Short Communication

Partial duplication of chromosome 20(pter→q12)

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Abstract

Partial duplication of chromosome 20 (20pter→20q12) resulting from a maternally inherited translocation t(14;20)(q11;q13) is described in a female child with neuropsychomotor retardation and multiple congenital anomalies. To our knowledge this is the largest duplication of chromosome 20 that includes segments of both the short and the long arms thus far described in a live-born child.

INTRODUCTION

Numerical and structural abnormalities involving chromosome 20 are extremely rare in live-born babies. Most cases of partial duplications involve the short arm and just a few affect the long arm (Sanchez et al., 1977; Pawlowitzki et al., 1979; Sax et al., 1986). Some partial duplications of chromosome 20 showed involvement of the entire short arm and part of the long arm, associated with small duplications or deletions of other autosomal segments (Krmpotic et al., 1971; Marcus et al., 1979; Rudd et al., 1979; Schinzel, 1980; Delicado et al., 1981).

We describe a girl with partial duplication of chromosome 20 (20pter→20q12) which resulted from a maternally inherited translocation t(14;20)(q11;q13). To our knowledge, this is the largest partial duplication of chromosome 20 described hereto.

CLINICAL REPORT

The patient (Figure 1), a black female child, is the first daughter of healthy, unrelated parents, the mother and father being 23 and 25 years old, respectively, at her birth. The mother’s second pregnancy resulted in a spontaneous abortion after two months of gestation. The girl was born post-term by cesarean section, after an uneventful pregnancy. At birth, weight was 2,450 g and length, 44.5 cm. Her health was good, but she was noted to be hypotonic. Motor development was delayed: she held up her head at 5 months of age and sitting without support occurred only at 10 months.

At 3 years and 6 months of age, the patient presented psychomotor retardation; she could not stand up or walk; there was no speech development and generalized hypotonia was observed. Her height was 77 cm (below the 3rd centile) and weight, 12 kg (3rd centile). She had a round face and a narrow forehead. The frontal bones presented a prominent metopic suture and lateral depressions. Facial dysmorphisms included a mongoloid slant of palpebral fissures, apparent hypotelorism, bilateral convergent strabismus, a depressed broad nasal bridge, a short nose with upturned tip and large nares, a long philtrum, a thin upper lip, and retrognathia. She had a highly arched palate. The ears were low set and posteriorly angulated, with over-folding of the helices. The neck was short, and the abdomeen appeared normal except for an umbilical hernia. The distal phalanges of digits and toes, especially of the thumbs and halluces, were broad. The hands showed single palmar flexion creases. Pes planus and a prominent calcaneum were observed bilaterally. Radiological examination documented thoracic kyphosis. An electroencephalogram at 3 years of age did not reveal any abnormalities.

At 6 years and 9 months of age, the patient was a healthy girl with severe neuropsychomotor retardation. She sat down without support, but could not stand up or speak. Comprehension of simple orders was rather poor, irritability was constant and sphincter control had not developed. Her height was 92.5 cm (below the 3rd centile) and the head circumference, 47.5 cm (below the 2nd centile).

CYTOGENETIC STUDIES

Chromosomal analysis was performed on peripheral blood leukocytes after G banding. In the propositus, a chromosome 14 was replaced by a derivative chromosome, resulting from a translocation of almost the entire long arm of chromosome 14 to the long arm of chromosome 20 at band 20q13.1 (Figure 2a). Examination of parental chromosomes revealed a normal 46,XY paternal karyotype and an abnormal 46,XX,t(14;20)(14pter→14q11.2::20q13.1→20qter;20pter→20q13.1::14q11.2→14qter) maternal karyotype (Figure 2b).

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The karyotype of the child was 46,XX,-14,+der(20),
t(14;20)(q11.2;q13.1)mat, and she was diagnosed as having a
duplication of the segment 20pter→20q12 and a deletion of
14pter→14q11.1.

