A boy with developmental delay and a maternally inherited deletion in 15q11q13

M King, C Hardy, B Asenbauer, M Kilpatrick, T Webb

Abstract
A boy was referred at 8 weeks of age for failure to thrive. Cytogenetic and molecular studies showed that he had a large proximal deletion of the maternally derived chromosome 15q. He did not have Angelman syndrome, but at 2 years of age was severely globally delayed. He died at 2½ years of age.

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Key words: developmental delay; chromosome 15; deletion.

Cytogenetic deletions involving 15q11q13 are associated with both Prader-Willi syndrome (PWS)1 2 and Angelman syndrome (AS).3 The origin of the deleted chromosome is paternal in PWS and maternal in AS, which is characterised by severe developmental delay, an ataxic gait, hand flapping, inappropriate laughter, tongue thrusting, and seizures.4

Molecular studies have confirmed the parental origin of the deleted chromosome 15 in both syndromes and have also shown frequent cases of maternal uniparental disomy in non-deletion PWS5 6 and occasional paternal uniparental disomy in AS,7 confirming that the region is imprinted. Molecular investigation has also determined that the Prader-Willi critical region (PWS/AS) is located proximal to that for AS (ASCR), but that the majority of cases of either syndrome have common breakpoints lying between D15S18 and D15S9 on the proximal side and between D15S12 and D15S24 on the distal (the PWS/ASCR).7 The gene for oculocutaneous albinism (D15S12), unlike several others in the region, is not imprinted,8 and its inclusion within the common deletion may account for the hypopigmentation which is observed in both PWS and AS.7 This locus in 15q13 marks the proximal breakpoint of the common deletion.7

Proband who have proximal 15q deletions but who do not have a typical PWS or AS phenotype have occasionally been reported. Generally, if the deleted region is large and reaches into 15q14 or beyond then a more severe phenotype may result.5

Case report
The proband was the third child of healthy, unrelated parents who were both in their third decade at the time of his birth. He was born at term by spontaneous vertex delivery, birth weight 3500g, length 53 cm, head circumference 37 cm. There were no neonatal problems and he breast fed well. At 4 weeks of age his feeding deteriorated and he was admitted to hospital at 8 weeks of age because of failure to thrive.

On examination, his length and head circumference were on the 50th centile, while weight had dropped to the 3rd centile. He did not have any specific dysmorphic features (fig 1). He had marked dyskinesia with dystonic posturing, choreoathetoid movements of the limbs, tongue thrusting, and jaw tremor on sucking. Limb tone and tendon reflexes were increased. There was marked truncal hypotonia. He did not fixate or track or smile. There was horizontal jerky nystagmus and pale optic discs. The primitive reflexes were exaggerated.

INVESTIGATIONS
A normal karyotype was reported from the referring hospital and so initially a metabolic or structural brain disorder was suspected.

Biochemistry
In blood the following investigations were normal: acid base balance, urea, electrolytes, creatinine, creatine kinase, ammonia, lactate/pyruvate ratio, uric acid, cholesterol, triglycerides, B12, folate, thyroid function studies, biotinidase, platelet DHAPAT, vitamin E, very long chain fatty acids, bile acids and lysosomal enzymes.
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cerebral atrophy. There was a symmetrical area of abnormal myelination involving the deep white matter around the posterior horns of the lateral ventricles.

**Electrophysiology**

Visual evoked responses were absent while electroretinogram was of small amplitude. Electroencephalogram (EEG) between the ages of 9 weeks and 2 years showed diffuse high voltage slow activity (400 μV) with associated spike and slow wave discharge during sleep. The typical EEG features of AS were not seen.

**CYTOGENETIC STUDIES**

Prometaphase chromosomes were obtained from blood lymphocytes both by synchronisation and by intercalation of DNA with ethidium bromide. The proband had the karyotype 46,XY,del(15)(q11q13)(mat). The deletion was observed to be larger than that generally found in either Prader-Willi or Angelman syndromes (fig 2). Study of polymorphisms associated with the short arm of chromosome 15 showed the deleted chromosome to be maternal in origin. No other additional karyotypic abnormality was observed.

![Figure 2](https://example.com/figure2.png)

*Figure 2* Two chromosomes 15 from the proband. The abnormal, deleted homologue is on the left hand side of the pair. Band 15q12 is clearly visible in the normal homologue (arrowed) but is deleted in the abnormal chromosome 15.

In urine the following were normal: sulphite test, amino acids, bile acids, and organic acids.

In cerebrospinal fluid the following were normal: protein, amino acids, and lactate.

**Imaging**

Abdominal and cerebral ultrasound, skeletal survey, and CT brain scan were all normal. However, MRI of the brain showed generalised cerebral atrophy. There was a symmetrical area of abnormal myelination involving the deep white matter around the posterior horns of the lateral ventricles.

