Deletion 2q: two new cases with karyotypes 46,XY,del(2)(q31q33) and 46,XX,del(2)(q36)

ROBERT S YOUNG*, STEVEN D SHAPIRO*, KATHRYN L HANSEN*, L KENNEDY HINE*, DELBERT E RAINOSEK*, AND FERNANDO A GUERRA†

From *the University of Texas Health Science Center, and †Santa Rosa Medical Center, San Antonio, Texas, USA.

SUMMARY  We describe the clinical and cytogenetic findings of two patients with deletions of the long arm of chromosome 2. One has an interstitial deletion identical to that found in a previously reported patient, although they are phenotypically dissimilar. The other patient has a terminal deletion, the first such deletion reported to date.

Although translocations involving chromosome 2 are by no means uncommon,1 to our knowledge isolated long arm deletions of chromosome 2 have been reported in only five patients.2-6 We have recently observed two patients with isolated deletions of the long arm of chromosome 2, one an interstitial deletion and the other most probably a terminal deletion.

Case reports

CASE 1
Case 1 is a 4-month-old male infant, the second child born to a 27-year-old G2, P2 woman and a 27-year-old man. The first child, a female, was delivered by caesarian section because of fetal heart rate deceleration; she is, however, physically and developmentally normal. The family history revealed a 62-year-old maternal great uncle who is mentally retarded from congenital hydrocephalus, a second maternal great uncle who died at birth of unknown causes, and a maternal aunt who died at 6 weeks of age from an unspecified heart defect. The paternal family history is non-contributory.

The antenatal period was unremarkable except for persistent nausea and vomiting from the third to the eighth month, for which Bendectin (doxylamine succinate and pyridoxine hydrochloride) was prescribed. There was also moderate maternal exposure to commercial insecticide fumes during the second week of gestation. There were no other known exposures to potential teratogenic agents of any kind.

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lobes, bilateral iris colobomata, mild microphthalmia and narrow palpebral fissures, left corneal opacity, beaked nose, maxillary alveolar ridge hyperplasia, borderline micrognathia, pilonidal dimple, shawl scrotum, bulbous tip to the penis, complete syndactyly of the 2nd and 3rd fingers of the left hand, bilateral simian creases, syndactyly of the second to the fifth toes bilaterally, and bilateral equinovarus. He had generalised jaundice and a weak cry, but manifested no cardiopathy.

His dermatoglyphics showed an excess of digital whorls. He has six whorls and four ulnar loops, while his parents have ten ulnar loops each. His axial triradius is in the r' position on the left and is absent on the right. A hypothenar radial loop is present on the left palm. He has bilateral hallucal fibular arches and his big toe patterns are a fibular loop on the left foot and an arch on the right foot.

During the first day of life the patient had two episodes of poor feeding with associated vomiting and apnoeic cyanotic spells without bradycardia. A barium swallow was normal except for a dysco-ordination of the soft palate upon swallowing. A chest x-ray revealed infiltrates in the left upper and lower lobes. Intravenous ampicillin and gentamycin plus orogastric tube feeding were begun and the patient improved markedly over the following 2 days. His jaundice resolved quickly with phototherapy and he was discharged aged 11 days. Orogastric tube feedings were discontinued 2 weeks later. A developmental assessment at 11 weeks of life revealed behaviour characteristic of an 8-week-old and was interpreted as normal considering his prematurity.

Cytogenetic evaluation of peripheral blood lymphocytes revealed a consistent interstitial deletion of bands 2q31→q33 in all 50 cells examined (fig 2a). Parental karyotypes were normal. His chromosomal constitution is therefore 46,XY,del(2)(pter→q31::q33→qter) de novo.

**Case 2**

Case 2 (fig 3) is an 8-month-old Hispanic female born to a non-consanguineous 30-year-old G4, P4 woman and a 31-year-old man. All sibs are reported to be physically and mentally normal. The patient's father abandoned the family shortly after her birth, although her mother recalls no member of her husband's family with birth defects, mental retardation, or a history of multiple miscarriages. The same history applies to the maternal side of the family with the exception of one mentally retarded cousin, aetiology unknown.

The mother denied any illness or exposure to unusual chemicals, drugs, or radiation throughout the gestational period. The patient was born by spontaneous vaginal delivery after 37 weeks' gestation by dates. The placenta and amniotic fluid were normal except for slight meconium staining.

Apgar scores were 6 and 9 at 1 and 5 minutes, respectively. She weighed 2300 g (−3 SD), measured 48·3 cm in length (25th centile), and had a head circumference of 30·5 cm (−3·5 SD). Physical
examined revealed a microcephalic infant with
a high pitched weak cry, bilateral epicanthus,
downward slanting palpebral fissures, extremely
long eyelashes, a flat nasal bridge, low set pointed
ears with preauricular protrusion, a cupid’s bow
mouth, a cleft of the soft and hard palates with
absent uvula, inverted nipples, arachnodactyly with
extra mesophageal flexion creases on the fourth
digit of the left hand and digits 2 to 5 on the right
hand, genu varum, and bilateral syndactyly of the
second and third toes. All other systems appeared
clinically normal.

