Three siblings with complete androgen insensitivity syndrome

Abstract

A 27-year old woman presented with primary amenorrhoea and infertility. On examination, she was found to have palpable inguinal gonads, normal female external genitalia, a blind-ending vagina with no cervix, almost complete absence of axillary and pubic hair, and good breast development. Further investigations confirmed the absence of internal female genitalia, the presence of a 46,XY karyotype and testosterone levels in the high normal male reference range. A diagnosis of complete androgen insensitivity syndrome was made, based on this clinical picture and laboratory findings. Two of her younger siblings were subsequently also diagnosed with this condition. She underwent an orchidectomy and is currently on female hormone replacement therapy. At the time of writing up this case study, her two younger sisters were still awaiting surgery.

Case study

A 27-year old female African patient from Mpumalanga was referred to the Endocrinology Clinic at Steve Biko Academic Hospital with primary amenorrhoea and infertility. She was referred from the cardiology department after admission for a supraventricular tachycardia. A diagnosis of Wolff-Parkinson-White syndrome was made. She had sought help, four years prior to this referral, at the gynaecology clinic, but was lost to follow-up before the workup was completed.

She was one of six siblings, with a 26-year old brother and four younger sisters. There was no history of consanguinity in the family. Of her younger sisters, the youngest, aged 11, was prepubertal and premenarchal, and the 16-year old sibling had normal menses. Both the 19-year old and the 21-year old siblings experienced primary amenorrhoea. The patient reported a usual female gender role and female gender identity, and had strong maternal feelings. She had already adopted a child, but wanted biological children with her husband.

On examination, it was found that she had a normal female phenotype, but was relatively tall at 1.71 m. She reported that she was much taller than her mother, and was the same height as her father. Her body mass index was 26. She was found to have palpable inguinal gonads which caused some discomfort, and normal female external genitalia, with no cliteromegaly or ambiguous genitalia present. Her vagina was short and blind-ending, with no palpable cervix. The patient reported having experienced severe dyspareunia for the first two years after becoming sexually active. She had good breast development, but an almost complete absence of axillary and pubic hair.

A transvaginal sonar confirmed the absence of a uterus and intraabdominal ovaries. The inguinal sonar detected a right gonad measuring 2.9 x 2.8 x 1.2 cm.

Table I: Laboratory findings in the index case

<table>
<thead>
<tr>
<th>Laboratory test</th>
<th>Value</th>
<th>Normal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Follicle-stimulating hormone</td>
<td>3.2</td>
<td>Female: 1.8-22.5 IU/l</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Male: 1.3-19.3 IU/l</td>
</tr>
<tr>
<td>Luteinising hormone</td>
<td>18.9</td>
<td>Female: 1.2-103 IU/l</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Male: 1.2-8.6 IU/l</td>
</tr>
<tr>
<td>Testosterone</td>
<td>22.9</td>
<td>Female: 0.34-2.6 nmol/l</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Male: 6.07-27.1 nmol/l</td>
</tr>
<tr>
<td>Sex hormone-binding globulin</td>
<td>158.1</td>
<td>Female: 18.2-135.7 nmol/l</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Male: 13.2-89.5 nmol/l</td>
</tr>
<tr>
<td>Calculated bioavailable</td>
<td>3.31</td>
<td>Female: 0.13-1.3 nmol/l</td>
</tr>
<tr>
<td>testosterone</td>
<td></td>
<td>Male: 2.8-15.5 nmol/l</td>
</tr>
<tr>
<td>Oestradiol</td>
<td>341</td>
<td>Female: 99-1 590 pmol/l</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Male: &lt; 73-173 pmol/l</td>
</tr>
</tbody>
</table>

*: Depending on the stage of the menstrual cycle
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and a left one measuring 2.5 x 1.1 x 2.2 cm (Figure 1 a-c). The patient’s laboratory findings are summarised in Table II.

