Gonadal mosaicism 45,X/46,X,psudY(q11.2) resulting in a Turner phenotype with mixed gonadal dysgenesis

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ABSTRACT
A two-year-and-eight-month-old girl was referred with clitoromegaly and short stature. Two cell lines, 45,X and 46,X,psudY(q11.2), were observed. Cytogenetic and fluorescence in situ hybridisation investigations were carried out on her peripheral lymphocytes and gonadal cells, to determine the genotype-phenotype effect with respect to differential tissue distribution, effects of the sex determining region of the Y chromosome, and the break-points in the azoospermia factor region.

Keywords: fluorescence in situ hybridisation, gonadal dysgenesis, mosaicism, sex chromosome aberrations, Turner syndrome

INTRODUCTION
As dicentric chromosomes are inherently unstable, isodicentric Y chromosomes are usually found in association with a monosomic X (45,X) cell line. Consequently, clinical presentations in these patients can range from a classical Turner phenotype, through mixed gonadal dysgenesis, to phenotypically-normal males.1-2 Such variability in the sexual phenotype is thought to be related to the tissue distribution and relative proportions in the developing gonads of the respective cell lines, particularly those with functional copies of the SRY gene (sex determining region of the Y chromosome). However, studies on gonadal tissue are hindered by the fact that it is rarely available for analysis, and a more easily-accessible tissue is usually studied. A further factor influencing the phenotype may be the breakpoint in the Yq long arm, resulting in the duplication or absence of the regions bearing genes for azoospermia factor (AZF) a, b and c. We have karyotyped gonadal skin, right and left gonads as well as peripheral lymphocytes to determine the distribution of these two cell lines. Fluorescence in situ hybridisation (FISH) probes have been used to detect the SRY, as well as to localise the breakpoint in the Yq11.2 region to determine the presence/absence, as well as the effect of the AZF region on phenotype.

CASE REPORT
A two-year-and-eight-month-old girl was referred with clitoromegaly and short stature (height 81 cm: < third percentile). She was the fifth child of a non-consanguineous marriage with uneventful birth history, and her developmental milestones were normal. Examination of the external genitalia revealed clitoromegaly (phallic length 3 cm; diameter 1 cm), with separate urethral and vaginal orifices (Fig. 1). Pelvic ultrasonography demonstrated a vestigial uterus with a patent vagina, no ovaries and no adnexal mass. Laboratory findings included normal levels of 17-hydroxy progesterone (0.34 nmol/L), thus excluding simple virilising congenital hyperplasia. The serum testosterone was unmeasurable at < 0.14 nmol/L. Laparoscopic exploration revealed a left streak ovary and a right ovotestis. She underwent clitoroplasty and a bilateral gonadectomy. Histopathology showed the left gonad to contain streak ovarian elements, while the right gonad consisted mainly of immature testicular tissue with some rudimentary ovarian tissue in the adjacent connective tissue.

Chromosomal analysis of peripheral blood lymphocytes in our index patient revealed two populations of cells. One cell line (10%) was hypodiploid with only one X chromosome, while the other cell line (90%) was diploid, having one X chromosome and an isodicentric Y with a breakpoint in Yq11.2 (Fig. 2). Fibroblasts from the genital skin were cultured and two cell lines were observed in the percentage of 45,X[72]/46,X,psudY(q11.2)[28]. Gonadal tissue taken from the right gonad showed 45,X[69]/46,X,psudY(q11.2)[31] and the left gonad showed only the 45,X cell line (Fig. 3). For FISH analysis, probes on both the long and short arms of the Y chromosome were used (Wessex Regional Genetics Reference Laboratories, Salisbury, UK and Sanger Centre, Cambridge, UK). The proband’s abnormal Y chromosome showed two FISH signals with the Ypter, SRY(Yp11.2), DYZ3(Ycen), RP11-70G12 (Yq11.23), RP11-539D10 (Yq11.2), while for the aberrant cell line only the SRY probe showed two signals with the Ypter, DYZ3(Ycen), RP11-70G12 (Yq11.23), and RP11-539D10 (Yq11.2). Consequently, the SRY gene and the Yq11.2 breakpoint were present in only one cell line, which was hypodiploid. This was thought to be a consequence of the abnormal Y chromosome with an isodicentric Y, resulting in a Turner syndrome with a common deletion of the AZF region.
(Yq11.23) probes (Fig. 3), but no signal was seen with the probe CTA959A10 (Yq11.23) and DYZ1(Yqh) probe indicating that the breakpoint lies in the 800 kb interval between RP11-539D10 and CTA959A10 in the AZFc region (Figs. 4a & b).

