Case reports

Trisomy 13 in monozygotic twins discordant for major congenital anomalies

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SUMMARY The occurrence of trisomy 13 in twins is very rare. We report a pair of genotypically identical twins with trisomy 13 discordant for major anomalies. This case contributes to the already published data on the contribution of non-genetic factors to the aetiology of congenital malformations in monozygotic twins.

Bartolin in 1657 was the first to describe the pattern of anomalies which in 1960 was identified by Patau et al as being caused by the extra chromosome in the D (13 to 15) group. Since then, numerous reports have been published focusing on the wide range of clinical features in this syndrome.\(^2\)\(^3\) We report the occurrence of trisomy 13 in twins who were identical genotypically and yet different phenotypically.

Case report

Male twins were spontaneously delivered at 32 weeks of gestation to a 38 year old gravida 8 mother and a 45 year old father. Amniocentesis had been refused on religious grounds. No hereditary

FIG 1 Major malformations in twin A: (a) bilateral cleft lip and palate, and (b) left diaphragmatic hernia.

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chromosomal disorders were noted in the family. Twin A, a breech presentation, was born with an Apgar score of 2 at one minute and weighed 1800 g. Because of respiratory difficulty he was immediately intubated and ventilated. Twin B, a vertex presentation, had Apgar scores of 5 and 8 at one and five minutes and weighed 1850 g. He was given oxygen and was intubated and ventilated at two hours of life for deteriorating respiratory distress. The initial physical examination of twin A showed trigonocepha- lty, dysplastic low set ears, microphthalmia, severe bilateral cleft lip and palate (fig 1a), postaxial polydactyly of the left hand, a simian line on the left hand, hyperconvex fingernails, syndactyly of the left second, and third toes, bilateral rockerbottom feet, and micropenis. Chest x ray showed left dia- phragmatic hernia (fig 1b) with the heart displaced to the right. Echocardiographic studies showed a structurally normal dextroposed heart. Twin B had dysplastic low set ears, microphthalmia, postaxial polydactyly of the left hand, a left simian line, hyperconvex fingernails, syndactyly of the left second and third toes, bilateral rockerbottom feet, and micropenis. Chest x ray showed dextrocardia without situs inversus (fig 2). The diagnosis of isolated dextrocardia was confirmed by echocar- diography.

Chromosome studies were performed on peripheral lymphocytes by a modification of the banding technique of Seabright4 and showed the non-disjunction type of trisomy 13 in both twins.

Concordance was shown for both HLA antigens and blood groups (O+;C+;E--;c--;N+), establishing monozygosity. The condition of both twins rapidly deteriorated. Twin A died on the second day of life and twin B on the fifth day. Parental consent for necropsy was refused.

Discussion

It has been suggested that the incidence of congenital malformations in monozygotic twins is higher than in dizygotic twins and singleton infants.5-7 Although monozygotic twins share an identical hereditary genetic constitution, discordance with respect to congenital anomalies has been reported previously.5-7 The reported concordance rate of monozygotic twins for cleft lip or palate, central nervous system malformations, and congenital heart disease is only 10 to 40%, suggesting a relatively low genetic contribution to the aetiology of these anomalies.8-10 The occurrence of Down’s syndrome in twins has been extensively studied. The estimated concordance rate for trisomy 21 in monozygotic twins is 80 to 100%.6,11 Although concordant for the genotype, a different phenotype (major congenital malformation in only one twin) has been previously reported,12 emphasising the role of non-genetic factors in the development of malformations.

To the best of our knowledge, this is the first published case of trisomy 13 in twins. Although in this case concordance was proven for the genotype, the twins differed markedly in their major congenital malformations. Twin A had severe cleft lip and palate, left diaphragmatic hernia, and trigonocepha- lty, while twin B had isolated dextrocardia without recognisable cardiac defects.

Two basic explanations for congenital malforma- tions in twins exist: (1) the germ cell theory, which assumes the defect to be present in the gametes before fertilisation, and (2) the environmental theory, which assumes that the defect is produced after fertilisation has taken place.11 As this phe- nomenon in monozygotic twins cannot be explained on the basis of dissimilar genetic constitution, intrauterine environmental influences and the action of a variety of unknown exogenous agents must be considered as the possible mechanism for this condition.11,13

References

Partial trisomy 17q and a generalised bone dysplasia in a 12 week fetus

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SUMMARY A fetus, which was spontaneously aborted at 12 weeks' gestation, was found to have a generalised bone dysplasia and an unbalanced karyotype with trisomy for 17q23-1→qter due to a maternal translocation: 46,XX,t(5;17)(p15.3;q23-1)mat.

We present the clinical details of a fetus, spontaneously aborted at 12 weeks' gestation, which had an unbalanced translocation between chromosomes 5 and 17 and a generalised bone dysplasia.

Case report

The fetus was the product of the third pregnancy of a non-consanguineous Scottish couple. The pedigree is shown in fig 1. Their first pregnancy was terminated at 20 weeks' gestation when anencephaly was diagnosed in the fetus by ultrasound examination. The second pregnancy aborted spontaneously at eight weeks' gestation. No pathological or cytogenetic studies were performed on either abortus. Chromosome analysis was performed on both parents and the proband was found to be the carrier of a balanced reciprocal translocation, 46,XX,t(5;17) (p15.3;q23-1) which she had inherited from her mother (fig 1). The present fetus aborted spontaneously at 12 weeks' gestation. Macroscopic examination revealed abnormally short limbs and a cleft palate and the phenotypic sex was male (fig 2).

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