the severe ptosis, however, has made only a slight difference in the child's visual responsiveness.

Comparison of the two cases indicates that monosomy 3p forms a clinically recognisable pattern of malformation which should be considered in the face of prenatal onset growth deficiency, small ears, ptosis, and postaxial polydactyly.

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Addendum
Since this article was submitted for publication, we have become aware of two further cases with identical deletions of chromosome 3. The children, both females, aged 11 and 3 years, also had severe growth and mental deficiency, microcephaly, ptosis, micrognathia, and altered ear structure. The findings further support the hypothesis that monosomy 3 (p25) has a distinct clinical phenotype.

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**Chronic renal disease, myotonic dystrophy, and gonadoblastoma in XY gonadal dysgenesis**

**SUMMARY** A patient with XY gonadal dysgenesis and gonadoblastoma showed myotonic dystrophy and chronic renal disease of unknown aetiology. The coexistence of renal disease and XY gonadal dysgenesis in this and two other subjects suggests a presently obscure aetiological relationship between the phenomena.

XY gonadal dysgenesis is a genetically heterogeneous group of disorders, the most common form being an X linked recessive or male limited autosomal dominant disorder in which phenotypic females have bilateral streak gonads and fail to undergo normal secondary sexual development. The pathogenesis is uncertain and H-Y antigen may or may not be present. Gonadoblastomas or dysgerminomas have been reported in 20 to 30% of cases.

Blanchet et al reported a patient who, in addition to XY gonadal dysgenesis, showed renal failure resulting from “interstitial nephritis”. While the patient was receiving immunosuppressive therapy, a gonadoblastoma was discovered after renal transplantation. However, the authors did not attribute the development of neoplasia to immunosuppression. Harkins et al also observed renal failure of unknown origin. Similarly, we have observed renal failure in a patient with XY gonadal dysgenesis who not only had gonadoblastoma but also myotonic dystrophy, an autosomal dominant trait.

**Case report**
The proband was born to a 35-year-old father and a 32-year-old mother. The patient, her father, and her

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paternal aunt had muscle disease, cataracts, and other stigmata of myotonic dystrophy. The diagnosis of myotonic dystrophy was established in the proband by muscle biopsies and electromyography. Several other males in the family were said to show premature balding, but none complained of weakness. The proband’s male and female sibs were unaffected.

Infancy and early childhood development were considered normal in the proband. Although active and healthy at that time, routine urine analysis at 17 years of age revealed albuminuria. Over the next 2½ years the patient was admitted to hospital on several occasions because of persistent proteinuria; haematuria was not present. Intravenous urography and cystoscopy showed no abnormalities, but proteinuria continued. At the age of 20 years, the patient developed ankle oedema. One year later she noted easy tiring, occasional nausea, headaches, blurred vision, and hypertension that was controlled with methyldopa. Renal function (creatinine clearance tests) was less than 5% of normal and peritoneal dialysis was required. At the age of 21 years the patient was admitted for renal transplantation.

She had undergone no spontaneous secondary sexual development by the age of 17 years. At that time, administration of birth control pills led to withdrawal uterine bleeding, but cessation of hormones resulted in no subsequent uterine bleeding. No further hormones were administered, but at
the age of 19 years the patient experienced spontaneous uterine bleeding for several consecutive months. Breast development and pubic hair development also occurred.

At 21 years the patient was 162 cm tall. Blood pressure was 130/80. Facial muscles were atrophic and her face was expressionless. Her grasp showed typical myopathic behaviour, confirmed in the lower extremities and in the upper right extremity by electromyelography. Slit lamp examination showed no cataracts. Scalp hair distribution was normal. Breast development was at Tanner stage 2. External genitalia were normal and the uterus was small but well-differentiated. Initially, a pelvic examination revealed no abnormalities. However, pelvic x-rays, made as a result of genetic consultation, revealed a calcified pelvic mass, following which a right adnexal mass was detected on pelvic examination. Chromosomal analysis of lymphocytes revealed a 46,XY complement, without evidence of mosaicism. Buccal epithelial cells contained Y chromatin but not X chromatin.

At the age of 21, the patient received a renal transplant from her mother. The graft was placed in the patient’s right iliac fossa through a retroperitoneal approach. Bilateral nephrectomy was subsequently performed. The left and right kidneys weighed 44 and 45 g, respectively; both showed advanced total glomerular sclerosis with marked tubular atrophy and interstitial fibrosis. The walls of the small arteries and arterioles were characterised by marked fibrous intimal thickening and medial hypertrophy consistent with hypertensive nephropathy. The findings were those of an end-stage renal disease. No definite aetiological designation was possible.

After normal renal function was established by the transplanted kidney, exploratory laparotomy for the calcified mass revealed a left streak gonad, a right gonadoblastoma, and a small uterus and cervix. The 2 cm left gonad consisted of an irregular fibrous stroma, hilar cells, and rete ovarii; neither oocytes nor testicular structures were observed. The 13 g right gonad consisted of a pink-tan, glistening nodule of rubbery consistency; no definitive ovarian or testicular tissue was identified. Two-thirds of the mass was fibrotic, hyalinised, and calcified (figure a). The non-calcified areas showed sheets of primitive germ cells; small lymphocytes were present in the fibrous tissue stroma. The germ cells were large and showed abundant pale cytoplasm and darkly stained nuclei (figure b), a pattern similar to that of a dysgerminoma or seminoma. Overall, the features suggest a ‘burned-out’ gonadoblastoma characterised by the overgrowth of the germ cells with lymphocytic stroma, extensively replaced by fibrosis, hyalinisation, and calcification.

The patient now receives hormonal replacement therapy and has normal renal function.

Discussion

The patient reported here had three rare disorders: XY gonadal dysgenesis, myotonic dystrophy, and end-stage renal disease. The coexistence of the first two could be merely coincidental, but the third requires comment. In particular, this is the third patient with XY gonadal dysgenesis who has shown end-stage renal disease requiring kidney transplantation. The patient we report had end-stage renal disease of unknown aetiology, as did that of Harkins et al. The patient reported by Blanchet et al had “interstitial nephritis”. Different pathological processes thus may or may not have occurred in these patients. If the processes are similar, genetic heterogeneity or pleiotropy for the XY gonadal dysgenesis mutant could exist. Possibly of relevance is the occurrence of a nephrotic syndrome in subjects with XY gonadal dysgenesis, and coexistence of XY gonadal dysgenesis, renal disease (glomerulonephritis or nephrotic disease), and Wilms’ tumour.

Interestingly, the patient reported here showed spontaneous menstruation and breast development, possibly attributable to hormone production by the gonadoblastoma. Patients with XY gonadal dysgenesis ordinarily show no evidence of oestrogen production. Thus, puberal feminisation associated with XY gonadal dysgenesis strongly suggests the presence of a gonadoblastoma or dysgerminoma, either of which should be extirpated.

References


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