Humeroradioulnar synostosis in a patient with lambdoid synostosis

T J C Edwards, E A Haan, I J Humphrey

Abstract

We report on a patient with humero-radioulnar synostosis and lambdoid synostosis. The case differs from three previously described cases in minor details, but the upper limb abnormalities are strikingly similar. (J Med Genet 1993;30:81-2)

Cases of upper limb oligoelectrosoyndactyly have previously been described by Hersh et al,1 Leroy and Speeckaert,2 and Gollop and Coates.3 The cases described by these authors all exhibited bilateral upper limb deformities. In addition, two of the three cases showed mild plagiocephaly. The infant described in this article was born with a unilateral upper limb deformity that was similar to those previously described, along with plagiocephaly owing to right lambdoid craniosynostosis.

Case report

The proband was the first child of non-consanguineous parents. He was born at term after a normal pregnancy, labour, and delivery with a birth weight of 2273 g. The mother was of Maori and the father of European extraction.

The child was referred at 7 months of age for assessment of plagiocephaly with right occipital flattening. Weight (5-62 kg) and head circumference (41-5 cm) were below the 3rd centile. Both upper limbs were abnormal (fig 1). On the right there was a dimple in the skin of the upper arm overlying a bony exostosis from the medial side of the midshaft of the humerus. There was no elbow joint and the forearm bones were absent. There were three digits, a rudimentary thumb attached by a skin tag and two other digits joined along their lengths by soft tissue (fig 2). On the left, the upper arm, elbow, forearm, and wrist were normal but the thumb was hypoplastic. The facies, the ears, anal placement, cardiovascular examination, and developmental assessment were normal.

Radiographs showed premature synostosis of the right lambdoid suture, an abnormally large anterior arch of the atlas, and fusion of the posterior elements of C2 and C3. The bases of the left fourth and fifth metacarpals were fused and all elements of the left thumb were hypoplastic. The right humerus was abnormally long with a 3 cm exostosis arising medially from the junction of the middle and distal thirds. The radius and ulna were absent. An ossification centre was present in the carpal region and appeared to be composed of two fused elements. There were two complete rays (metacarpals and phalanges) and a rudimentary thumb with two phalanges (fig 3). Radiographs of the lower limbs were normal.

Discussion

The three cases reported previously1-3 had almost identical limb deformities to the case presented here, the only difference being that in the case presented here the limb deformity is unilateral, whereas the previous three have shown bilateral deformity. The association of radial aplasia and craniosynostosis led us initially to entertain the diagnosis of Baller-Gerold syndrome. However, the very mild degree of craniosynostosis and the striking...
similarity of the limb deformity in these four cases suggested that this case is more closely aligned to the humeroradioulnar synostosis group. The most intriguing factor of all four cases is the apparent bifurcation of the long bone of the arm. The pathogenesis of this is unclear.\(^1\)

This case therefore represents a variation of the three previously reported in that the upper limb defect is essentially unilateral and the plagiocephaly is represented by lambdoid synostosis. While this case broadens the spectrum of this entity it fails to shed any light on its aetiology.

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