Clinical outcomes of adjacent 1 segregation in a familial translocation t(8;18)(p21.3;p11.23)

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Abstract
We report a reciprocal translocation t(8;18)(p21.3;p11.23) in which both unbalanced products of adjacent 1 segregation were observed within a family. The proband was originally referred because of short stature and a webbed neck, but further investigations showed that she had mental retardation and a congenital heart defect, and had inherited an unbalanced form of the maternal translocation, 46, XX, der(8)t(8;18)mat. The proband's sister spontaneously aborted an 11 week fetus with multiple major system malformations, which was found to have a 46,XY, der(18)t(8;18)mat karyotype. The phenotypic findings of the affected subjects are discussed.

Key words: adjacent 1 segregation; chromosome 8; chromosome 18.

Balanced reciprocal translocations occur with a frequency of approximately 1 in 658 in the general population, and the breakpoints involved are nearly always unique, one notable exception being the t(11;22) translocation. However, it is often difficult to advise carrier couples on the a priori risk of giving birth to an abnormal child following malsegregation of a translocation. In an attempt to resolve this difficulty, Jalbert et al. described a means by which the least unbalanced malsegregation of a translocation could be calculated, so that those gametes containing the least amount of duplication/deletion material, and therefore most likely to produce unbalanced and perhaps viable concepures, could be predicted. Nevertheless, empirical observations of the malsegregants of balanced translocations provide the most compelling evidence for the relationship between the amount and content of genetic imbalance and phenotypic effects. We report a family in which an 8;18 translocation segregated by both alternate and adjacent 1 modes. In the resulting unbalanced family members, the amount of deletion in one is equivalent to the extent of the duplication observed in the other. The proband with the derived 8 chromosome was not cytogenetically investigated until 15 years of age, whereas a fetus with the derived 18 spontaneously aborted at 11 weeks' gestation.

Materials and methods
CASE REPORT
The proband (fig 1, III-7) was first investigated cytogenetically aged 15 years, and was originally referred because of a webbed neck, short stature, and menorrhagia. A more detailed clinical history showed that at 8 weeks of age she had a correction of a tetralogy of Fallot with a transannular patch, and at that time was found to be microcephalic. At 13 years of age epileptiform seizures began and she was noted to be mentally retarded; a brain scan showed no structural abnormalities, although it was felt that the pattern of cortical sulcation might be accentuated. More recently she was referred to a cardiologist after an episode of retrosternal chest tightness, shortness of breath with pallor, and peripheral cyanosis brought on by climbing a hill near her home. On examination she was noted to be very obese and a chest x ray showed cardiomegaly. She is currently being investigated for long term pulmonary regurgitation with a degree of right ventricular dysfunction.

Her phenotypically normal older sister (fig 1, III-2) was counselled in early pregnancy that there was a 10 to 20% risk of her having a liveborn aeploid offspring and she declined the offer of prenatal diagnosis. She had a spontaneous abortion (fig 1, IV-1) at 11 weeks' gestation. Necropsy showed a severely macerated fetus, crown-rump 20 mm, crown-head 22 mm, consistent with 9 weeks' gestation. The head was small in comparison to the length of
the body, the upper and lower limb buds were present, but only at the stage of finger and toe rays. Pigmented eyes were present, but there was no clear evidence of nasal and lip development. Although the posterior neuropore was closed, there was very delayed development of the cranial end of the CNS and of the thoracic organs, although the liver, adrenals, kidneys, and gut were clearly identified. Owing to some disruption of the fetus, probably arte-factual, it was uncertain if there were any anterior thoracic or abdominal wall defects. A cystically dilated segment of umbilical cord was identified which contained only two vessels.

The proband's mother (fig 1, II-4) has also had four miscarriages (III-3–6) all between 6 and 12 weeks of gestation. She was one of twins, the other (II-3) apparently died during early pregnancy. Her two male sibs had also died. One (II-1) was born at 28 weeks' gestation and died at 24 hours, which at that time may have been because of prematurity, and the other brother (II-2) was born at term and died aged 18 months. The wife of her maternal cousin (II-6) has also had two early miscarriages.

Results

The results are shown in the table. The absence of a maternally inherited allele at D8S264 and D8S201 confirmed that a deletion of distal 8p in the proband was of maternal origin.

Discussion

Rarely are both products of an unbalanced segregation seen within a single pedigree. In our family the translocation has segregated both alternately, giving rise to balanced and presumably normal products, and by adjacent 1 segregation, the latter causing the unbalanced karyotype, which by using the method of Jalbert et al it is predicted to be the mode of segregation giving rise to the least imbalance. The pachytene configuration for the 8;18 translocation is shown in fig 2B. The proband's karyotype is deleted for the segment 8p21 to 8pter and, as expected, she has features associated with the 8p- syndrome, which include significant mental retardation, postnatal growth retardation, microcephaly, and congenital heart defects. More recently Hutchinson et al and Pettenati et al described five and three patients respectively.
and reviewed a further four cases in which they reported a much milder phenotype when the breakpoint is at 8p23.1. However, these latter cases were terminal deletions in which no other chromosomes appeared to be involved, whereas our case may have been modified by the duplication of a portion of 18p. A review of duplication 18p by Wolff et al. found that only 14 cases have been reported previously; nine were detected because of family studies initiated by the diagnosis in another family member of monosomy or tetrasomy 18p, that is, these unbalanced cases had been ascertained indirectly and may otherwise have remained undetected. Four cases had chromosome 18 as the sole rearranged chromosome and were not mentally impaired. However, five of the remaining 10 cases which had another chromosome involved in the imbalance were associated with mild mental retardation and non-specific dysmorphic features. Wolff et al. concluded that duplication 18p does not appear to be associated with significant phenotypic abnormalities, so the phenotype of the proband in our case seems to be caused mainly by monosomy 8p with the congenital heart defect as the only life threatening feature.

In contrast, the other adjacent 1 product seen in the spontaneous abortion (IV-1) gave rise to a chromosome imbalance resulting in partial monosomy 18p and partial trisomy 8p, and as could be predicted, a much more severe phenotype. The 18p-syndrome is a well characterised and relatively common autosomal deletion syndrome in which holoprosencephaly, described in over 10% of reported cases, is probably the most significant congenital malformation. In addition, short stature, mental retardation, and dysmorphic facies are also observed. However, because of developmental delay observed in the fetus (IV-1), it is likely that the above abnormalities would have been more apparent if the fetus had survived for longer. Most of the 17 published cases of duplication 8p11-13 were born at term, unlike our spontaneous abortion (IV-1) which did not survive the first trimester. The majority of these 17 cases were unbalanced owing to mal-segregation of familial translocations, and all were severely mentally retarded, some with brain abnormalities including dilatation of the lateral ventricles or absence of the corpus callosum or both. The fetus in our case had very delayed development of the cranial end of the CNS even by 11 weeks gestation, but because of severe maceration, it is difficult to make further comparisons to those duplication 8p patients that survived life in utero. Therefore, both chromosomal imbalances carried by the spontaneous abortion appeared to contribute towards the severe phenotype, in particular, brain abnormalities, which probably accounted for the fetus’s early demise.

When the proband’s family history is reviewed, it seems that the translocation may have been present in past generations. The proband’s mother lost three sibs, one of which was her twin; and a cousin lost two offspring as spontaneous abortions. Unfortunately no other family members were available for study.