this case and that of Gripenberg et al (1972) resemble syndrome I more than syndrome II. An infant recently reported to have monosomy 22 had congenital heart disease but lacked the epicantthus and syndactyly of syndrome II (DeCicco et al, 1973). Hence these distinctions need further refinement and correlation with karyotypes if they are to prove useful for clinical diagnosis.

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REFERENCES


Karyotype 45,XX,-21/46,XX,21q- in an infant with symptoms of G-deletion syndrome I

Summary. An infant with antimongoloid eye slants, achalasia, broad nose, one low-set large and one rudimentary ear lobe, and rudimentary nails with a retarded psychomotor development showed an 45,XX,-21/46,XX,21q- karyotype. By fluorescence and Giemsa staining it was shown that the missing or deleted chromosome 21 was of paternal origin.

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Since Lejeune et al (1964) first reported a case with a mosaic of cells with 45 chromosomes and a missing G chromosome and cells with 46 chromosomes with a deleted G, a number of such mosaic

FIG. 1. The proposita at 6 weeks of age. Note large low-set hypoplastic ear and receding chin.

FIG. 2. The proposita at 6 weeks of age. Note antimongoloid slants, blepharochalasia, and broad nose bridge.
cases have been reported (Greenwood and Sommer, 1971; Berger, 1972). This report presents another
infant with a similar chromosome abnormality and a 45,XX,−21/46,XX,21q− karyotype.

Case report

The proposita was born in April 1973. She is the second child of unrelated parents. The first child is a
healthy 7-year-old girl. The father is 36 years old. He had mumps half a year before conception and worked
with radar for a period of five years until 1962. The mother is 31 years old and has hay fever. She had been
vaccinated during the pregnancy and had been treated with ephedrine. The pregnancy was normal.

The child was born one week before term. She weighed 1650 g and was 41 cm long. The head circum-
ference was 29 cm. The Apgar score was 2/1 min and 4/2 min.

Physical examination revealed a premature infant with a
rudimentary right ear lobe without any external audi-
tory meatus, and a large, low-set left ear lobe (Fig. 1). The nose was broad with a prominent nasal bridge. On
the tip of the nose a dimple was seen and from this
dimple a light streak ran downwards in the midline to the
philtrum. There was an antimongoloid slant of the
eyes and bilateral blepharochalasis (Fig. 2); ophthalmo-
scopy was normal. The mouth was broad; a cleft
 palate was seen in the midline, and the mandible was
small and receding. The hands and fingers were very
slender; on the right hand, the second and fourth finger
overlaid the third finger. The left foot was in a calcaneo-
valgus position and the right foot was inverted. The
nails on all fingers and toes were rudimentary (Fig. 3).
A rough systolic murmur was heard to the left of the
sternum.

Skeletal radiology revealed 13 ribs on the left side and
hemivertebrae at the 10th and 11th thoracic vertebrae.

Psychomotor development has been very slow. At 4
months she weighed only 3000 g and was severely re-
tarded.

The peculiar appearance of the infant prompted a
chromosome analysis shortly after birth.

| TABLE I |
| CHROMOSOME COUNT DISTRIBUTION |

<table>
<thead>
<tr>
<th></th>
<th>44</th>
<th>45</th>
<th>46</th>
<th>47</th>
<th>Total No. of Cells Examined</th>
<th>Karyotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proposita Blood</td>
<td>2</td>
<td>47</td>
<td>11</td>
<td>11</td>
<td>60</td>
<td>45,XX,−21/46,XX,21q−</td>
</tr>
<tr>
<td>Skin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>50</td>
<td>45,XX,−21/46,XX,21q−</td>
</tr>
<tr>
<td>Mother Blood</td>
<td>20</td>
<td></td>
<td></td>
<td></td>
<td>20</td>
<td>46,XX</td>
</tr>
<tr>
<td>Father Blood</td>
<td>1</td>
<td>54</td>
<td>1</td>
<td></td>
<td>57</td>
<td>46,XY</td>
</tr>
<tr>
<td>Sister Blood</td>
<td></td>
<td></td>
<td></td>
<td>50</td>
<td>50</td>
<td>46,XX</td>
</tr>
</tbody>
</table>

* 47,XXY and 47,XY,+G.
fibroblasts the 46 line with the deleted G was the predominant line (Table I). The breakpoint of the deleted 21 chromosome was determined by fluorescence and Giemsa banding to be located at q22 close to the medium intense band. The deleted chromosome had non-fluorescing satellites which were well-stained by Giemsa. The mother and the sister had normal karyotypes. The father had two hyperdiploid cells, one with an XXY karyotype and one with a supernumerary G chromosome (Table I). A comparison of parental cells with the two cell lines from the proposita showed that the respectively missing, or deleted 21 chromosome was of paternal origin. One of the father's 21 chromosomes showed non-fluorescing but Giemsa stained marker satellites. This chromosome was missing in the proposita's cell line with 45 chromosomes and it was deleted in the 46 cell line (Fig. 4).

