Partial monosomy for chromosome 22 in a patient with del(22)(pter→q13.1::q13.33→qter)

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Abstract
An 18 month old girl with partial monosomy for the long arm of chromosome 22 is described. The karyotype was 46,XX,del(22)(pter→q13.1::q13.33→qter). To our knowledge this is the first report of monosomy for this specific segment of chromosome 22. Clinical features include developmental delay in all areas, hypotonia, macrosomia, full cheeks, eyebrows, and eyelids, mild epicanthus, wide nasal bridge, long philtrum, and thick lower lip. Parental chromosome studies were normal.

Case report
The proband was an 18 month old girl born to a 34 year old mother and a 28 year old father. Pregnancy was uneventful with birth weight 3600 g at 42 weeks' gestation. At birth she had a positional talipes, but no other anomalies were noted. At the age of 11 months her head circumference was 44.6 cm (50th centile), her weight was 9 kg (50th centile), and her length was 79 cm (97th centile). She had a rather unusual, low pitched, growling cry. There was a small umbilical hernia and coccygeal dimple. Other than a constant running nose and dribbling, her general physical appearance was unremarkable. At 18 months a Denver developmental assessment showed delays in all areas, particularly in fine motor and gross motor development. She is hypotonic, is not yet walking, and is unable to feed herself. Her hearing is thought to be normal, although she has only three or four recognisable words. Her length is now 87 cm (just above the 97th centile), weight 11.4 kg, and head circumference 47.5 cm (both just above the 50th centile). She is a floppy child with a relatively expressionless face, full cheeks and eyebrows, fullness of the upper eyelid, ptosis, and mild epicanthus.

There is no obvious strabismus. She has a wide nasal bridge with short columella and slightly long philtrum, a V shaped upper lip, and a thick, pronounced lower lip (fig 1). She has prominent auricles with some transverse creases on the lobes. The external genitalia are normal although the labia majora are rather prominent. Full blood count, thyroid function, serum amino acids, and routine biochemistry screen were normal. Cytogenetic studies showed a complement of 46,XX,del(22)(pter→q13.1::q13.33→qter) (fig 2). Parental chromosome studies were normal.

Discussion
Partial monosomy for chromosome 22 is associated with ring chromosome 22 and frequently with the DiGeorge malformation.1 Deletions of other segments of 22q include 22q12→qter reported by Watt et al.,2 22(pter→q12::) described by Kirshenbaum et al.,3 and 22q13.31 reported by Herman et al4 in a patient with the Goldenhar complex.

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Figure 1 Frontal facial appearance of index case.
Comparison of our patient with the one of Kirshenbaum et al is more difficult as no high resolution chromosome banding studies were presented in that report. Nevertheless, from the information available the case appears similar, presenting with developmental delay, epicanthic folds, and other characteristics cited by the authors as observed in G-deletion syndrome. Various normal and disease genes have now been mapped to the 22q13→qter region, namely the meningioma chromosome region, β-galactosidase-2, arylsulphatase A, NAOH-diaphorase-1, and preproacrosin (ACK). It would be of interest in such cases as ours to study the enzyme arylsulphatase A (ARSA) activity for example. A reduced activity would be expected, consistent with a gene dosage effect, owing to deletion of one of the two homologous genes.

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Clinically, as well as overall developmental delay, our patient shares many of the features described in the case of Watt et al, namely full cheeks and eyebrows, epicanthic folds, slightly long philtrum, wide, flat nasal bridge, and long trunk. These findings are also frequently seen in r(22) patients who show, in addition, like our case, muscular hypotonia, thick, full lips, ptosis, and large ears with poorly folded helix.