anteverted nostrils, a long philtrum, very narrow external auditory meatus, bilateral moderate mixed hearing loss, a repaired cleft palate, and camptodactyly. The serum calcium, phosphorus, magnesium, and alkaline phosphatase were all normal. The child has been followed up for three years, with no essential change in his condition.

Clinical, radiological, and ophthalmological assessment of the parents showed them to be phenotypically normal.

CASE 2
A younger sister (III.9) was delivered normally at term. She was first examined at two months of age. Her weight and height were below the 3rd centile and she had marked rhizomelic shortening with a similar facial appearance to her brother, together with a posterior cleft palate and camptodactyly of

FIG 4 Narrow chest and mild platyspondyly seen in the early months.
the little finger. Failure to thrive and recurrent gastrointestinal and chest infections were major problems and she died outside Kuwait at four months of age after severe gastroenteritis.

CASE 3
A younger brother (III.10) was delivered by caesarean section for brow presentation and had a birth weight of 4010 g, with Apgar scores 5 and 8. He also had marked rhizomelic shortening, 'dish-like' facies, a posterior cleft palate, and camptodactyly of the little finger, but in addition he had micrognathia and glossoptosis. He too had recurrent chest infections and failure to thrive. Haematological, biochemical, and immunological tests were normal. He died at eight months of age after fulminating bronchopneumonia.

RADIOLOGICAL FINDINGS
The three patients described above all showed similar findings. The spine showed a lordosis in the cervicothoracic region instead of the normal kyphosis and only mild generalised platyspondyly with anterior wedging of the thoracolumbar vertebrae, together with posterior vertebral scalloping in the lumbar region. The long bones were all shortened with striking enlargement of the metaphyses and loss of bone moulding, particularly during the first year of life (figs 3 and 4). This was less marked in the six year old (figs 5, 6, 7, 8).

Discussion
The disorder appears to be of autosomal recessive
inheritance; parental consanguinity was not a known feature, but is nevertheless likely in a country such as Kuwait with a very high consanguinity rate (54.3%), high average inbreeding coefficient (F=0.0219), and large family size.6

The predominant clinical and radiological findings were short limbed dwarfism, characteristic facies, deafness, cleft palate, camptodactyly, and striking radiographical changes both in the spine and long bones. Gross enlargement of the metaphyses, as seen here, in the early months of life is a feature of hereditary arthro-ophthalmopathy (Stickler syndrome), as is the deafness, 'dish-like' facies, and mild platyspondyly.7 However, the Stickler syndrome is predominantly an epiphyseal disorder accompanied by a high degree of myopia, retinal detachment, and blindness, none of which was present here.

Metatropic dysplasia has similar dwarfism and radiographical changes in the long bones, but here the platyspondyly is more severe with 'paper thin' vertebrae.8 Kniest disease is the more likely diagnosis, with its characteristic facies and mild platyspondyly. However, Kniest disease is usually described as being of autosomal dominant inheritance, although in fact very few cases are known and there could well be a dominant type.

The lateral spine radiograph of the older child has characteristics of classical achondroplasia, with its short pedicles, posterior scalloping, and wedged vertebrae at the thoracolumbar junction, but other clinical and radiographical features do not suggest this disorder.

It is not possible to make a firm diagnosis with certainty here, but if the disorder is not Kniest disease with autosomal recessive inheritance, then it is likely to be a 'new' spondyloepimeta physeal dysplasia.
A family with spondyloepimetaphyseal dwarfism

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References


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