Familial translocations involving 15q11–q13 can give rise to interstitial deletions causing Prader-Willi or Angelman syndrome

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Abstract
A de novo interstitial deletion of 15q11–q13 is the major cause of Prader-Willi syndrome (PWS) and Angelman syndrome (AS). Here we describe two unrelated PWS patients with a typical deletion, whose fathers have a balanced translocation involving the PWS/AS region. Microsatellite data suggest that the deletion is the result of an unequal crossover between the derivative chromosome 15 and the normal chromosome 15. We conclude that familial translocations involving 15q11–q13 can give rise to interstitial deletions causing PWS or AS and that prenatal diagnosis in such families should include fluorescence in situ hybridisation or microsatellite studies or both.

Key words: Prader-Willi syndrome; chromosome 15; translocation.

Prader-Willi syndrome (PWS) and Angelman syndrome (AS) are caused by the loss of function of imprinted genes in 15q11–q13. Approximately 70% of patients have a deletion of 15q11–q13, which in PWS is always of paternal origin and in AS always of maternal origin. Most deletions are de novo interstitial deletions, which probably arise by an unequal crossover between repeated DNA sequences flanking the PWS/AS region. In a few percent of cases, the deletion is the result of an unbalanced translocation. Unbalanced translocations can occur de novo or by segregation of chromosomes involved in a familial balanced translocation. Most PWS patients and a few AS patients without a deletion have two maternal or two paternal chromosomes 15, respectively. Uniparental disomy (UPD) most often results from the postzygotic correction of a meiotic non-disjunction event.

Smeets et al. described a family with a balanced translocation t(6;15)(p25.3;q11.1), in which one cousin had PWS and another had AS. Cyogenetic and molecular studies showed that the PWS patient had a de novo deletion of 15q12 on the paternally derived derivative chromosome 6 and the AS patient had uniparental paternal heterodisomy for chromosome 15. Here we describe two unrelated families in which the father has a balanced translocation involving 15q11–q13. In both cases, a de novo interstitial deletion of 15q11–q13 occurred, which resulted in a child with PWS. The findings reported by Smeets et al. and in this report show that familial translocations involving the PWS/AS region on chromosome 15 can give rise to interstitial deletions and uniparental disomy.

Case reports
FAMILY S (GERMANY)

The proband (III.6) is the second child of young and unrelated parents (age of parents at birth, 29 years) (fig 1). Cytogenetic analysis was performed because of clinical signs of Down syndrome. Chromosome analysis showed trisomy 21 and a balanced translocation (karyotype 47,XY,+21,t(15;18)(q12;q12)). The translocation was also present in the father (II.3) and the paternal grandfather (I.1) (karyotype 46,XY,t(15;18)(q12;q12)). A metaphase of the father is shown in fig 2A. The first son (III.4) is healthy and no cytogenetic analysis was done. A second pregnancy resulted in an abortion (III.5). During the fourth pregnancy, which was uneventful apart from reduced fetal movements, a prenatal diagnosis was performed. Amniotic fluid cells indicated a normal male karyotype (46,XY). Delivery was by caesarean section in the 38th week of pregnancy because of a pathological CTG. At birth, the boy (III.7) had a weight of 2420 g (10th centile), length 49 cm (50th centile), and head circumference 34 cm (50th centile). Pronounced muscular hypotonia, pale skin, high arched palate, abnormal EEG, and cryptorchidism were observed. Tube feeding was necessary for three months.

The father has two healthy sibs. His sister (II.1) has a normal female karyotype (46,XX). She had three abortions, which were not investigated cytogenetically. His brother (II.2) has not been investigated. He has no offspring. Their mother had one abortion in the first trimester. Sibs of the paternal grandfather (I.1) could not be investigated, but there are no signs of developmental retardation or Prader-Willi syndrome.

FAMILY E (THE NETHERLANDS)

Cytogenetic analysis of a mentally retarded boy (III.1) and his normal father (II.2) showed a balanced translocation (46,XY,t(6;15)(q11.2;q13)). The karyotype of the boy is shown...
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Figure 1 Abbreviated pedigrees of families S and E.

in fig 2B. The cause of the mental retardation is unknown. Pregnancy and birth were uneventful. He has no dysmorphic features or focal neurological signs. Metabolic studies were normal. Uniparental disomy 15 or a deletion 15q11–q13 were excluded by DNA analysis. The first pregnancy of a new relationship of this father, a twin pregnancy (III.2, III.3), ended in a miscarriage at 12 weeks’ gestation. In the second pregnancy, amniocentesis was per-

