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An infant with ring 17 chromosome and unusual dermatoglyphs: a new syndrome?

SUMMARY A case of ring 17 chromosome in a 5-month-old male infant is investigated and compared with five previously reported cases. The findings commonly observed in these patients include mental and motor retardation, seizures, short stature, muscular hypotonia, and microcephaly among others. Dermatoglyphic studies showed an increased number of ulnar loops. More interestingly, bilateral transverse hypothenar creases were noted. Two of the reported cases also had unspecified genital abnormalities. The variation in clinical findings among these patients may be explained by a difference in the breakpoints on chromosome 17.

To the best of our knowledge this is the third report of a liveborn child with ring 17 chromosome without mosaicism confirmed by banding. We present here the clinical findings in our patient compared to those observed in other cases with and without mosaicism as described in published reports.

Case report

The proband (fig 1) was the first child of non-consanguineous parents born after an uncomplicated 42 week pregnancy. The birthweight was 1290 g. The mother was 32 and the father was 34 years old.

The patient was seen at 5 months of age with a weight of 4432 g, head circumference 36·6 cm, and length 53·3 cm (all below the 3rd centile). Inner canthal distance was 30 mm and outer canthal distance was 76 mm (below the 97th centile). Anterior fontanelle size was 5·5 cm (greater than the 97th centile). The forehead was flattened and he had unruly hair. The nasal bridge was broad and depressed with anteverted nostrils. The palpebral fissures had a downward slant with epicanthal folds. He had a disconjugate gaze and an alternating esotropia. The ears were low set, with the left ear being larger than the right. Flattening of the helix,

FIG 1 Proband at 5 months of age.
antihelix, and tragus was noted on the left. Both upper and lower lips appeared thin. Testes were palpable in the scrotum and the urinary meatus was normally placed. There was fifth finger clinodactyly bilaterally.

Dermatoglyphic analysis showed ten ulnar loops on the finger tips and r triradii located in the hypothenar areas of both palms. There was considerable ridge dotting and some dissociation. Transverse hypothenar creases were also noted bilaterally (fig 2).

At 2½ months of age, the patient was admitted to hospital with gastroenteritis. Five days after admission he had two grand mal seizures. The electroencephalogram was normal, but computerised tomography of the brain showed hydrocephalus involving the anterior portion, occipital, and temporal horns of both lateral ventricles. He was started on phenobarbital and has had no subsequent seizures. Developmental testing at 4 months of age showed a Developmental Quotient of 40 with gross and fine motor ceilings of 6 weeks.

**FIG 2** Dermatoglyphic patterns of the proband.

**FIG 3** E group chromosomes of the proband from fibroblast (A) and peripheral lymphocyte (B and C) cultures.

**CYTOGENETIC STUDIES**

Chromosomal analysis of 30 peripheral lymphocyte and 50 skin fibroblast metaphases by G banding, using a modification of Seabright's technique, showed a 46,XY,r(17) karyotype with breakpoints located at p13 and q25 (fig 3). The karyotypes of the parents were normal, except that the mother had an enlarged satellite on one chromosome 14, which was not seen in the child.

**Discussion**

This is the sixth report of a patient with ring 17 chromosome and the third such case without mosaicism (to be confirmed by banding studies) in a liveborn infant. The patient had mental retardation, short stature, low weight, hypotonia, microcephaly, seizures, hydrocephalus, and abnormal dermatoglyphs.

Petit and Koulisher reported a child with the mosaic karyotype 46,XX/46,XX,r(17), confirmed by autoradiography, who had mental retardation, short stature, and low birthweight. A father and son were described by Burdea et al to have the karyotype 46,XY,r(17), which was not, however, confirmed by banding. The father was reported to have moderate microcephaly, while the son had mental retardation and minor phenotypic abnormalities. Ono et al described a 10-year-old girl with a 46,XX,r(17)(p13;q25) karyotype who had short stature, microcephaly, mental retardation, hypotonia, and seizures. At the age of 12, her seizures became uncontrollable with subsequent mental deterioration (Nakagome, 1980, personal communication). In the fourth report by Weinberg et al banding studies indicated a 46,XX,r(17) karyotype in amniotic fluid cells. A necropsy after a therapeutic abortion at 20 weeks' gestation showed no abnormalities either grossly or microscopically. However, it was recorded that subtle abnormalities of facial development might not be apparent at this stage.

Qazi et al reported a mentally retarded 6-year-old boy with speech delay, seizures, abnormal facial features, and abnormal dermatoglyphs. Another case of a mosaic 46,XX/46,XX,r(17) is listed in Borgaonkar et al but no clinical information is available.

A comparison of our findings with those reported in previous cases of ring 17 chromosome, both with and without mosaicism, are shown in the table. On the basis of the common findings in the present case and previously reported cases, with the exception of the father in Burdea et al, a pattern emerges suggesting a recognisable syndrome associated with this chromosome abnormality. The non-specific findings present in the majority of the cases were:

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*Case reports*
mental and motor retardation, growth retardation, microcephaly, flattened nasal bridge and anteverted nostrils, epicanthus, clinodactyly, and hypotonia. Seizures and café-au-lait spots, not commonly seen in other chromosomal syndromes, were present in the majority of patients. Most interestingly, the dermatoglyphic findings show the presence of a transverse hypothenar crease in the present case. This crease has previously been described only in Coffin-Lowry syndrome (Plato, 1980), personal communication. Only two illustrations of dermatoglyphic patterns of the hands were available in the reported cases and, of these, one case reported by Qazi et al.6 also had transverse hypothenar creases.

With the limitations of the present banding techniques, it is not always easy to determine the breakpoints which result in a ring chromosome. Certainly a difference in the breakpoints, and thus the area involved in the terminal deletions, could account for differences in phenotypes. However, the size of the ring chromosomes and the location of the breakpoints appear to be quite similar in our case and those of Weinberg et al.,6 Ono et al.,4 and Qazi et al.,6

More accurately, one way to determine the extent of the deletion of this chromosome would be through the use of biochemical markers. Karyotype-phenotype correlations will be difficult to achieve until more individual gene products can be studied.

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