**DISCUSSION**

The patient’s karyotype was the result of a type 2 ad-
jaent segregation of the translocation chromosomes and
their homologues in maternal meiosis. The material lost
from chromosome 14 comprised the short arm, censtro-
mere and a small pericentric segment of the long arm. The
loss of such segments in Robertsonian translocations does
not cause phenotypic abnormalities so that it is reason-
able to assume that the patient’s clinical picture was the
result of chromosome 20 duplication (20pter→20q12).

The first report of a child with trisomy of chromo-
some 20 (Pan *et al*., 1976) involved a neonate with unusual
facial features and multiple congenital malformations who
died 4 hours after birth. However, Steele (1990) reanalyzed
the chromosomes from a frozen fibroblast culture and iden-
tified the extra chromosome as an isochromosome 12p. In-
deed, based on the clinical findings, Schinzel (1980) had
already suggested that this case represented partial trisomy
of an autosomal segment with a banding pattern similar to
that of chromosome 20. The same explanation would ac-
count for the other presumed trisomy of chromosome 20
reported by Wahlström *et al.* (1976) in a girl who had an
abnormal appearance and cat’s cry at birth, and later on
showed poor weight gain and psychomotor retardation.

A presumptive 20p and partial 20q duplication was
reported by Krmpotic *et al.* (1971), who were unable to
precisely localize the breakpoint on the long arm. More
recently, duplications of chromosome 20, involving the
short arm and the proximal part of the long arm, have been
identified by banding patterns (Marcus *et al*., 1979, Rudd
*et al*., 1979, Schinzel, 1980; Delicado *et al*., 1981). In
these cases, the propositi had an extra-rearranged chro-
mosome 20 that included small segments of other auto-
somes. The 20q duplication affected only the band 20q11
(Marcus *et al*., 1979; Schinzel, 1980; Delicado *et al*.,
1981) or comprised part of 20q12 (Rudd *et al*., 1979).
Our patient had a larger duplication, that included at least
the major part of band 20q12.

Table 1 summarizes the clinical signs of these pa-
tients. Most of these signs are associated with 20p duplica-
tion (for review, see Grammatico *et al*., 1992). It is
noteworthy that the two patients with the largest 20q du-
plications (Rudd *et al*., 1979, and the present case) are
the only individuals with severe growth retardation, mi-
crocephaly and broad distal phalanges of thumbs and toes.

**ACKNOWLEDGMENTS**

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RESUMO

Descrevemos uma duplicação do cromossomo 20 (20pter→20q12), resultante de uma translocação t(14;20)(q11;q13)mat, em uma menina com retardo do desenvolvimento neuropsicomotor e anomalias congênitas múltiplas. Trata-se da mais extensa duplicação do cromossomo 20 presente em indivíduo nascido vivo até agora publicada.

REFERENCES


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Table I - Clinical signs in patients with dup20p and proximal dup20q.

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<td>Deletion 14pter→14q11</td>
<td>dup(?)</td>
<td>12q24.3→qter</td>
<td>13p11→qter</td>
<td>11q25→qter</td>
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<td>male</td>
<td>male</td>
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<tr>
<td>Age</td>
<td>3 6/12 years</td>
<td>13 weeks</td>
<td>13 months</td>
<td>2 10/12 years</td>
<td>6 months</td>
</tr>
<tr>
<td>Birth weight</td>
<td>2,450 g</td>
<td>2,553 g</td>
<td>2,640 g</td>
<td>2,500 g</td>
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<td>Growth retardation</td>
<td>&lt;3rd%</td>
<td>&lt;3rd%</td>
<td>25th-30th%</td>
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<td>Psychomotor retardation</td>
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<td>Speech impediments</td>
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<tr>
<td>Head circumference</td>
<td>&lt;2nd%</td>
<td>&lt;3rd%</td>
<td>&lt;50th%</td>
<td>3rd-10th%</td>
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<tr>
<td>Lateral depression of frontal bones</td>
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<td>Prominent metopic sutures</td>
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<td>Narrow forehead</td>
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<td>Round face and full cheeks</td>
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<td>Thin upper lip</td>
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<td>Broad distal phalanges of thumbs/toes</td>
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<td>Overriding toes</td>
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