**DISCUSSION**

Isodicentric Y chromosomes appear to be formed by a single break in one of the Y chromatids, followed by a fusion of the broken ends of sister chromatids and the loss of the acentric fragment during gametogenesis before formation of spermatids. Such rearrangements are generally unstable and an additional 45,X cell line is frequently present. This study aimed to establish the influence of sex chromosome mosaicism on the resulting phenotype with respect to differential tissue distribution, effects of the SRY and the breakpoints in the AZF region. The ratio of the Y-bearing cells to the monosomic X cell line in blood and gonadal tissues was highly skewed and not at all representative of each other. The gonadal skin had 28% idicY-bearing cells, the rest being monosomic for the X chromosome. The right gonad, which was an ovotestis, had all cells with 45,X and no idicY cells. The peripheral blood lymphocytes showed 90% of cells with an idicY chromosome. The differing representation is understandable as gonads develop from the endoderm, while blood is derived from the mesoderm. As gonadal tissues are not easily available, there are few reports on such comparisons, and most studies correlate phenotype with peripheral lymphocyte karyotypes, which leads to the current uncertainty with predicting the phenotype resulting from this mosaic karyotype.

Formation of the testis from the undifferentiated embryonic gonad depends on the presence of the short arm of the Y chromosome, containing SRY-sequences. Testosterone production stimulates development of the Wolffian system and induces male development of the external genitalia, failing which, differentiation proceeds along female lines and Müllerian structures are formed. There seems to be the necessity of a minimal amount of SRY to be present for the undifferentiated gonad to become a testis. Apparently, in our case, the threshold of SRY-containing cells (31%) required for the development of the embryonic gonad into a testis was obviously not adequate to enable complete differentiation of the right gonad into a testis resulting in clitoromegaly. A report by Reddy et al, showed the presence of a testis with only 21% Y-bearing cells. Another illustrative case, reported by Sugarman et al, described a male individual with mixed gonadal dysgenesis and a 45,X/46,X,dic(Yp) karyotype. These authors had the opportunity to karyotype cells of the streak gonad on one side, and the normal looking testis on the other side. A 45,X karyotype was found in metaphases of tissue cells derived from the streak gonad, while a 46,X,dic(Yp) karyotype was found in tissue cells from the normal testis on the other side, which obviously had adequate SRY products to differentiate the gonad into...
a testis. An attempt to refine the breakpoint in the Y long arm using FISH probes was successfully carried out in this patient. The fluorescently-labelled probes RP11-70G12 and RP11-539D10, used in this study, lie within the two clusters of DAZ (deleted in azoospermia) genes, absence of which can cause infertility in males.\(^{(8,9)}\) In our patient, the breakpoint lies between 539D10 and CTA959A10, which is beyond the DAZ gene cluster, indicating presence of the DAZ gene (Fig. 5). However, this would not have any bearing on the proband as she had streak gonads, which were later removed.

In conclusion, the cells predominantly present in the gonads seem to be influencing the phenotypic sex, irrespective of the percentage of Y-bearing cells in the blood or buccal mucosa. The 45,X cell line may exert a more dominant effect on sexual differentiation, only if it is predominant in the gonads. This seems to be supported by a recent article on prenatally diagnosed 45,X/46,X,idic(Y)\(^{(10,11)}\) where the outcome of the cases was a normal male phenotype, despite differing percentages of the X- and Y-bearing cell lines in the peripheral blood. The presence of testes in these individuals obviously suggests that the Y cell line with adequate SRY product must have been present in the gonadal tissue—enabling differentiation along male lines, though infertility may be a distinct possibility. It is, however, clear that the percentages of the Y-bearing cell line in the peripheral blood remains an unreliable indicator of the sexual phenotype. This is exemplified in our proband where the presence of only the 45,X cell line leads to the formation of ovarian tissue in the left streak gonad, whereas the mosaic presence of the idic(Y) cell line together with the 45,X cell line, leads to formation of testicular tissue in the right streak ovoidistis. In our patient, the 90% Y cells in the peripheral blood are not a suitable indicator for the resulting phenotype.

REFERENCES