![Figure 3](https://example.com/figure3.png)

*Figure 3* Southern blot of RFLP analysis at D15S11 showing lack of maternal inheritance in the proband (left). Autoradiographs showing maternally derived deletions in the proband at three loci in 15q11q13 (GABRA5, GABRB3, D15S24) (centre) and the presence of both maternal and paternal contributions at loci ACTC and D15S86 (pMS620) (right).
Informative haplotypes at loci within the 15q11q13 region for the proband and his parents

<table>
<thead>
<tr>
<th>Locus</th>
<th>Proband</th>
<th>Mother</th>
<th>Father</th>
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<tr>
<td>D15S11</td>
<td>1-2kb</td>
<td>1-0 kb</td>
<td>1-2, 1-2kb</td>
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<tr>
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<td>3-</td>
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<td>1.3</td>
</tr>
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<td>D15S113</td>
<td>2-</td>
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</tr>
<tr>
<td>GABRB3</td>
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<td>1.2</td>
<td>2.3</td>
</tr>
<tr>
<td>GABRA5</td>
<td>1-</td>
<td>3.3</td>
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</tr>
<tr>
<td>D15S24</td>
<td>B-</td>
<td>A, A</td>
<td>B, C</td>
</tr>
</tbody>
</table>

MOLECULAR STUDIES

The boy and his parents were typed with polymorphic markers for chromosome 15 loci to define the deletion at the molecular level (table). As his father was homozygous for the 1.2 kb band and his mother homozygous for the 1.0 kb band, while the proband showed only the 1.2 kb band, RFLP analysis using RsaI for locus D15S11, which lies within the Prader-Willi chromosomal region (PWCR), showed no maternally derived material in the proband (fig 3, left), as did analysis with microsatellite markers for D15S11, D15S128, and D15S113 loci, and the GABRB3 and GABRA5 genes (fig 3, centre), which have also been assigned to the Angelman/Prader-Willi region. Analysis of the D15S24 locus, which lies distal to the PWCR, again showed no maternally derived material in the proband (fig 3, centre). Analysis with a CA repeat marker for the cardiac muscle actin gene locus (ACTC), which lies distal to D15S24, however, did show the presence of a maternally derived allele (fig 3, right). This defined the distal boundary of the deletion as between D15S24 and ACTC. The family was also fully informative for the telomeric marker pMS620 (D15S86) which showed the presence of both maternally and paternally derived alleles in the proband (fig 3, right). Fig 4 shows an ideogram of chromosome 15 with deleted informative loci in 15q11q13 shown in order. It can be seen that the deletion includes all of the PWS/AS critical region and extends distally beyond it.

PROGRESS

There was gradual improvement in feeding but no neurodevelopment. At 2½ years he was severely globally delayed. He had no head righting or head control and was not sitting unsupported, reaching, smiling, or communicating. He did not make contact with people or display any interest in his surroundings. The dyskinesia remained prominent. He had developed mild epilepsy which was controlled by sodium valproate.

At 2½ years he was found dead in his cot. General necropsy did not show any abnormality. Neuropathological examination showed features of mild hypoxic ischaemic encephalopathy of recent onset. There was also mild non-obstructive ventricular dilatation. In particular there was normal cortex, subcortical white matter, striatum, brain stem, and cerebellum with no evidence of demyelination or dysmyelination. It is likely that death occurred as a result of prolonged apnoea during a seizure.

Discussion

A boy with gross developmental delay but without Angelman syndrome has been found to have a maternally derived deletion of chromosome 15 at 15q11q13. The deletion was microscopically very large and was shown by molecular methods to include D15S24, a locus which is known to lie distal to the PWS/ASCR and is very rarely deleted in either Angelman or Prader-Willi syndromes. The deletion in this proband was not found to extend into 15q21 either cytogenetically or by the finding of heterozygosity at the locus for ACTC. The distal limit of the deletion was thus defined as lying between ACTC and CMW-1 (D13S24). Proximally it included D15S11, shown both by Southern blotting and by PCR methodology. Paternal uniparental isodisomy was excluded both cytogenetically and by the inheritance of a maternal contribution in D15S86 and ACTC. A combination of cytogenetic and molecular estimations of the size of this deletion suggests that it may represent at least 10% of the total length of chromosome 15 or approximately 10 megabases and extend beyond the imprinted region associated with Prader-Willi and Angelman syndromes. The severity of the clinical symptoms culminating in the death of the proband is most probably consequent upon the size of the deletion.

This would, however, imply that haploid inefficacy of an important gene located between D15S12 and ACTC could have such dire consequences or that there are further imprinted genes lying beyond D15S12.

The combination of clinical signs in this case has not, to our knowledge, been reported in association with deletion of maternally derived chromosome 15q11q13. In particular, the lack
of any neurodevelopment whatsoever, early death, striking dyskinesia, and eye findings were not typical of Angelman syndrome. In addition, the characteristic craniofacial, neurological, and EEG features of this syndrome were not present. Although a maternally derived deletion in 15q11q13 which encompasses the critical region would be expected to result in a phenotype with manifestations of Angelman syndrome, the influence of an increased chromatin loss is concomitant with a more severe phenotype and the early death of this boy, so that direct developmental comparisons were precluded.

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