Mosaic trisomy 16 in a live newborn infant

N J Gilbertson, J W Taylor, I Z Kovar

Abstract
Trisomy 16 is thought to be incompatible with fetal survival. A boy with mosaic trisomy 16 who lived for 11 weeks is reported. Chromosome analysis was carried out on skin fibroblasts grown during life and confirmed on samples taken at necropsy. We believe that this is the first report of mosaic trisomy 16 that has been confirmed by cytogenetic banding.

Trisomy 16 is the most common autosomal anomaly seen in early aborted fetuses, but has been thought to be incompatible with full embryonic development; empty sacs and disorganised embryos are usually seen.1 There are reports of putative full and mosaic trisomy 16 but none confirmed by cytogenetic banding studies. We report an abnormal baby boy born alive with mosaic trisomy 16 who survived for 11 weeks.

Case report
A baby boy was born at 39 weeks' gestation to healthy Chinese parents. During the pregnancy symmetrical intrauterine growth retardation and polyhydramnios were noted; delivery was by caesarean section because of fetal distress and the growth retardation.

The baby was initially cyanosed with regular shallow respiratory efforts and a heart rate of 60 beats/minute; he responded well to oxygen given by face mask. Craniofacial and musculoskeletal dysmorphic features were immediately apparent as was clinical growth retardation; he weighed 1590 g at birth, was 40 cm long, and had a head circumference of 31·5 cm (all below the third centile). The abnormalities noted are listed in the table; the infant is shown in the figure.

The baby required ventilatory support for the first three weeks because of respiratory distress, and was then maintained with oxygen given by headbox for a further three weeks. The respiratory insufficiency was thought to be caused by restriction from kyphoscoliosis. A ventilation-perfusion lung scan carried out at 8 weeks showed equal ventilation bilaterally, but with reduced ventilation in the upper zones. There was a discrepancy between ventilation and perfusion in the right upper zone, which was underperfused.

A loud pulmonary second sound and forceful left parasternal impulse were noted at birth; a pansystolic murmur was audible at the lower left sternal edge from 4 weeks. An echocardiogram showed a small perimembranous ventricular septal defect, which seemed to be closing by apposition of the tricuspid valve tissue.

Phenotypic abnormalities

Craniofacial:
- Cranial asymmetry
- Prominenence of left side of forehead
- Abnormal scalp hair patterning with high frontal hairline on left
- Hypertelorism
- Swelling of right upper eyelid giving appearance of ptosis and micro-ophtalmia
- Partial coloboma left iris
- Short nose, flat nasal bridge
- Thickened upper lip with groove in upper gingival margin, left of midline
- Preauricular pit
- Low set malformed auricles

Musculoskeletal:
- Thoracoabdominal scoliosis convex to left
- Prominent left hemithorax
- Hypoplastic left nipple
- Single palmar crease, left hand
- Middle finger overlapping index
- Wide space between first and second toes
- Dorsiflexed toes
- Bilateral talipes calcaneovalgus

Genital:
- Undescended testis
- Left inguinal hernia
- Non-communicating hydrocele

Cardiac:
- Ventricular septal defect

Appearance of infant with mosaic trisomy 16

The infant was fed initially by nasogastric tube, and subsequently by bottle from the age of 8 weeks. Despite a maximal milk intake his weight gain was slow and erratic. He gained just
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over a kilogram during the 11 weeks of his life, and weighed 2610 g at the time of his death. He was always socially unresponsive and was never seen to smile. He was hypotonic with no head control (‘rag doll-like’). He was visually attentive and had normal non-conveal electroretinograms, with bilateral cortical visual evoked potentials to flash stimuli. He did not respond to sound; auditory evoked potentials at eight weeks were inconsistent. An electroencephalogram showed no abnormality.

At 11 weeks his inguinal hernia became difficult to reduce and surgical repair was indicated. He was by this time clinically stable, and had been breathing air spontaneously for five weeks. The endotracheal intubation for general anaesthesia was extremely difficult, and the patient became hypoxic and had a cardiorespiratory arrest. Resuscitation was unsuccessful.

Necropsy was performed at the request of the coroner and left sided hypertrophy of the cornua of the hyoid bone and thyroid cartilage was found, together with pronounced hypertrophy of the left arytenoid cartilage which seemed to have caused partial obstruction of the glottis. There was a deep cleft between the hypertrophic left arytenoid area and the epiglottis. The vocal cords were asymmetrical, particularly on the left, and there was stenosis in the midsection of the trachea. A small aberrant left bronchus that was barely patent was present above the left main bronchus, and the right bronchus was elongated. There were no other findings of note except for the small perimembranous ventriculoscutal defect previously noted, and the dysmorphic features.

Discussion

We believe that this is the first report of a mosaic trisomy 16 confirmed by cytogenetic banding, and we have described phenotype abnormalities in addition to those noted previously in potentially similar cases.

Trisomy 16 has been reported as the most common autosomal trisomy in aborted fetuses. There have been isolated reports of 16– mosaicism in adults; a physically and mentally normal Japanese father of an aborted fetus that was trisomic for chromosome 16 had 3% of his white cells and 15% of his skin fibroblasts trisomic for chromosome 16. Some of the abnormalities that were seen in our proband—including kyphoscoliosis, breast hypoplasia, and asymmetric musculoskeletal development—have been described previously, but in these reports of mosaic trisomy 16 and in two of full trisomy 16 chromosome analysis was carried out without modern banding techniques. 4 5

There have been reports of infants surviving with trisomy of either 16p or 16q material confirmed by GTG banding6; partial trisomy 16 may also occur with familial 16:21 translocation. Patients with partial trisomy 16 all have multiple congenital anomalies and with one exception have died within the first year of life. As both trisomy 16p and 16q are compatible with intrauterine life and a limited degree of postnatal survival Roberts and Duckett concluded that it must be the combined effects of additional long and short arms which render the full trisomy invariably lethal. 6 It is noteworthy that trisomy 16 has been detected in chorion villus biopsy specimens from a number of cases in which the fetus itself was unaffected, indicating that the mosaic is formed in the early embryo and that the trophoblast can function normally in the presence of this trisomy. Our proband has some features in common with those seen in partial trisomies, namely skull asymmetry, auricular abnormalities, single palmar crease, cryptorchidism, inguinal hernia, ventriculoseptal defect, failure to thrive, and developmental retardation. Other features of this infant that have not been noted previously include coloboma, right inguinal hernia, scalp hair, and asymmetry of laryngeal structures.

This infant had a higher proportion of abnormal cells than suggested by the unconfirmed reports of mosaic trisomy 16. It remains to be determined what proportion of abnormal cells can exist before a mosaic trisomy becomes lethal, or if a full trisomy 16 is compatible with survival. Consideration of this baby’s anomalies may assist in delineation of a phenotype for the trisomy of chromosome 16 material.

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