An interstitial deletion of chromosome 7(q35)

Kerry Fagan, Christine Kennedy, Laurence Roddick, Alison Colley

Abstract
We describe a patient with developmental delay, mild dysmorphic features, and monosomy of 7q35. Only one other patient with an interstitial deletion of this band has been previously reported. A review of clinical features of these two children did not show similarities in dysmorphic features. Reports of patients with other 7q interstitial deletions are listed.

Interstitial deletions resulting in partial monosomy for different segments of the long arm of chromosome 7 have been described in cytogenetic publications. Interstitial deletions have been broadly classified according to the deleted segment into three main groups: deletions of 7(q11), del(7)(q21), and del(7)(q31). Fewer cases have been reported with deletions of 7(q33) and 7(q34). The phenotypic features of our patient with deletion of band q35 are presented and other patients with deletions involving the 7(q3) region are reviewed.

Case report
The proband was the second child of non-consanguineous parents. Her mother was 28 years and her father 33 years at the time of her birth. Her mother had two first trimester spontaneous abortions and has two normal sons. Labor was induced at 42 weeks of gestation after an uncomplicated pregnancy; birth weight was 2970 g and length was 49.5 cm. There was neonatal jaundice owing to Rh incompatibility with no major complications. Physical milestones were within normal limits; she sat by 7 months, crawled by 8 months, and walked by 14 months, but used only a single word until aged 2½ years. At 3½ years she still had only a few single words and speech therapy, which included Makaton signing, was started. Apart from bilateral hypermetropia and a convergent squint, her general health was good.

From the age of 2 years the proband attended special education and language classes. At 4½ years the Reynell verbal comprehension scale put her expressive language at only 2½ years. At 7 years a WPPSI assessment gave a full scale of 85 and at the age of 10 years a Stanford Binet assessment showed her to be functioning in the mildly intellectually handicapped range. Biochemical studies including thyroid function tests, mucopolysaccharide screen, and urinary amino and organic acids were all normal, as was her cerebral CT scan. Bone age was appropriate to chronological age.

When reviewed at the age of 11 years her head circumference was on the 50th centile with height and weight between the 90th and 95th centiles. Dysmorphic features included epicantic folds and almond shaped eyes (fig 1). Her inner and outer canthal distances and interpupillary distance were all on the 50th centile. There was a marked left convergent squint, a high palate, and a bulbous nasal tip. She had low set, posteriorly rotated ears and apparent micrognathia. Her speech was markedly hypernasal; her palate and uvula appeared normal; however, velopharyngeal incompetence was shown on video fluoroscopy. She attends special classes in a normal school.

Cytogenetic studies
Chromosome studies performed on the patient at birth and at the age of 2 years showed an apparently normal female karyotype. On re-examination at 11 years using high resolution cytogenetic techniques, a small interstitial deletion of band q35 was detectable on one chromosome 7 (fig 2). This constitutes a loss of only 0.18% of HAL (haploid autosome length). The karyotype can be described as: 46,XX, del(7)(pter→q34:q36→qter). The chromosomes, painted with a chromosome 7 specific probe (Cambio) by fluorescent in situ hybridisation, did not show evidence of a cryptic translocation or insertion. The parental chromosomes were normal.

Discussion
Deletions of interstitial segments of the long arm of chromosome 7 are still relatively rare. A review of published reports indicates several broad groups of 7q deletions which have breakpoints and phenotypic features in common (fig 3).

A deletion which includes the G band negative region of q11 results in severe growth and developmental retardation and a poor prognosis. Spasms and seizures are common and most patients remain non-ambulatory.

Four patients with interstitial deletions of chromosome 7 which included loss of band q21 have been reported. These patients all had split hand/split foot malformations among other dysmorphic features.

The most common interstitial deletion reported for chromosome 7, del(7)(q22→q31), has been described in 12 patients so far and a characteristic clinical picture is now emerging, including severe developmental retardation, an unusual cry in infancy, a large mouth, seizures, and feeding or gastro-oesophageal difficulties.

Patients with interstitial deletions in the q3 region of chromosome 7 include three members of one family with monosomy for bands

Cytogenetics Unit, Hunter Area Pathology Service, The John Hunter Hospital, Locked Bag No 1, Newcastle Mail Exchange, NSW 2305, Australia
K Fagan
C Kennedy
Department of Paediatrics, The John Hunter Hospital, Newcastle, NSW, Australia
L Roddick
Regional Medical Genetics Unit, PO Box 84, Waratah, NSW, Australia
A Colley

Correspondence to Dr Fagan.
Received 11 February 1994 Revised version accepted for publication 8 April 1994

J Med Genet 1994;31:738-739

J Med Genet first published as 10:1136/jmg.31.9.738 on 1 September 1994. Downloaded from http://jmg.bmj.com/ on July 15, 2023 by guest. Protected by copyright.
An interstitial deletion of chromosome 7(q35)

1. Their deletion resulted from malsegregation of an inversion carrier and the patients all had similar phenotypic features, including mental and developmental retardation, external ear malformation, hypertelorism, flat nasal bridge, a bulbous nose tip, and a wide mouth. A single case report of a deletion of bands q33 and q34 (patient 3) had microcephaly, prominent forehead, mongoloid slant of the eyes, bilateral simian creases, and cleft oral-malga, as well as severe growth and developmental retardation. One patient described had a deletion of bands 7q33→q35. Her features included microcephaly, neurodevelopmental delay, a cleft lip and palate, conductive deafness, and protruding tongue.

One patient has monosomy for band 7q35, similar to our patient, but it resulted from a de novo translocation t(3;7)(q27;q35). This patient was severely hypotonic at birth with absent Moro reflex; she had microbrachycephaly, ptosis of the left eyelid, alternate squint, small nose with a flat nasal bridge, and microstomia. The growth rate had been below the 3rd centile. Seizures occurred shortly after birth and a profound neural hearing deficit was documented. By comparison, our patient shows only mild dysmorphism but has a squint in common with their patient.

Conclusions
The clinical features of patients described with interstitial deletions of different segments of the long arm of chromosome 7 vary. More case reports are needed with deletion breakpoints accurately defined using high resolution banding to confirm whether a pattern of phenotypic characteristics can be described. Our case reiterates the need to repeat testing when a chromosome anomaly is suspected, despite an initially normal karyotype.

EBV transformation of the patient’s lymphocytes is in process and will be available on request.