Familial Transmission of a Chromosomal Translocation t(2q+;Cp−)

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Reciprocal translocations involving duplications of different regions of chromosomes of group C have been reported in patients with multiple congenital malformations. Some of these cases died at 2 or 3 months of age (de Grouchy et al., 1967a, b; Gray et al., 1966). Others were discovered at ages ranging from 12 months to 12 years (Edwards et al., 1962; Rohde and Catz, 1964; de Grouchy and Canet, 1965; Lindsten et al., 1965; Punnett et al., 1966; Lejeune et al., 1966; Lord, Casey, and Laurance, 1967). Each case had different phenotype probably because they represent partial trisomies for different regions of different chromosomes. Therefore, reports of all the new cases and attempts to identify the chromosomes involved will contribute to characterize new syndromes when enough patients with duplications for similar regions of the same chromosome become known.

In this paper, we present a detailed description of the multiple congenital anomalies observed in a stillborn child, who had a duplication for almost all the short arm of one chromosome C (probably No. 8) and a small deletion for the long arm of a No. 2. The father of this child was found to be the balanced translocation heterozygote, which may be written 46, XY,t(2q+;Cp−) according to the nomenclature adopted at the Chicago Conference (1966).

Case Report

The propositus was the first child of a 27-year-old Negro mother and a 29-year-old Negro father. Pregnancy was normal. The infant was delivered by caesarean section because of cephalo-pelvic disproportion. At delivery he showed symptoms of respiratory depression. Resuscitation was attempted but the child died. Birthweight was approximately 3.2 kg. Gross examination revealed a flat occiput, mongoloid slanted eyes, an incarcerated right inguinal hernia, and disarticulation of both knees and hips.

External examination of the infant at necropsy revealed meconium over the skin and several prominent deformities. The head had a flat occiput, malformed external ears, bilateral palpebral oedema, haemorrhagic sclerae, and submucosal haemorrhage in the lips (Fig. 1a). The caput succedaneum was present over the right posterior parietal area. The upper extremities were essentially normal except for the hands, which had abnormal finger-nails, a palmar simian crease, a single crease of the fifth finger, and a high axial triradius (Fig. 1b). The trunk was normal. However, a bulge due to a right inguinal hernia was noted. In the lower extremities, both knees had dislocation of the tibia anterior to the condyles of the femur. Both ankles were dislocated laterally (Fig. 1c).

Internal examination of the infant disclosed multiple petechiae over the surface of the thymus, lungs, and heart. The right lobe of the liver was paler than the left. The ductus arteriosus measured 4 mm. in diameter and its intimal surface was wrinkled. The cranial sutures had fused and the bone of the skull was calcified. The brain itself had a communicating hydrocephaly (Fig. 1d).

Histological studies showed aspiration of amniotic debris in the lungs, fatty metamorphosis in the right lobe of the liver, focal hepatic necroses, and hyperplasia of the islets of Langerhans.

In review, the multiple deformities of the infant caused protracted labour and death from the resultant intrauterine anoxia.

Cytogenetic findings. Chromosome analyses were made from blood leucocyte cultures, using a slight modification of the method of Moorhead et al. (1960). The blood for the study of the propositus was obtained 10 hours after death by puncture of the heart. Microscopical examination of 75 well-spread metaphases showed a modal number of 46 chromosomes in 71 cells. Two cells had 45 chromosomes and two had 47 chromosomes. All the cells, including those with abnormal chromosome counts, showed that one chromosome, No. 2, was unusually long (Fig. 2). The analysis of 13 karyotypes confirmed the difference in the size of chromosome No. 2. The two cells with 45 chromosomes presented a random loss of one chromosome of groups E and G, respectively. The two cells with 47 chromosomes showed an extra chromosome similar to those of group D. The abnormal length of one of the No. 2 chromosomes suggested the possibility of this chromosome being one product of a reciprocal translocation.

Karyotype analysis was performed on two blood cul-
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The two translocated chromosomes were measured and compared to the measurements of the chromosomes of group A and some chromosomes of group C. Chromosome measurements were made on photographic enlargements of 20 karyotypes, 10 from the propositus and 10 from the father, and the relative chromosome lengths were expressed as percentages of the haploid autosomal complement. The data are presented as a mean ± one standard error (Table I). Differences of means were tested for statistical significance by the t-test and p values were calculated.

The following relationships can be seen from the data:

1. The sum of the total length of the two translocated chromosomes is equal (p = 0.6) to the sum of the length of the normal chromosomes No. 2 and one chromosome C presumably classified as No. 8.

\[ 2c (9.82 \pm 0.12) + C2 (3.66 \pm 0.06) = 13.48 \]
\[ 2l (8.46 \pm 0.14) + 8 (4.99 \pm 0.06) = 13.45 \]

2. No significant differences (p = 0.4) exist between the short arm of 2c and the short arm of No. 2, or between the long arm of C2 and the long arm of No. 8:

\[ 2c \text{ s.a.} (3.20 \pm 0.05) = 2 \text{ s.a.} (3.24 \pm 0.09) \]
\[ C2 \text{ l.a.} (3.36 \pm 0.06) = 8 \text{ l.a.} (3.44 \pm 0.05) \]

3. The long arm of 2c is about equal (p = 0.50) to the sum of the long arm of No. 2 plus the short arm of No. 8, less the short arm of C2:

\[ 2l (5.22 \pm 0.09) + 8 \text{ s.a.} (1.54 \pm 0.05) - C2 \text{ s.a.} (0.29 \pm 0.009) = 6.48 \]
\[ 2c \text{ l.a.} = 6.62 \pm 0.09. \]

