The Phenotypic and Cytogenetic Spectrum of Partial Trisomy 9

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A new patient with trisomy for the chromosome segment 9pter→q22 is compared to 19 previously reported cases of partial trisomy 9. Manifestations such as microcephaly, prominent nasal root, bulbous nose, and down-turned corners of the mouth are common to patients with trisomic segments extending from 9p21 to 9q13, while intra-uterine growth retardation, cleft lip/palate, skeletal anomalies, and heart defects are more common with trisomic segments extending through 9q22-9q32. A graphic method illustrates this progression in the partial trisomy 9 malformation spectrum as the triplicated chromosome region extends from bands 9p21 to 9q32. More severe and random defects are observed with complete trisomy 9 or tetrasomy 9p, suggesting an extreme excess of material greatly increases developmental variability.

Key words: Rethoré syndrome, partial Trisomy 9, multiple congenital anomalies/mental retardation (MCA/MR) syndrome

INTRODUCTION

Since the description by Rethoré et al [1973] of a characteristic syndrome due to partial trisomy 9, over 150 reports of partial [Young et al, 1982] or complete [Annerén and Sedin, 1981] trisomy 9 have appeared. We report on a new case of partial trisomy 9 and discuss the phenotypic and cytogenetic correlations now possible for trisomy 9 disorders.

CLINICAL REPORT

The propositus was born at term after an uncomplicated pregnancy to a 36-year-old mother with a history of 5 first trimester abortions by her current husband and 2
healthy children by a previous marriage. At 10 hours the infant had a weight of 2.0 kg (<3rd centile) and a length of 44 cm (<3rd centile). There was microcephaly (OFC 30.5 cm, <3rd centile) with abnormally wide sutures. The child had an abnormal facial appearance (Fig. 1A) with an antimongoloid slant of palpebral fissures, bulbous nose, left unilateral cleft lip and palate, and proportionate hypertelorism with an inter-pupillary distance of 3.8 cm (50th centile). The ears were antverted and anteflexed with prominent helices. There was no cardiac murmur and the kidneys had a normal contour. There was a simian crease on the right palm and a single crease on each fifth digit with clinodactyly. There were simple arch patterns on all fingers and hypoplastic nails on hands and feet. The feet had a rocker-bottom appearance with metatarsus adductus bilaterally. The serum immunoglobulin A level was less than 3.8 mg% and the immunoglobulin M level was 5.1 mg% (normal). Skeletal survey: bilateral acromioclavicular synostosis and calcifications in the right upper quadrant suggestive of meconium peritonitis. A chest radiograph showed an enlarged heart with prominent pulmonary vasculature. The child died at 36 hours due to cardiorespiratory failure; autopsy was not permitted.

RESULTS AND DISCUSSION

Figure 1B shows the patient’s 9 chromosomes as obtained from the peripheral blood karyotype (47,XX,+9pter→q22:) using GTG staining. The extra chromosome was identified as 9pter→q22: based on its asymmetric appearance by both GTG and CTG banding. The mother’s chromosomes were normal; the father refused to be studied.

As summarized in Table I, the patient’s manifestations resemble those of 19 previous cases of partial trisomy 9, of which 8 were de novo occurrences and 11 occurred by 3:1 nondisjunction from a maternal translocation. The extra chromosome in our case may have been derived from a paternal translocation based on the history of miscarriage but the father refused analysis. The preponderance of females (14 of 20) with partial trisomy 9 has been noted as a general characteristic of partial trisomy 9 [Young et al, 1982] suggesting that males are generally more severely affected with higher likelihood of prenatal death.

