Partial deletion 21: case report with biochemical studies and review

NANCY J CARPENTER, JARY S MAYES, BURHAN SAY, AND DON P WILSON
H Allen Chapman Research Institute of Medical Genetics, Children’s Medical Center, Tulsa, Oklahoma, USA

SUMMARY An unbalanced translocation of a portion of the long arm of chromosome 21 to the short arm of chromosome 4 resulted in a partial deletion of chromosome 21 (pter→q21-05) and in the loss of the telomere of 4p. The phenotype of the child included asymmetrical facies, microcephaly, short stature, hypotonia, and psychomotor retardation associated with frequent infections. Normal SOD-1 activity in red blood cells and fibroblasts and normal cystathionine β synthase activity in fibroblasts suggest that these gene loci are distal to 21q21-05.

Most structural abnormalities of chromosome 21 are rings resulting from terminal deletions, although unbalanced translocations and interstitial deletions of portions of the long arm have occasionally been reported.1 We describe a child with an unbalanced translocation involving chromosomes 4 and 21 resulting in deletion 21pter→q21-05, who exhibited some of the features reported in other cases of proximal monosomy 21q.

Case report

The proband, a female, was the third child born to a mother and father who were 33 and 32 years old, respectively. The infant was born at term by breech delivery after a rapid labour.

The mother's previous two pregnancies resulted in two healthy daughters, now 14 and six years old. There is no family history of congenital malformations, mental retardation, or multiple spontaneous abortions.

At birth, the patient's weight was 2155 g, length 48 cm, and head circumference 32.5 cm. Apgar scores were 7, 7, and 6 at one, five, and 10 minutes. She had mild respiratory distress and was hypotonic. An intermittent grade I/IV short systolic murmur was present, thought to be due to a persistent duc tus arteriosus. The ECG was normal except for sinus bradycardia. A right subependymal cerebral haemorrhage resolved by the fifth day of life. A brain scan showed no dilatation of the ventricles, but an EEG showed episodic focal dysrhythmia of the left temporal area. Visual evoked potential studies showed P100 values typical of an infant of 36 to 37 weeks' gestational age. She responded to auditory stimuli. Additional clinical findings included a hoarse cry, small facies with micrognathia, low set ears, and long toes with a long midline crease in each foot. She was discharged from hospital at 11 days of age.

An EEG at 40 days of age showed suppression of activity in the right posterior hemisphere and rhythmic slow activity in the right frontal area. The child had recurrent, mostly viral, infections. At nine months tubes were inserted in both tympanic membranes. Serum immunoglobulin levels at 23 months were normal.

At 34 months of age the child’s weight was 7.6 kg, height 81.2 cm, and head circumference 42 cm, all below the 3rd centile for age. She was unable to crawl or walk and she had no speech but was able to use sign language to a limited extent. She had facial asymmetry with the left side being smaller than the right and had a tendency to tilt her head to the left. Other facial features included a prominent forehead, head, horizontal palpebral fissures, high nasolabial bridge, micrognathia, and low set, posteriorly rotated ears with prominent antihelices (fig 1). No heart murmurs could be detected. Her hands were slender with ulnar deviation of the second and third fingers bilaterally. Her feet were narrow with prominent heels. The results of the Developmental Sequencing Performance Inventory administered at 49 months of age were at the 23 month level for gross motor, fine motor, and social/self-help skills and at the 27 month level for cognitive and language skills.

CYTOGENETIC STUDIES

Giemsa-trypsin banding of prophase and prometa phase chromosomes from the child’s cultured blood lymphocytes and skin fibroblasts showed 45 chromosomes with a single normal chromosome 4 and a single normal chromosome 21. The distal segment of the long arm of the other chromosome 21 was translocated to the telomeric region of the short arm of the second chromosome 4 (fig 2). The child was therefore monosomic for the short arm, centromere and proximal part of the long arm of chromosome 21.

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and for the telomere of chromosome 4p. Her karyotype was 45,XX,-4,-21,+der(4)t(4;21) (4qter→4p16-3::21q21-05→21qter). Fibroblasts from the child have been contributed to the NIGMS Human Genetic Mutant Cell Repository (GM 8210). The karyotypes of the parents were normal.

**Dermatoglyphics**
Dermatoglyphic analysis of the hand showed nine ulnar loops and one whorl. The total finger ridge count (87), add angle (86°), and the palmar creases were normal. Loops were observed in the fourth interdigital spaces bilaterally.

**Biochemical Studies**
Superoxide dismutase-1 (SOD-1) was assayed by inhibition of the epinephrine autoxidation. The activity of SOD-1 in red blood cells was 3550 U/ml of RBC (controls 3080 to 4300) and in fibroblasts was 9-0 U/mg of cell protein (controls 8-3 to 9-9).

Cystathionine β synthase (CBS) was assayed by Dr Jan Kraus, Yale University. Activity was 25 mU/mg of cell protein which is in the normal range for fibroblasts (10 to 30 mU/ml).

**Endocrine Studies**
Standard provocative stimulation tests for growth hormone were conducted using arginine hydrochloride and glucagon. Quantitative levels of growth hormone were normal with a peak response of 14 ng/dl. Somatomedin C was 0-75 U/ml (controls 0-14 to 1-44 U/ml). Cortisol and thyroid function were normal.

