Lethal osteogenesis imperfecta associated with 46,XY,inv(7)(p13q22) karyotype

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SUMMARY An infant who died of complications of osteogenesis imperfecta (OI) at 22 days of age had a 46,XY,inv(7)(p13q22) karyotype. His mother carried the same inversion. One breakpoint of the inversion was within the region of the gene for α2(I) procollagen. The product of this gene is a component of type I collagen, the principal collagen synthesised by osteoblasts. Karyotypic abnormalities involving type I collagen gene sites have not previously been reported in association with OI.

Case report

A 24 year old, gravida 2 woman presented in the 26th week by dates of gestation for sonographic evaluation of fetal age. The limbs of the fetus were shortened and bowed and the thoracic diameter was decreased. Short limbed dwarfism was diagnosed. Labour began spontaneously at 38 weeks and a caesarean section was performed. The limbs of the infant (fig 1), a male, were severely deformed. Caput membranaceum was present. The facies was rounded, with a fine, pointed nose, and the sclerae were white. Other findings included a single transverse palmar crease on the left and a complexly abnormal palmar pattern on the right (fig 1). X rays showed a radiolucent skull and spine, beaded, irregular ribs, and thick, crumpled long bones (fig 2). Osteogenesis imperfecta (OI) was diagnosed. Supportive care without oxygen supplementation or ventilatory assistance was provided. The infant died at 22 days of age after several hours of tachypnoea with increasing cyanosis. Necropsy showed many rib fractures that bowed the chest wall inward. The lungs weighed 48 g (expected normal weight 65±17 g†). No pneumonia was found. Death was attributed to respiratory insufficiency owing to pulmonary hypoplasia, possibly aggravated by chest wall instability. Fibroblasts cultured at necropsy did not grow.

Both parents, recent immigrants from the same town in the Cape Verde islands, were of black

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FIG 1(a) Photograph of face and anterior torso. The habitus is that frequently encountered in lethal osteogenesis imperfecta.
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Portuguese background; consanguinity was denied. Their first child, a male, was three years old and well. None had signs or symptoms of OI. All had normal palmar crease patterns. No history could be obtained of osteochondrodystrophy in the family of either parent.

Giemsa banded preparations of cultured peripheral blood lymphocytes from the infant showed a 46,XY,inv(7) karyotype. The inversion was pericentric with breakpoints at p13 and q22 (fig 3). Similar preparations of peripheral blood lymphocytes from the mother showed a 46,XX,inv(7)(p13q22) karyotype. Preparations from the father showed no chromosomal abnormalities. Permission to draw blood from the sib was refused. No other family members were available for study.

FIG 1(b)  Photograph of right palm. The crease pattern is complexly abnormal.

FIG 2(a)  Radiograph (mammography technique) of anterior portion of membranaceous calvarium, showing innumerable minute Wormian bones.

FIG 2(b)  Radiograph of chest. The ribs are broad and show many fractures.
FIG 2(c) Radiograph of pelvis, legs, and feet. The multiple fracture sites and resultant deformities are evident.

FIG 3(a) Diagram of G banding pattern for normal chromosome 7; arrows indicate breakpoints. (Reproduced with permission from fig 9 in Harnden DG, Klinger HP, eds. ISCN 1985. An international system for human cytogenetic nomenclature. Basel: S Karger, 1985: 53. (b) Photomicrographs of chromosome 7 (right) containing pericentric inversion and of karyotypically unremarkable chromosome 7 (Giemsa, 500 band level). Arrowhead indicates p13 breakpoint, arrow indicates q22 breakpoint.

Discussion

Type I collagen, a heterotrimer containing two α1(I) procollagen molecules and one α2(I) procollagen molecule, constitutes approximately 95% of the collagen produced by osteoblasts.2 The gene for α2(I) procollagen has been assigned to the long arm of chromosome 7 at a site between bands q21 and q22.3 This region may have been disrupted by the chromosomal inversion found in the infant and his mother.

Abnormal karyotypes have been reported in only four other cases of OI. One infant, with features of chondrodystrophy and a karyotype of 46,XY,−5, −D,t(5q+:Dq−)+;cen+, died during delivery; his tissues were so fragile that his limbs were avulsed. Because the karyotype was unbanded, the chromosomes involved could not be identified precisely.4 Another patient with severe disease who died in infancy had an unbanded karyotype of 46,XX(92%)/
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47,XX,+G(8%). This low percentage mosaic state for a small acrocentric chromosome was probably coincidental. A karyotype of 46,XX,r(18)(p11q23), also probably an unrelated finding, was found in a third infant with lethal OI. Finally, a child with less severe disease had a karyotype of 46,XY,del(12)(p12p13). His father, who had a normal karyotype, also had moderately severe OI, indicating that OI in the child was probably unrelated to the partial deletion. We know of no previous report of OI associated with karyotypic abnormalities involving type I collagen gene sites.

Damage to one α2(I) procollagen allele caused by the inversion might have contributed to disease in the infant if a mutation affecting the other allele was present. Such a mutation might either have occurred de novo or have been inherited from the father; compound heterozygosity for abnormalities in the α2(I) procollagen gene, one of which was phenotypically recessive, has previously been reported in lethal 'broad boned' OI. It is also possible that a different de novo mutation caused OI in the infant by affecting one α1(I) procollagen allele and that the observed inversion was not related to his disease. The risk that other children born to these parents will have OI is increased only if each parent is heterozygous for an α2(I) procollagen gene abnormality. Attempts are now in progress to characterise α2(I) procollagen genes and gene products in the parents for purposes of genetic counseling.

References


Absence of a vagina and right sided adnexa uteri in the Waardenburg syndrome: a possible clue to the embryological defect

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SUMMARY An 18 year old single Jewish woman with the Waardenburg syndrome and absence of a vagina and right sided adnexa uteri is reported. Other congenital malformations associated with the Waardenburg syndrome are mentioned and it is postulated that they may be the result of an altered invasion of neurones or altered neurones in certain organ systems early in embryogenesis.

Over the years our group has described various congenital malformations associated with the Waardenburg syndrome. Recently, we had the opportunity to re-evaluate one of our patients that we had seen many years ago, and to our surprise we learned that she was born without a vagina and right adnexa uteri. The purpose of this brief report is to discuss the above observations in relation to other hypoplastic or aplastic congenital malformations seen in the Waardenburg syndrome.

Case report
An 18 year old single Jewish woman was referred to

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