Features of di George syndrome in a child with 45,XX, -3, -22, +der(3),t(3;22)(p25;q11)

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SUMMARY A child with an unbalanced translocation resulting in monosomy for chromosomes 22 (q11→pter) and 3(p25→pter) is described. Although no immunological dysfunction could be demonstrated, the abnormalities found are similar to those seen in the di George syndrome which has been associated with monosomy for the same region of chromosome 22.

The clinical features of di George syndrome (DGS) include neonatal hypocalcaemia, defective cellular immunity, absent parathyroid and thymus glands, atypical facial features, and cardiovascular abnormalities. More recent reports suggest a greater variability of the syndrome than originally described and Lischner advocated the name partial DGS to include these patients, some of whom have small amounts of thymic tissue and little, if any, immunological deficiency. There is evidence for an association of a partial deletion of chromosome 22 and DGS. We report here a child with an unbalanced chromosome translocation (3;22) with some features of DGS.

Case report

The child was the second daughter of non-consanguineous Asian parents. There is no family history of illness. The pregnancy was complicated by a urinary tract infection at five weeks and bleeding due to placenta praevia at 36 weeks. Fetal growth slowed considerably during the last part of pregnancy to such an extent that a Caesarian section was carried out at 37 weeks. Marked meconium staining was noted and respirations were established at five minutes. Her birth weight was 2120 g (<10th centile), occipitofrontal circumference 29.5 cm (<10th centile), and length 47.2 cm (25th centile). A number of dysmorphic features were noted soon after birth and included broad thumbs, upward slanting eyes, micrognathia, fish-like shape of mouth, short philtrum, and low set ears. A soft grade 4/6 pansystolic murmur was also heard. Tachypnoea caused feeding difficulties and prolonged her stay in hospital. Feeding caused significant difficulty throughout infancy requiring gavage feeding. Her weight gain was initially slow but by six months was on the 10th centile, although her head circumference continued to increase at a constant rate below the 10th centile.

INVESTIGATIONS

Chest x ray during the neonatal period, and tomography later, failed to show a thymic shadow. The values for serum calcium (2.56 mmol/l) and phosphate (1.84 mmol/l) were normal. Urea was 2.5 mmol/l and creatinine 37 mmol/l. IgG (4.7 g/l) and IgA (0.31 g/l, assessed at 13 weeks, were within the normal range for age although IgM (1.27 g/l) was raised. Lymphocytes showed a normal response to in vitro stimulation with phytohaemagglutinin, concanavalin A, or pokeweeds mitogen and no abnormality was detected in the proportion of lymphocytes identified by T3, T4, T8, B, or M markers. Phagocytosis of 131I labelled zymogen particles by polymorphonuclear cells was normal. Rubella HAI titre was less than 8 and CMV IgG antibody titre (FA) was less than 64. Cardiac catheterisation showed a small VSD and subsequent angiography demonstrated a complete absence of the left pulmonary artery.

CYTOGENETIC STUDIES

Chromosome preparations of lymphocytes and fibroblasts revealed an unbalanced translocation between...
chromosome 22

The normal chromosome 3, the derivative 3, and chromosome 22 stained by trypsin-Giemsa (top), orcein (middle), and C banding (bottom).

Discussion

This child, investigated because of dysmorphic features noted in the neonatal period, was found to have an unbalanced chromosome translocation involving loss of material from both chromosomes 3 and 22.

Children with congenital abnormality and unbalanced inversion products resulting from familial pericentric inversions of chromosome 3 with a breakpoint at p25 have been reported, but recognition of features specifically due to p25→pter deletion is difficult in these cases. A child with an unbalanced translocation resulting in partial trisomy 17 and deletion of 3p25→pter showed phenotypic abnormalities characteristic of trisomy 17. Pure monosomy for 3p25 has been reported in four cases (reviewed in Higginbottom et al) and these show similar congenital abnormalities, but of an unspecific nature.

De la Chapelle et al reported a child with the di George syndrome in association with an unbalanced translocation between chromosomes 20 and 22. The balanced form of the translocation was found to be familial and three earlier children had died with similar symptoms. The suggestion that the di George syndrome is associated with deletion of 22q11→pter was supported by reports from Kelley et al and Greenberg et al of four unrelated cases of patients with a di George-like phenotype, a variable degree of clinical expression, and an unbalanced translocation involving chromosome 22 with chromosomes 3, 4, 10, or 20. The mother of the child with the 4;22 translocation also carried the unbalanced translocation, but was asymptomatic although investigation showed her to have a partial T cell deficiency. Thus, even within one family, there is a marked variation in clinical expression.

There is some difficulty in establishing the exact breakpoint of these translocations involving the pericentric region of chromosome 22. De la Chapelle et al have argued that the absence of any evidence of di George type symptoms in patients with 21;22 translocations or 22p deletion would suggest that the deletion of significance is at q11 and not on the short arm. In our patient, there is no evidence of a second centromere on the derivative chromosome but it would appear that most, if not all, of 22q is present, implying a breakpoint at q11.

The clinical features of this child, which are shared with those of patients with DGS, include upward slanting eyes, short philtrum, fish-like mouth, micrognathia, low set ears, an absent left pulmonary artery, and ventricular septal defect. Our patient lacks the classical features of hypoparathyroidism and defective cellular immunity described by di George although no thymus was visible on x ray and she has no facial features and cardiac anomalies consistent with the partial DGS.

Although the specific breakpoint may be important in some cases the variation in the clinical features in cases with similar deletions may say more about the activity of the alleles for which the patient is effectively hemizygous than about those that are missing.

References

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A child with partial monosomy 6q secondary to a maternal direct insertional event

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SUMMARY We report a child, now aged two years, who is monosomic for the region 6q23.1–q24.2. Her mother, older sister, and twin sister have a balanced chromosome complement with this region of 6q being inserted into 11q.

Case report

The proband was one of twins born to healthy non-consanguineous parents; the mother was aged 35 and father 38. The twins were born at term with the proband weighing 2.6 kg and her sister 3.1 kg.

The mother had had two previous pregnancies. The first resulted in a healthy daughter and the second in a son who died aged six weeks from cardiac failure. He was reported as being dysmorphic with micrognathia and a large neck and unfortunately was not karyotyped. Necropsy (Dr R I K Elliott) showed cardiomegaly with a patent ductus arteriosus and a small patent foramen ovale but no explanation for the cardiomegaly. The skin was unusually thick and indurated.

There were no neonatal problems but the proband was slower to gain weight than her healthy sister. At the age of five months she had the first of several hospital admissions for recurrent chest infections, persistent collapse of the right upper lobe, and failure to thrive. At the age of two years it was noted that length, weight, and head circumference growth rates had all fallen and the liability to infections persisted. She was globally delayed and barely able to sit unsupported. She transferred objects from hand to hand but said no meaningful words.

Her face was slightly dysmorphic (fig 1) with hypertelorism, prominent bridge to the nose, tented nares, and a small mouth with narrow upper lip. Overall her facies was reminiscent of that seen in the Wolf-Hirschhorn syndrome. Her palate was normal, her ears were low set, and her neck short, but no other abnormalities were noted.

Investigations have shown recurrent iron deficiency anaemia but normal immune function and sweat tests.

CYTOGENETIC FINDINGS

Peripheral blood lymphocytes from the proband, her twin sister, and both parents were cultured for chromosome studies with trypsin G banding. The proband showed a small interstitial deletion of the long arm of chromosome 6. Examination of the maternal chromosomes showed a direct insertional event between chromosome 6 and the long arm of chromosome 22.

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