Fig. 1. Photographs of the propositus: (A) facies with malformed external ears and palpebral oedema, (B) hand with abnormal finger-nails and palmar simian crease, (C) infant at necropsy—note dislocation of both knees and ankles, (D) macroscopical sections of the brain showing communicating hydrocephaly.
These measurements and the above relationships suggest that the chromosome C involved in the translocation may be one chromosome 8. We have classified as No. 8 the chromosome which appeared to be less metacentric than those considered as No. 7 and 11, and more metacentric than chromosomes 9 and 10. Unfortunately, autoradiographic studies were not carried out. These, if combined with the measurements, might

![Metaphase and karyotype of propositus.](image)

Note that the long arm of one of the chromosomes No. 2 is significantly longer than the other member of this pair.

| TABLE I |
|------------------------------|----------------------------------|
| AVERAGE CHROMOSOME MEASUREMENTS* FROM 20 CELLS: 10 FROM PROPOSITUS, 10 FROM FATHER |

<table>
<thead>
<tr>
<th>Chromosome No.</th>
<th>Father of Propositus</th>
<th>Propositus</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Long Arm</td>
<td>Short Arm</td>
</tr>
<tr>
<td>1</td>
<td>4.49 ± 0.04</td>
<td>4.48 ± 0.09</td>
</tr>
<tr>
<td>2</td>
<td>5.22 ± 0.09</td>
<td>3.24 ± 0.09</td>
</tr>
<tr>
<td>3</td>
<td>3.39 ± 0.06</td>
<td>3.38 ± 0.05</td>
</tr>
<tr>
<td>4</td>
<td>3.44 ± 0.05</td>
<td>1.54 ± 0.05</td>
</tr>
<tr>
<td>5</td>
<td>3.35 ± 0.06</td>
<td>1.38 ± 0.02</td>
</tr>
<tr>
<td>6</td>
<td>3.14 ± 0.06</td>
<td>1.27 ± 0.04</td>
</tr>
<tr>
<td>7</td>
<td>6.62 ± 0.09</td>
<td>3.20 ± 0.05</td>
</tr>
<tr>
<td>8</td>
<td>3.36 ± 0.06</td>
<td>0.29 ± 0.009</td>
</tr>
</tbody>
</table>

*Mean values and standard errors of arm measurements expressed as percentages of haploid autosomal complement.
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**Family Data.** The proband was the product of the first pregnancy of phenotypically normal parents. Chromosome studies show that the father is the balanced carrier of the translocation and the mother is normal. The only sib of the father, a sister, is married, but she has had no children or abortions. Her karyotype is normal. The paternal grandparents are also phenotypically normal. No family history of abortions or malformations could be ascertained in other members of the family. Chromosome studies have not been performed on the paternal grandparents or other members of the family because of lack of co-operation.

**Discussion**

The genetic constitution in the progeny of a balanced translocation heterozygote depends on the meiotic segregation of four chromosomes: the two translocated chromosomes and their corresponding unaltered normal homologues. These chromosomes form a cross-shaped configuration (Fig. 5), during pairing at pachytene of the first meiotic division. The translocation here described is very asymmetrical, and the most likely configuration at metaphase 1 is a chain IV. Three kinds of segregation could occur: alternate, Adjacent I, and Adjacent II. The results observed in maize and drosophila have shown that the relative frequency of each type of segregation is related to the size of the interchanged segment and the length of the two...
chromosomes involved (Burnham, 1950, 1956, 1962). In our case, the two chromosomes affected are long and the translocated segments are relatively small. We would expect, therefore, that alternate segregation would be more frequent than Adjacent I, whereas the occurrence of type Adjacent II would be very low.

Six types of gametes may be formed by the three types of segregation. After fertilization, the combination of these gametes with normal ones would give rise to six kinds of zygotes, as shown in Table II. Two of these combinations involve duplication or deletion for the largest portion of a long autosome, No. 2, and both are known to be lethal in humans (Hall and Källén, 1964; Kelly et al., 1965; Kerr and Rashad, 1966). Furthermore, these two kinds would be derived from Adjacent II type of segregation which is expected to be infrequent in the present case. Therefore, four classes are likely to be found among the offspring of a balanced carrier such as the father of the propositus: the normal, the carrier, the deletion of the small arm of one chromosome C with a small duplication of chromosome No. 2, and the duplication of the small arm of the chromosome C with a small deletion of the long arm of a No. 2. The last class was found in the propositus. The severity of the defects observed in this child suggests that the duplication here described might result in abortion, stillbirth, or neonatal death.

The clinical manifestations caused by the deletion of the short arm of chromosomes of the C

**TABLE II**

*Gametes and possible zygotes that would be formed by each type of segregation*

<table>
<thead>
<tr>
<th>Gametes</th>
<th>Alternate</th>
<th>Zygotes</th>
<th>Adjacent</th>
<th>II</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2C</td>
<td>2C</td>
<td>2C</td>
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<td>2C</td>
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</table>

*Proband.*
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Fig. 5. Cross configuration in prophase of meiosis I.

lateral dislocation of the ankles. The chromosomes involved in the translocation were identified visually and by measurements. The father was the carrier of the balanced form of the translocation. The meiotic behaviour of the translocated chromosomes and the genetic risk of the balanced carrier are discussed.

The authors thank Mrs. Carolyn Meiller for valuable technical assistance, and Mrs. Helen Carroll for help in the statistical evaluation of chromosome measurements. We are also most grateful to Dr. E. Chu for the review of the manuscript.

REFERENCES


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doi: 10.1136/jmg.6.2.174

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