Karyotype 46,XX in a Black Male

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SUMMARY

A clinical and cytogenetic report is made of a Black male child with the karyotype 46,XX. Although 46,XX males have already been reported, this is the first time this anomaly has presented in a Black.


In the late 1950s it was accepted that the normal male phenotype is associated with a 46,XY karyotype. Chapelle et al. reported a male with a 46,XX karyotype and since then a number of cases have been recorded. This report presents a case of a Black male child with the karyotype 46,XX.

CASE REPORT

The patient, a 13-month-old Black male of the Lomwe tribe from Malawi, was referred for investigation of congenital malformation of the genital organs. A hypospadic penis was present with the urethral opening at the base of the phallus. Partly undescended testes were palpable in the inguinal canal and could be brought down into the scrotum by gentle traction (Fig. 1).

No uterus or ovaries were seen at laparotomy. A small biopsy specimen was taken from each gonad and showed normal immature testicular tissue in each case (Fig. 2). The child appeared alert and unretarded.

Cytogenetic Studies

Chromosome analysis was performed on phytohaemagglutin-stimulated lymphocyte cultures obtained from peripheral blood. Suitable metaphase spreads were stained with buffered Giemsa, and for fluorescent studies by the method of Caspersson et al. A total of 50 cells were karyotyped from the first culture, and the same number from a repeat culture. A modal number of 46,XX chromosomes was found in each case.

Buccal smears were stained for Barr bodies by the method of Klinger and Ludwig, and for Y-body fluorescence by the method of Kim et al. Single Barr bodies were seen in 35% of cells examined, but Y-body fluorescence was not present. The patient was removed by his father before tissue for skin fibroblast cultures was obtained.

DISCUSSION

A number of cases have been reported suggesting that the Y chromosome is unnecessary for the development of testicular tissue. In the majority of reports of 46,XX males some clinical features of Klinefelter's syndrome were noted.

The hypospadic penis reported here appears to favour a diagnosis of true hermaphroditism, and was not found in other recorded cases of 46,XX males. In these cases a normal, although small, penis was usually present.
Shah et al., recorded a male pseudohermaphrodite, karyotype 46,XX, with ambiguous external genitalia, an enlarged phallus and a vagina with a separate opening from that of the urethra. No uterus or ovarian tissue was noted. The present case lacked the rudimentary vagina which is commonly found in pseudohermaphrodites. In addition, the absence of ovarian tissue discounts true hermaphroditism.

The possibility that the male determining loci may be on the short arm of the Y chromosome, leaves room for speculation that a translocation of a deleted and hence non-fluorescent Y chromosome could account for this phenomenon. Kasdan et al. showed evidence of an autosomal inheritance of a male-determining gene in a family with several male members having a 46,XX karyotype.

I wish to thank Dr E. Forbes for referring the patient.

Books Received: Boeke Ontvang


REFERENCES