Figure 2 Cytogenetic analysis. (A) Metaphase of II.3 in family S. The normal and derivative chromosomes are indicated. (B) Karyotype of III.1 in family E. (C) FISH analysis of the patient in family S. A probe for D1S11 showed a signal on one chromosome 15 (arrowhead), but not on the other (arrow). The PML probe gave a signal on both homologues. Chromosomes were counterstained with propidium iodide. (D) FISH analysis of the patient in family E. A probe for SNRPN showed a signal on one chromosome 15, but not on the other. The PML probe gave a signal on both homologues.
formed, and normal chromosomes (46,XY) were found. The child (III.4) was born at term with the aid of vacuum extraction because of bradycardia and meconium in the amniotic fluid. The umbilical cord was short (20 cm) and torn off during birth. Apgar scores were 4, 5, and 7 after one, five, and 10 minutes, respectively. He showed severe hypotonia and respiratory insufficiency, and needed artificial ventilation. There was no pulmonary explanation for his respiratory distress, which did not improve in the following days. He also showed no sucking reflex and needed gavage feeding. His birth weight was 3265 g (40th centile) and his head circumference was 35.5 cm (50th centile). He had a small face with a narrow bitemporal diameter, hypertelorism, micrognathia, cryptorchidism, scrotal hypoplasia, contractures of the extremities, and simian creases of both hands. EEG, EMG, and MRI of the cerebrum showed no abnormalities. Because of an extremely poor prognosis, the parents did not wish to continue treatment. Necropsy showed no abnormalities except for cryptorchidism.

The father’s sister (II.6), who had one intraterine death (III.6), two miscarriages (III.8, III.9), and two healthy daughters (III.5, III.7) carries the same translocation.

### Results and discussion

Subject III.7 in family S (patient S) and subject III.4 in family E (patient E) have typical Prader-Willi syndrome according to the consensus diagnostic criteria. In both families, the father has a balanced translocation involving 15q11–q13. Prenatal cytogenetic analysis of the patients had shown a normal male karyotype (46,XY) at the 50th band stage. After birth, metaphase chromosomes from peripheral blood lymphocytes were studied by fluorescence in situ hybridisation (FISH). Using a cosmid probe for D15S11, metaphase spreads from patient S were found to have a signal on one chromosome 15 only (fig 2C). A cryptic translocation between chromosomes 15 and 18 was excluded by hybridisation with library probes for these two chromosomes (not shown). Patient E was studied by FISH with a cosmid probe for SNRPN, which showed a deletion (fig 2D).

To determine the extent and origin of the deletions, we used the D15S63 methylation test and performed microsatellite analysis with five markers mapping within the typical PWS/AS deletion region (D15S541, D15S543, D15S11, D15S128, and GABRB3) and two markers mapping to 15q14–q21 (D15S144 and CYP19). The results are summarised in table 1. Both patients lack paternal alleles from D15S541 to D15S128. Patient E also lacks a paternal allele at GABRB3. Patient S is hemizygous or homozygous at this locus. These data suggest that the patients have a typical deletion. As the fathers are heterozygous at these loci, the deletions have occurred de novo.

Although we cannot distinguish between hemizygosity and homozygosity for D15S144 and CYP19, the cytogenetic data exclude an involvement of these loci in the deletions. Hence, D15S144 is the most proximal intact
and informative marker in family E. In family S, this is CYP19 because D15S144 was not informative. The PWS patients and their translocation carrying brothers have different paternal alleles at these loci. Assuming that there is no recombination between the translocation breakpoint and D15S144 in the brother of patient E (III.1), allele 3 is on the derivative chromosome 8 and allele 2 is on the normal paternal chromosome 15. In family S, allele 1 of CYP19 is on the derivative chromosome 18 and allele 2 is on the normal paternal chromosome, again assuming that there is no recombination between the translocation breakpoint and this marker in the brother of patient S (III.6). In both families, the patients inherited the alleles located on their father’s normal chromosomes. This suggests that the deletions are the result of unequal crossovers between the derivative chromosomes 15 and the normal homologues (mechanism B in fig 3). It should be noted, however, that we do not know the linkage phase in these families. Therefore, mechanism A (fig 3) cannot be formally excluded.

In addition to our families and the family reported by Smeets et al, we are aware of at least one other family in which a 15q11–q13 deletion occurred in a family segregating a familial translocation involving chromosome 15 (t(Y;15)(q12;p11)), although the translocation breakpoint in this family is outside the PWS/AS region (S Eliez, C DeLozier, personal communication). In the families identified by Smeets et al and Eliez and DeLozier, the deletion is present on a translocation chromosome. In our families, the deletion is present on an otherwise normal chromosome and was overlooked by conventional cytogenetic analysis. These findings show that interstitial deletions can occur de novo in families with a balanced translocation involving 15q11–q13. However, the number of families is too small to prove that translocations increase the frequency of unequal crossovers. Similarly, we do not know whether translocations involving other chromosomal regions will give rise to deletions at the same frequency as those involving 15q11–q13, or whether our findings are specific for chromosome 15. The presence of DNA repeats flanking the PWS/AS region might support the latter view. However, there are several examples of families in which one translocation carrier is healthy and another one is affected. The disease may be related to a de novo submicroscopic deletion or the presence of uniparental disomy.

Our findings have important implications for prenatal diagnosis. It is common practice to exclude risk, when a fetus is found to have the same balanced translocation as his parents or to have a normal karyotype. At least for translocations involving 15q11–q13, this can be wrong. We conclude that prenatal diagnosis in such cases should include FISH and microsatellite analysis to rule out a deletion or uniparental disomy. Depending on the parental origin, such a deletion leads to Prader-Willi or Angelman syndromes.

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