Distal 10q trisomy syndrome with unusual cardiac and pulmonary abnormalities

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Abstract
Since its description in 1965, distal 10q trisomy has become recognised as a well defined, although rare syndrome, almost always the result of an unbalanced translocation. Typical features consist of psychomotor delay, a distinctive dysmorphic appearance, growth retardation, and, in some cases, cardiac, renal, and ocular abnormalities. (J Med Genet 1998;35:72–74)

Keywords: distal 10q trisomy; cardiac abnormalities; pulmonary abnormalities

We report a child with partial trisomy 10q, displaying many of the usual phenotypic abnormalities, who also had an unusual combination of cardiac and pulmonary abnormalities. These consisted of mild hypertrophic cardiomyopathy, ventricular septal defect, pulmonary stenosis (valvar and infundibular) with a moderate sized bronchial collateral supplying the right lung, absent right pulmonary veins, and a narrow, hypoplastic left main bronchus. This combination of anomalies presents an interesting management challenge and, to our knowledge, has not been described in children with this chromosomal abnormality.

Case report
The child was born to a primigravid, healthy mother. The paternal grandparents had seven miscarriages but there was no history of phenotypically abnormal children. The pregnancy had been uneventful, with no teratogenic exposure. Intrauterine growth retardation was noted and delivery was by caesarean section for fetal distress at 42 weeks' gestation. Birth weight was 2300 g (<3rd centile) and dysmorphic features included low set ears, epicanthic folds, hypertelorism, bilateral ptosis, a high forehead, broad nasal bridge, and a bow shaped mouth (fig 1). He had clinodactyly and a dermoid cyst in the middle of his forehead.

Chromosomal analysis showed an unbalanced translocation resulting in trisomy of the distal one third of the long arm of chromosome 10 (q24–26) (fig 2). A balanced translocation of that segment was found in his father between chromosomes 10 and 18 (fig 3), with breakpoints at 10q24 and 18p11.31 or 32.

During his first two years, recurrent episodes of cardiac failure and respiratory tract infections with right upper lobe consolidation and collapse continued to be a problem. Normal investigations included immunoglobulin and IgG subclass levels, sweat electrolytes, neutrophil function studies, alpha-1-antitrypsin levels, viral and atypical serology, ciliary beat frequency, and cine barium swallow. Cincinnati chest X ray (penetrated for visualisation of airways) showed an extremely narrowed left main bronchus and some narrowing of the upper airways (fig 4).

Aged 5 years, he was continuing to show delay in his growth and development, but was walking well and had good comprehension. A CT chest scan showed early right sided bronchiectasis when investigated for haemoptysis associated with respiratory tract infections. There was absent perfusion and diffusely reduced ventilation on the right side on ventilation and perfusion scan. The left side appeared normal. Cardiac catheterisation showed absence of the right sided pulmonary veins. The left sided veins were normal, as were both pulmonary arteries. The right lung was also supplied by a moderate sized bronchial collateral. Valvar and infundibular stenosis of the right ventricular outflow tract was shown with suprasystemic right ventricular pressures (fig 5). At fibreoptic bronchoscopy, severe narrowing of the left main bronchus with complete cartilage rings was noted. In summary, virtually all the blood flow was to his left lung, whereas the main left bronchus was much narrower than the right.

Figure 1 Typical facial features of distal 10q trisomy syndrome with forehead dermoid cyst removed.
Following balloon dilatation of the right ventricular outflow tract, the child’s respiratory symptoms have improved. Currently he still has recurrent chest infections, often with small haemoptyses, but without hypoxia or cardiac failure and not requiring admission to hospital.

Discussion
Partial trisomy of the long arm of chromosome 10 was first described in 1965 and since then over 30 published reports have appeared of children with this disorder. The well recognised and defined type of trisomy of the long arm of chromosome 10 has been labelled the “distal 10q trisomy syndrome” to distinguish it from other forms of partial 10q trisomy. A recognisable phenotype has been suggested but critical attempts to correlate karyotype with phenotype are lacking. In this syndrome, of which our case is an example, the only common trisomic segment in these patients is the distal two chromosome bands 25 and 26. Patients with the trisomic segment situated either proximally or in the middle of 10q have been less frequently described and do not represent phenotypically distinct syndromes.

Table 1 details the common features of the distal 10q trisomy syndrome. Patients with trisomy 10q24—qter tend to have more severe clinical manifestations such as heart or renal abnormalities. Ocular abnormalities are particularly well described and consist of hypertelorism, blepharophimosis, and microphthalmia as well as occasional fundus changes. Developmental delay to varying degrees is invariably present, and most cases also display somatic growth delay.
Cardiac anomalies have been frequently reported. A publication in 1979 summarising all the previously described cases of this chromosomal abnormality found that seven of 13 children were thought to have some form of congenital heart disease.

There are no published cases, to our knowledge, describing similar pulmonary and vascular anomalies to those described here, although this may be because of lack of appropriate investigative tools at the time of reporting. Interestingly, however, several other papers report children with this chromosomal disorder who have died of pneumonia. The application of more modern investigative techniques may have provided more diagnostic information had they been available at the time.

This case is complicated by a partial monosomy of chromosome 18p which may account for some of the features seen. The amount of 18p material deleted in this case is extremely small compared with those who have the recognisable 18p- syndrome and is highly unlikely to contribute to the cardiac and respiratory abnormalities. Forabosco et al reported a case with very similar breakpoints to those described here with no cardiac or pulmonary involvement. Interestingly, in their case, the child had no impairment of growth or psychomotor development.

There are several long term therapeutic options in this case. Further balloon dilatation of his right ventricular outflow tract will probably be required. Surgery on his narrowed left main bronchus is too hazardous as this could potentially compromise his one functioning lung. Embolisation of the bronchial collateral supplying the right lung, the presumed site of haemoptysis, is an option should haemoptysis become a more significant problem. Currently he is well and managed conservatively.

We would like to thank Dr J W Taylor, Cytogenetics Department, St George’s Hospital, London, for his help in preparing this manuscript.

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*J Med Genet* 1998 35: 72-74
doi: 10.1136/jmg.35.1.72

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