A spectrum of clinical severity which correlates with the extent of triplicated chromosome 9 material can be derived by inspection of Table I. Mental retardation is virtually a constant feature. Lewandowski et al [1976] emphasized that partial trisomy 9pter→p21: patients have mild facial anomalies and few skeletal or visceral anomalies. The summary of partial trisomy 9pter→p11: patients provided by Young et al [1982] includes the patients of Lewandowski et al [1976] and a slight background of phenotypic effects due to unbalanced reciprocal translocations; this large number of patients has the typical face described by Rethoré et al [1973], i.e., microcephaly, enophthalmos, hypertelorism, antimongoloid slant of palpebral fissures, broad nasal root with a bulbous nasal tip, down-turned corners of the mouth, and anomalous ears. Partial trisomy 9pter→q11-13: patients also have the typical face seen in the Rethoré syndrome with an increased incidence of skeletal and heart defects. Tetrasomy 9p patients [Balestrazzi et al, 1983], have a similar facial appearance but are more severely affected; 33% have died in the first year and six of 10 had hydrocephalus (Table I). Partial trisomy 9pter→q22-32: patients, as exemplified by our propositus, have the characteristic face with increased incidence of intrauterine growth retarda-
tion, cleft lip/palate, micrognathia, cardiac anomalies, rocker-bottom feet, sacral dimple, and talipes equinovarus or congenital hip dislocation. The two cases of partial trisomy 9q reported by Turleau et al. [1975] had a very different face from trisomy 9p patients; this influence of the distal 9q region may explain why complete or mosaic trisomy 9 patients often do not show the characteristic facies of patients with Rethoré syndrome. Trisomy 9 patients are more severe and variable in their phenotype than partial trisomy 9 patients, and there is frequent mosaicism. Because of their growth retardation and dermatoglyphic arch patterns [Young et al., 1982], severe cases of partial trisomy 9 may be initially confused with trisomy 18 patients.

The catalogue of triplication 9 patients summarized in Table I is perhaps the best opportunity in man for matching clinical manifestations with a progressive increase of extra chromosome material. Using these data, the incidence of defects is graphically related to the extent of trisomic chromosome 9 in Fig. 2. Each square of the graph is shaded according to the percent of patients having a given defect. At the right is shown the mean percent for the five patient categories; the resulting hierarchy of defects represents the malformation spectrum for partial/complete trisomy 9 patients as a whole. Using this graphical method, there is an obvious progression in malformation spectrum with triplicated chromosome length through band 9q32. The apparent decline in malformations for complete trisomy 9 patients in part reflects the different phenotypic influence of the 9q terminal region described above [Turleau et al., 1975], but also the wide variety of less common defects in these patients which are not listed in Table I. Unusual malformations reported in patients with trisomy 9
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<sup>1</sup>Lewandowski et al [1976].
<sup>2</sup>Young et al [1982].
<sup>3</sup>Baccichetti et al [1979].
<sup>5</sup>Baccichetti et al [1979], Centerwall et al [1975], Fujita et al [1976], Kushnick and Adessa [1976], Neu et al [1979], Schwanitz et al [1974].
<sup>6</sup>Turleau et al [1975].
<sup>7</sup>Annerén and Sedin [1981], Romain and Sullivan [1983].
<sup>8</sup>Annerén and Sedin [1981], Katayama et al [1980], Sanchez et al [1982], Wilson and Barr [1983].
<sup>9</sup>Trait present/trait mentioned or excluded.
<sup>10</sup>These cases had hypotelorism.

<sup>1</sup>These cases had upslanting palpebral fissures.
<sup>2</sup>Hydrocephalus.
<sup>3</sup>Sacral dimple.
include cranial asymmetry and malrotation [Wilson and Barr, 1983], cloverleaf skull and leukomalacia [Katayama et al, 1980], perineal hypoplasia [Annerén and Sedin, 1981], ambiguous genitalia [Sutherland et al, 1976], and many others. In trisomy 9 and tetrasomy 9p patients an extreme of excess chromosome material seems to be reached that increases the variety and severity of malformation. The progression of findings illustrated by partial/complete trisomy 9 patients supports the hypothesis of Shapiro [1983] that one effect of extra chromosome material is to increase the range of variation permitted for embryologic and homeostatic mechanisms.

REFERENCES


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