Two successive partial trisomies for opposite halves of chromosome 22 in a mother with a balanced translocation

A 30-year-old white female, G2, P1, delivered a 2000 g female infant, measuring 42.5 cm, with the following anomalies: unilateral cleft lip with cleft palate, low set ears, microcephaly, simian creases, and rocker bottom feet. The child died a cardiac death 2 days after birth. At necropsy, the heart showed a truncus arteriosus with ventricular and atrial septal defects. There was also an absent left kidney. There was neither anal atresia nor micrognathia. Chromosomal analysis showed 46,XX,+22(q13→qter) (figure a). Karyotyping of the child’s parents showed the father to be 46,XY and the mother to be a balanced translocation carrier, 46,XX,t(7;22)(p22;q13) (figure b).

A diagnostic amniocentesis was done at 17 weeks menstrual age in her next pregnancy and showed a fetal chromosomal complement of 47,XX,+del(22)(q13) (figure c). The fetus had received both maternal chromosomes 22 through non-disjunction and had a partial trisomy 22 for that region of the chromosome not represented in the first child described in this report. The parents elected to continue the pregnancy and at 36 weeks, after spontaneous labour, a female infant was delivered. This infant weighed 2130 g and had multiple congenital anomalies. These included a flattened bridge of the nose with hypertelorism and marked microphthalmia, preauricular sinus, low set ears, and rocker-bottom feet. A soft systolic murmur was heard. The palate was intact and the genitalia were normal. The anus was patent. Progressive respiratory distress ensued and the infant died after 51 hours. Additional findings at necropsy included a bicuspid aortic valve, patent foramen ovale, and pulmonary atelectasis with hyaline membrane disease. Blood and skin for chromosomal analysis confirmed the prenatal diagnosis.

The clinical features of our two patients were compared with other reported patients who had partial or complete trisomy for chromosome 22.1 The only positive clinical features shared by our two newborns were low set ears and failure to thrive. Other characteristic findings, such as cleft palate, congenital heart disease, microcephaly, microphthalmia, and preauricular sinuses,2 were subdivided between the two (see above description). This is of interest because together our two infants represent a total trisomy owing to non-overlapping areas of chromosome 22. Neither child had coloboma, anal atresia, or micrognathia, commonly associated with the cat eye syndrome, which has been attributed to partial trisomy 22.2 These comparisons are valuable, not only to see where our patients fit in the described phenotypic spectrum, but also to highlight the difficulty in making a clinical diagnosis without clear cytogenetic support.

This report includes the first clinical description of a partial trisomy 22 for the distal end of the chromosome.

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References


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