Digital dermatoglyphic patterns include four
arches and six ulnar loops for a total ridge count of
34. This apparent increase in arch patterns must be
tempered by the fact that her mother has six arches.
A vestige pattern is present in the left thenar
eminence. The axial triradial are displaced distally,
located in the t’’’ position on the left and t’ on the
right. She has bilateral big toe arches and bilateral
hallucal distal loops.

Radiographic examination revealed pulmonary
infiltrates consistent with aspiration pneumonia and
a T1 to T8 spina bifida occulta. A barium swallow
was normal, as were her liver function and sweat
cloride tests.

The patient’s course has been complicated by
feeding difficulties secondary to her cleft palate and
two episodes of aspiration pneumonia, for which
she was admitted to hospital. She has developed
seizures which are partially controlled by pheno-
barbitral. Overall, her development, both physical
and mental, is significantly delayed.

Cytogenetic evaluation of peripheral lymphocytes
revealed a small long arm deletion of chromosome 2
in all 50 cells analysed (fig 2b). The deleted segment
appears to be terminal (2q36→qter), although
interstitial deletions of bands 2q33→q34, 2q34→q35,
or 2q33→q35 cannot be ruled out. Her karyotype is
most probably 46,XX,del(2)(qter→q36:). Her
mother’s karyotype is normal, but her father was
unavailable for study. It is unknown, therefore,
whether the patient represents a de novo deletion or
the unbalanced product of a balanced translocation
segregating in her father.

Discussion

From both a clinical and cytogenetic viewpoint
there is very little similarity between the five cases
reported so far and our two patients. The table
presents these seven cases from the most proximal 2q
deletion on the left to the most distal on the right and
lists those features shared by at least two patients
in the series. Two pairs of patients do, however,
have essentially identical cytogenetic defects and
would therefore be expected to resemble each other
phenotypically.

The first pair are the patients reported by Fryns
et al and McConnell et al, who are both mono-
somic for 2q32→q24. They have a number of
features in common, including low birth weight,
relatively small facial features, micrognathia, cleft
palate, great vessel abnormalities, and overlapping
fingers. The second pair with a common deleted
segment are the patient reported by Taysi et al and
and our case 1 who are monosomic for bands
2q31→q33. These two patients are strikingly dis-
similar having only developmental delay and low
set ears as consistent features.

<table>
<thead>
<tr>
<th>TABLE</th>
<th>Clinical features common to two or more patients with 2q deletions.</th>
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<tbody>
<tr>
<td></td>
<td>Antich et al</td>
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<tr>
<td>Deleted segment</td>
<td>q12→q14</td>
</tr>
<tr>
<td>Sex</td>
<td>M</td>
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<tr>
<td>Developmental delay/ mental retardation</td>
<td>+</td>
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<tr>
<td>Low birth weight</td>
<td>+</td>
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<tr>
<td>Sutual irregularities</td>
<td>+</td>
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<tr>
<td>Microcephaly</td>
<td>+</td>
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<tr>
<td>Large/bulging forehead</td>
<td>+</td>
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<tr>
<td>Small facial features</td>
<td>+</td>
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<tr>
<td>Neural tube defect</td>
<td>+</td>
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<tr>
<td>Microphthalmia</td>
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<td>Corneal opacity</td>
<td>+</td>
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<td>Potois</td>
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<tr>
<td>Low set ears</td>
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<tr>
<td>Micrognathia</td>
<td>+</td>
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<tr>
<td>Cleft palate</td>
<td>+</td>
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<tr>
<td>Cleft lip and palate</td>
<td>+</td>
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<tr>
<td>Short neck</td>
<td>+</td>
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<tr>
<td>Cardiopathy</td>
<td>+</td>
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<tr>
<td>Overlapping fingers</td>
<td>+</td>
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<tr>
<td>Long fingers/toes</td>
<td>+</td>
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<tr>
<td>Clefting between toes 1 and 2</td>
<td>+</td>
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</table>
As the table indicates, the phenotype of case 2 most closely resembles that of the patients of Fryns et al and McConnell et al, even though her deletion is considerably more distal than these others. One might anticipate some phenotypic overlap with the patient described by Warter et al if she actually has an interstitial deletion of band 2q35 or 2q36, since Warter's patient is monosomic for bands 2q34→q36, but this is not the case. Based on the cytogenetic and clinical evidence our case 2 appears to be the first reported case of a terminal long arm deletion of chromosome 2.

Given the paucity of 2q deletion patients and the clinical variability typically seen among persons with identical chromosomal abnormalities, it is fruitless to speculate at this point on the '2q monosomy phenotype'.

References


Correspondence and requests for reprints to Dr Robert S Young, Department of Pediatric Dentistry, University of Texas Health Science Center, San Antonio, Texas 78284, USA.
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R S Young, S D Shapiro, K L Hansen, L K Hine, D E Rainosek and F A Guerra

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