Familial Occurrence of Congenital Malformations and Ring Chromosome (46,XX,Cr)

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A ring chromosome in group C has been described only rarely (Atkins et al., 1960; Turner et al., 1962; Butler, France, and Jacoby, 1967). We here report a family where there have been numerous miscarriages and congenital malformations, and in one member of which a ring chromosome of the C group was detected.

Case Reports

The pedigree (Fig. 1) shows that in the first and second generations no aberration was found; it is only in the third generation that miscarriages, prematurity, and congenital malformations appear.

II.3. A healthy female was pregnant nine times but has only one healthy living child; two of her children had congenital malformations, one of these was stillborn, the other died a few days after birth. The other 6 pregnancies resulted in miscarriages.

III.9. Stillborn at full-term. Apart from congenital heart disease (agenesis of pulmonary valve), no other abnormality was found.

III.13. Premature (2400 g.). Died after 16 days. Necropsy revealed jaundice, congenital heart disease (atrial and ventricular septal defect, aortic hypoplasia), and polycystic kidney.

II.4. The mother of the proposita is healthy and has three normal boys. The third pregnancy resulted in twins (III.16, III.17).

II.5. The father of the proposita. No congenital malformation was found in him or in his family.

III.16. The 2-year-old girl (Fig. 2), birthweight 2800 g., height 82 cm., weight 8600 g., was microcephalic, with head laterally flattened, and protruding beak-like frontal bone. Slanting eyes, hypertelorism, epicanthus, horizontal nystagmus discoloured papillae. Ears low-set and dissimilar. Palate gothic, teeth regular, 6/6. Small, wide-apart nipples. Labia majora hypoplastic.

On both sides between toes II and III there was partial syndactyly. The skin was poorly pigmented. Psycho-motor development was delayed and mental development was greatly retarded. The urinary excretion of amino acids was normal on thin layer chromatography.

III.17. This twin sister of III.16, with birthweight 3000 g., died at the age of 2 months. The right side of face was normal, left side was hypotrophic with the following malformations: cleft lip, cleft palate, dysplasia of the mandible, microphthalmia, no external auditory canal; in place of the auricle there was a small bud of skin. There was a double thumb on the right side.

II.6. A healthy female; all her four pregnancies ended in miscarriages.

Cytogenetic Studies

The chromosome studies were performed on peripheral leucocyte culture using Moorhead's slightly modified method (Moorhead et al., 1960; Schuler, 1965). In III.16 (Fig. 3, Table I) most of the cells had a 46,XX,Cr chromosomal constitution. In the cells containing 45 chromosomes the ring chromosome was

<table>
<thead>
<tr>
<th>Case No.</th>
<th>No. of Cells Counted</th>
<th>45,Cr</th>
<th>45</th>
<th>46,XX,Cr</th>
<th>46</th>
<th>47</th>
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</thead>
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<tr>
<td>III.16</td>
<td>80</td>
<td>3</td>
<td>5</td>
<td>70</td>
<td>2</td>
<td>47</td>
</tr>
<tr>
<td>II.3</td>
<td>31</td>
<td>3</td>
<td>3</td>
<td>24</td>
<td>3</td>
<td>47</td>
</tr>
<tr>
<td>II.4</td>
<td>20</td>
<td>2</td>
<td>2</td>
<td>18</td>
<td>18</td>
<td>47</td>
</tr>
<tr>
<td>II.5</td>
<td>26</td>
<td>2</td>
<td>2</td>
<td>24</td>
<td>19</td>
<td>47</td>
</tr>
<tr>
<td>II.6</td>
<td>20</td>
<td>2</td>
<td>2</td>
<td>18</td>
<td>19</td>
<td>47</td>
</tr>
</tbody>
</table>

Table I

<table>
<thead>
<tr>
<th>Cells Scored</th>
<th>Labelled Cells</th>
<th>Late Replicating X</th>
<th>Labelling Over Ring Chromosome</th>
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</thead>
<tbody>
<tr>
<td>No. of cells</td>
<td>153</td>
<td>73</td>
<td>19</td>
</tr>
</tbody>
</table>

TABLE II

Numerical Data of Autoradiography

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Fig. 1. Pedigree of the family.

Fig. 2. Photograph of proposita.

Fig. 3. Karyotype of proposita (46,XX,Cr).
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Fig. 4. Autoradiography of group C chromosomes of two cells.

Discussion

A 2-year-old girl is described with multiple congenital malformations in whom most of the cultured peripheral blood cells contained a ring chromosome. As regards its size and the arrangement of other normal chromosomes, it belongs to the longer members of group C. Considering the sex chromatin positivity, the symptoms, and the autoradiography, the possibility of its development from the X chromosome can be excluded.

The clinical picture in other cases of a ring C chromosome differed from ours and from each other. Though the first case of Atkins et al. shows some similarities to the patient described here (low-set ears, epicanthus, gothic palate, nystagmus, mental retardation), we do not think that the presence of these symptoms would be sufficient to assume that the ring chromosome had originated in both cases from one and the same member of group C. The symptoms in our case were also different from those that have been presented up to now in connexion with other aberrations of chromosomes in the C group. Since no chromosomal aberrations were found in other members of the family, we cannot explain the great number of congenital malformations and miscarriages.

Summary

An infant is described with multiple congenital malformations who was shown to have a ring chromosome in group C. In the family there have been numerous cases of congenital malformations and miscarriages. The co-twin of the proposita was also affected. In the healthy members of the family who were examined, no chromosomal aberration was found.

References


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