**Discussion**
Lejeune et al were the first to report a partial monosomy of a G group chromosome in an infant mosaic for a ring (G). These investigators considered this case the countertype to Down’s syndrome and,
as other cases were reported with similar clinical findings, the term ‘antimongolism’ was used. Warren and Rimoy divided the G deletion syndromes into two categories, G deletion syndrome I later becoming identified as partial deletion of chromosome 21. Downward slanting palpebral fissures, a broad nasal bridge, microcephaly, significant growth retardation, hypertonia, and psychomotor and mental retardation are features of this syndrome. Complete monosomy 21 is rare but has been reported to include cleft lip or palate or both and multiple skeletal deformities with congenital contractures, in addition to the previous findings. The first observation of partial monosomy 21 involving the proximal q arm was reported by Rethoré et al. Other cases have since been described resulting from familial or de novo unbalanced translocations of chromosome 21 distal to band q21 onto other autosomes. Reported clinical findings included horizontal palpebral fissures, high forehead, aplastic nasal bridge, skeletal malformations, bone fragility, and mental retardation (IQ below 50). Some differences among the patients are expected, since the breakpoints in 21q probably differ in each case and the partial deletion of the other autosomes may influence the phenotype.

Our patient does not display the downward slanting palpebral fissures or the joint contractures found in the 21 deletion or monosomy 21 syndromes. She most closely resembles the patient reported by Schmidt et al. and later by Wahrman et al. who also had the breakpoint approximately in the centre of band 21q21, although this patient had normal growth. In the other cases of proximal monosomy 21q, either the breakpoint is difficult to ascertain from the photographs available or it is located more distally in band 21q22.

The locations of the breakpoints in this translocation make the present case particularly useful for regional localisation of DNA probes on chromosome 21 and possibly for the identification of DNA markers near the gene for Huntington’s disease and the probe G8, which has been mapped to 4p16.3 by in situ hybridisation.

The gene for SOD-1 has been assigned to chromosome 21 by linkage analysis in somatic cell hybrids. The activity of SOD-1 is about 1.5 times greater in cells from patients with trisomy 21. Studies initially indicated that the gene was localised in sub-band 21q22.1. In the present case, normal SOD-1 activity was measured in red blood cells and fibroblasts, suggesting that the gene is located distal to 21q21-05.

The gene for CBS has been assigned to chromosome 21 by somatic cell hybridisation and the activity of this enzyme is increased by about 50% in fibroblasts from patients with trisomy 21. Localisation of the CBS gene has been suggested to be between 21q21 and 21q22-1 by dosage effects of partial duplications of chromosome 21 in somatic cell hybrids and to be in 21q22 by in situ hybridisation. Normal activity of CBS in fibroblasts from the proband suggests that the gene for this enzyme is located distal to 21q21-05.

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References


Deletion 15q21.1→q22.1 resulting from a paternal insertion into chromosome 5

MOH-YING YIP*, MARK SELIKOWITZ†, NEVILLE DON†, ALEX KOVACIC*, S PURVIS-SMITH*, AND P R L LAM-PO-TANG*

*Cytogentic Unit, Prince of Wales Hospital, and †Department of Paediatric Medicine, Prince of Wales Children’s Hospital, Sydney, Australia.

SUMMARY A 15 month old boy with an interstitial deletion 15q derived from a paternal insertion (5;15)(q31;3;q21.1q22.1) is described and compared with one other reported case. A beak like nose with hypoplastic nasal alae, a thin upper lip, failure to thrive in infancy with later onset of obesity, and severe mental retardation are features common to both.

Insertions involving three breakpoints are the least common chromosome rearrangements in man. With banding studies, insertions have been detected between and within chromosomes. We are aware of 21 published reports of interchromosomal insertions. We describe here a case of an interstitial deletion of 15q21.1→q22.1 derived from a paternal insertion in a 15 month old boy.

Case report

The proband was the first live born child of healthy, unrelated parents. At his birth, the mother was 25 and the father 30. There were three previous miscarriages at six, eight, and 12 weeks.

The pregnancy with the proband was complicated by polyhydramnios and decreased fetal movement which developed at 30 weeks. Caesarean section was performed at 35 weeks’ gestation for intrauterine growth retardation. The Apgar scores were 8 and 9 at one and five minutes, respectively. The birth weight was 1680 g (less than the 3rd centile), head circumference 30.7 cm (10th centile), and birth length 40 cm (3rd centile). Dysmorphic features noted at birth were: narrow bifrontal diameter, a patent posterior fontanelle, low temporal hair implantation, and low set ears. The mouth had a thin upper lip and was usually kept open. The nose was beak shaped with poor development of the nasal alae (fig 1a). The hands were held in ulnar deviation and there was bilateral clinodactyly of the fifth fingers and simian creases. Dermal ridge patterns were present on all 10 fingers. Creases were decreased around the elbows and knees which, however, had a full range of movement (fig 1b). The testes were bilaterally undescended with an under-developed scrotum. Bilateral inguinal herniae were present and the penis was of normal size but hooded with coronal hypospadias and chordee.

There was hypotonia and poor feeding. He had occasional apnoeic episodes during the neonatal period. A chest x ray showed a raised anteromedial portion of the right hemidiaphragm which was inserted high on the posterior surface of the sternum. X ray of the spine showed that the vertebral bodies had a square outline and reduction of interpedicular distances in the lumbar region.

From the age of six months the patient has gained weight rapidly despite dietary joule restriction. At present, at the age of 15 months, his weight is on the 90th centile and his length on the 10th centile. His psychomotor development corresponds to a six month level. He has developed a mild degree of scoliosis and strabismus associated with myopia.

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