**Discussion**

Only few G-deletion cases have so far been analysed by banding methods. They have shown a variety of chromosome abnormalities. Mosaics, deletions, and rings of both chromosome 21 and chromosome 22 were found (Table II). There is a considerable difference in the amount of chromosome material lost and the clinical pictures are widely different. However, two clinical syndromes

**TABLE II**

**BANDING STUDIES IN DELETION SYNDROMES**

<table>
<thead>
<tr>
<th>Reference</th>
<th>Karyotype</th>
<th>Deletion Syndrome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crandall <em>et al</em> (1972)</td>
<td>46,XY,r21</td>
<td>I</td>
</tr>
<tr>
<td>Case 1</td>
<td>46,XY,r22</td>
<td>II</td>
</tr>
<tr>
<td>Case 2</td>
<td>46,XY,r22</td>
<td>II</td>
</tr>
<tr>
<td>Case 3</td>
<td>46,XY,r22</td>
<td>II</td>
</tr>
<tr>
<td>Gripenberg <em>et al</em> (1972)</td>
<td>45,XX, - 21</td>
<td>?</td>
</tr>
<tr>
<td>Magenis <em>et al</em> (1972)</td>
<td>46,XY,r21</td>
<td>I</td>
</tr>
<tr>
<td>Case 1</td>
<td>46,XY,r22</td>
<td>II</td>
</tr>
<tr>
<td>Case 2</td>
<td>46,XY,r22</td>
<td>II</td>
</tr>
<tr>
<td>Richmond <em>et al</em> (1973)</td>
<td>46,XX/45,XX, - 21 (blood)</td>
<td>I</td>
</tr>
<tr>
<td>46,XX/46,XX,rG (skin)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shibata <em>et al</em> (1973)</td>
<td>45,XX, - 21/46,XX,r21</td>
<td>I</td>
</tr>
<tr>
<td>Stoll <em>et al</em> (1973)</td>
<td>46,XX,r22</td>
<td>II</td>
</tr>
<tr>
<td>Warren <em>et al</em> (1973)</td>
<td>46,XY,r21</td>
<td>I</td>
</tr>
<tr>
<td>Case 1</td>
<td>46,XY,r22</td>
<td>II</td>
</tr>
<tr>
<td>Case 2</td>
<td>46,XY,r22</td>
<td>II</td>
</tr>
<tr>
<td>Case 3</td>
<td>46,XY,r22</td>
<td>II</td>
</tr>
</tbody>
</table>
may be recognized: the clinical picture of the ‘antimongolism syndrome’ (Lejeune et al, 1964) or deletion syndrome I (Warren and Rimoin, 1970) which shows hypertonia, blepharochalasia, antimongoloid slants, large poorly lobulated ears, microretrognathia, retarded growth, and mental retardation. All cases with a clear cut deletion of chromosome 21 or a mosaic with a missing 21 have shown most of these symptoms.

The clinical symptoms were less characteristic in cases where chromosome 22 was affected. Hypertonia, epicantus, ptosis, syndactyly, and mental retardation were frequently observed (Reisman et al, 1967; Berger 1972; Warren, Rimoin, and Summitt, 1973). Complicated rearrangements with G chromosomes with additional or missing chromosome material were reported (Rethoré et al, 1972; Koivisto et al, 1973). As well as a missing chromosome 21 an unidentified small metacentric chromosome was also found in Koivisto’s case. The clinical picture, therefore, can not be compared with the two deletion syndromes.

Only a few cases of apparently pure monosomy G are known (Thorburn and Johnson, 1966; Al-Aish et al, 1967; Hall, Fredga, and Svenningsen, 1967; Böhm and Fuhrmann, 1969; Gripenberg, Elfving, and Gripenberg, 1972). Only in Gripenberg et al’s case was the missing chromosome identified by special staining as chromosome 21. The other cases were examined before special staining methods were available. Only lymphocytes were examined in the cases of Thorburn and Johnson and of Böhm and Fuhrmann. In Böhm’s and Fuhrmann’s case 4% normal cells were found in blood culture. In both cases, mosaicism with a normal cell line might have been present. The clinical picture of these cases showed considerable variability. The minor symptoms found in the cases of Al-Aish and Gripenberg might be explained by a hidden translocation of a small part of chromosome 21 material onto another chromosome. Such a case was demonstrated by Rethoré et al (1972); unfortunately no clinical description was given stating only that the symptoms were different from the deletion syndrome I. In Rethoré’s case the distal end of a chromosome 21 was translocated onto the short arm of chromosome 9. While in deletion syndrome I, the distal part of chromosome 21 is missing, in the translocation case the proximal part of chromosome 21 was missing. This type of anomaly is impossible to detect with conventional chromosome techniques.

A thorough clinical examination with a comparison of cytogenetic findings with special staining methods may add to our knowledge of phenotype-genotype association and may establish with certainty two or even more G-deletion syndromes.

The expert technical assistance of Mrs Hanne Poulsen and Mrs Birthe Jespersen is gratefully acknowledged. This study has been supported by a grant from the Research Committee of the Danish Mental Retardation Service.

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References


The trisomy 8 syndrome: two additional mosaic cases*

Summary. Two patients with trisomy 8 mosaicism, confirmed by trypsin-Giemsa banding are described. While the majority of patients with this aneuploidy have been mosaics, the phenotypes of the complete and mosaic trisomies closely resemble each other. Mosaic trisomy 8 results in specific clinical findings which include skeletal dysplasia, particularly absent patellae, deep furrowing of the soles of the feet, and periarticular changes resulting in camptodactyly and progressive limitation of joint mobility.

Trisomy C has been described in approximately 30 patients and as expected, the phenotypes have shown considerable variation. Identification of the particular C-group chromosome involved was not possible until the advent of the chromosome banding techniques. With these techniques, a specific trisomy 8 syndrome has emerged and eight patients with this syndrome have been described (de Grouchy, Turleau, and Léonard, 1971; Bijlsma, Wijffels, and Tegelaers, 1972; Caspersson et al., 1972; Kakati, Nihill, and Sinha, 1973). We would like to report two additional cases of trisomy 8 mosaicism, confirmed with trypsin-Giemsa banding, both of whom showed specific physical changes.

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FIG. 1. Case 1. Note dysplastic ear, minimal micrognathia, and severe acne.
Karyotype 45,XX,−21/46,XX,21q in an infant with symptoms of G-deletion syndrome I
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