Unusual dual genital duct remnants in true hermaphroditism

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SUMMARY A case of true hermaphroditism is reported in which a 46,XY karyotype was associated with a testis and an ovotestis. The dual presence of a Fallopian tube and a vas deferens on the side of the ovotestis is documented as a previously unreported finding.

In true hermaphroditism both ovarian and testicular tissue are present in the same subject. The nature of the genital ducts adjacent to the gonads is of particular interest; it is generally accepted to be related to the ability of testicular tissue to stimulate the development of the Wolffian ducts and induce regression of the Müllerian ducts. A case of true hermaphroditism in which both vas deferens and Fallopian tube were present on the same side as an ovotestis is described.

Case report

The proband was the first child of healthy, unrelated parents. The pregnancy was uneventful and spontaneous delivery occurred at term. Birth weight was 3.05 kg. There was no relevant family history.

Ambiguous genitalia were noted at birth. There was an enlarged clitoris surrounded by fused rugose labial folds. No gonads were palpable. There was a single perineal urethral opening. The remainder of the examination was normal.

The peripheral karyotype determined on two separate occasions was 46,XY with no evidence of mosaicism. A total of 78 mitoses was counted and no abnormality was detected using G, C, and Q banding techniques. The plasma 170H progesterone concentration was normal (8 nmol/l) thus excluding 21-hydroxylase deficiency. Plasma testosterone concentrations increased from 3.1 to 6.7 nmol/l after three daily injections of human chorionic gonadotrophin (HCG 2000 IU/day). A perineal sinogram showed a left cornuate uterus and Fallopian tube. An intravenous pyelogram was normal. Laparotomy performed at the age of 11 months by Mr J Lari showed a normal sized uterus for age to which a left Fallopian tube was attached. Both gonads were found in the ovarian position. The gonads and adnexal ducts were excised and the uterus left in place. A reduction clitoroplasty was also performed.

The left gonad was an irregular white nodule measuring 1.5x1.0x0.3 cm to which a 1 cm length

FIG 1 Ovotestis. Ovarian stroma containing a number of primordial follicles.
of cylindrical tissue 0·3 cm in diameter was attached. Histologically the bulk of the gonad consisted of ovarian tissue containing primordial follicles (fig 1). A small amount of testicular tissue in the form of tubules lined by Sertoli cells was present at one margin (fig 2). The line of demarcation between ovary and testicular tubules was abrupt. The tissues around the gonad showed both normal Fallopian tube and normal epididymis and vas deferens (fig 3). The ductal structures were separated by a short amount of loose connective tissue. The more distal regions of the adnexal duct showed only a Fallopian tube.

The right gonad had the appearance of a small immature testis. The seminiferous tubules were lined by immature Sertoli cells and contained small numbers of germ cells. The associated ducts were an epididymis and vas. Karyotype analysis of the testis based on five mitotic counts was 46,XY.

Discussion

The classification of true hermaphroditism is based upon the lateral distribution of gonadal tissue. While an ovotestis is the most common gonad in true hermaphroditism, the combination of an

FIG 2 Ovotestis. Seminiferous tubules lined by Sertoli cells.

FIG 3 Gonadal ducts. Dual presence of a Fallopian tube (left) and epididymis and vas deferens (right).
ovotestis with a testis is rare and occurs in only 10% of cases. A 46,XY karyotype is also uncommon, present in 11-6% of all cases. This case fulfils the criteria of true hermaphroditism because of the presence of a testis on the right and an ovotestis containing primordial follicles on the left side. The condition must be differentiated from mixed gonadal dysgenesis. This is characterised by a unilateral testis, a streak gonad (lacking follicles or testicular tubules) on the contralateral side, and persistent Müllerian structures, frequently in the form of a uterus and bilateral Fallopian tubes. The karyotype usually shows XO/XY mosaicism.

The nature of the genital duct associated with an ovotestis is a matter of considerable interest. In the series of 48 patients reviewed by van Nierkerk, either a Fallopian tube was present (in 65%) or there was a vas deferens (in 35% of cases). The normal development of the internal genital ducts is determined by the presence or absence of adjacent testicular tissue. Müllerian duct regression in males is mediated between day 62 and 77 of human gestation by anti-Müllerian hormone (AMH), a protein derived from fetal Sertoli cells. There is a period of maximum sensitivity to AMH during this early phase of gestation. The process precedes Wolffian duct stabilisation which is dependent upon the production of a high local concentration of testosterone between day 71 and 74 of gestation.

The persistence of Müllerian ducts in otherwise normal males is well recognised, occurring either as a sporadic or familial disorder. Retention of both Müllerian and Wolffian ducts adjacent to an ovotestis has not been described previously. Inappropriate timing of adequate AMH release coupled with insufficient Leydig cell synthesis of testosterone is one possible reason for this unique observation, but the hypothesis is impossible to prove postnatally. Perhaps the indifferent gonad failed to grow to a critical size before differentiation into a testis. It has been proposed by Cunha et al that complex hormonal effects involving epithelial morphogenesis result from cell-cell communication between mesenchyme and epithelium. An alternative hypothesis is that cells (perhaps mesenchymal) responsible for Müllerian duct regression did not establish appropriate mesenchymal/epithelium relationships at the critical time.

The gonads and internal genital ducts in all cases of true hermaphroditism should be examined carefully to document whether the coexistence of Müllerian and Wolffian duct remnants adjacent to the same gonad occurs more often than previously recognised.

We thank Mr S Roberts and Miss E Little for the cytogenetic studies and Dr R Prosser for referring the patient.

References


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doi: 10.1136/jmg.25.3.206

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