The serum testosterone values were in the adult male range, with an elevated luteinising hormone (LH). Chromosomal studies detected a 46,XY karyotype. This confirmed the diagnosis of a form of 46,XY disorder of sex development. Taking into account the clinical picture, especially the lack of ambiguous genitalia, the lack of secondary sexual hair, her female phenotype and the sonar findings, in combination with testosterone levels in the high normal male range, the diagnosis was most likely to be that of complete androgen insensitivity syndrome (CAIS). Ovotesticular disorder of sex development (especially in view of her high oestradiol level), and 5-alpha-reductase type 2 deficiency were also considered, but were much less likely in view of the clinical findings and special investigations.
CAIS was previously called complete testicular feminisation. The reported incidence in the literature varies from between approximately one in 20 000 genetic males in Denmark, to one in 99 000 in the Netherlands.1,2 No data are available on how common this condition is in South Africa. Only one large case series has been described.4 It forms part of a wide spectrum of disorders caused by mutations in the gene coding for the androgen receptor, leading to resistance to androgens.1 Patients can present as phenotypically normal females, or with ambiguous genitalia in the case of partial androgen insensitivity syndrome, or in milder forms, as normal men with undervirilisation or fertility problems. These conditions are classified under the 46,XY disorders of sex development.2 The rest of this discussion will focus on CAIS.

Aetiology

Sentinel work was performed by Marcelli, Tilley and Griffin, and McPhaul, Marcelli and Zoppi, who identified the different mutations that caused CAIS.4,5 Since the androgen receptor gene is on the X chromosome, the inheritance of this condition is X-linked recessive. It is caused by numerous mutations in the gene coding for the androgen receptor, leading to severe impairment in androgen receptor function. More than a thousand mutations have been identified so far.6 Unfortunately, we could not obtain sequencing of the gene mutation in this family.

Reported abnormalities include missense mutations, premature stop codons, deletions and splicing abnormalities. Loss-of-function mutations in the coding sequence of the androgen receptor can be found in most patients with CAIS. Several other defects have been identified in recent years.9,10 In a large kindred from Soweto, the genetic cause was found to be the missense mutation, D732Y, in exon 5 of the androgen receptor ligand-binding domain.4 Whereas mutations in the androgen receptor are natural, environmental endocrine disruptors have been implicated in some cases of PAIS.11

Clinical features

Occasionally, babies are diagnosed with CAIS, even prenatally or at birth, because of a discrepancy between the karyotype determined through amniocentesis and the anatomical gender.1,5 Babies with CAIS have phenotypically normal female external genitalia which are not ambiguous. A small percentage of these children present with inguinal hernias. In retrospective studies, the incidence of CAIS was estimated to be approximately 0.8–2.4% in girls with inguinal hernias. This
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was confirmed in a prospective study which found an incidence of 1.1%. Some patients are also diagnosed at a young age because of the presence of palpable gonads in the inguinal area or in the labia majora. The majority present later during puberty with primary amenorrhoea. It is probably not unusual for patients to present much later in developing countries, as occurred in our index case. The hallmark of this condition on examination is normal breast development, normal external female genitalia, with the absence of or scanty pubic and axillary hair. Very few other disorders of sex development have this combination of clinical findings. Other confirmatory findings are an absent, short or blind-ending vagina, an absent cervix and often palpable gonads in the inguinal area or in the labia majora. Testes can also be intra-abdominal. Special investigations confirm the absence of a uterus, a 46,XY karyotype, and demonstrate serum testosterone levels in the normal adult male range. Body habitus and breast development is similar to that of normal females. These patients have a usual female gender identity with normal maternal instincts. They tend to be taller and larger than would be expected following a calculation of mid-parental height. The mean height is 171.5 cm, as was also observed in these three sisters. Usually, the growth pattern follows that of girls, but the final achieved height is between that of average males and females. It is postulated that the Y chromosome may have a direct effect on growth and body size which is not mediated through the androgen receptor. The bone mineral density is decreased in the lumbar spine, even in subjects where gonadectomy, followed by oestrogen replacement, is performed early. This suggests an important role for androgens in bone mineral density that cannot be replaced by oestrogen.

Laboratory findings

Our three patients had typical laboratory findings of CAIS. The serum testosterone level was in the normal adult male range, or slightly higher. Serum sex hormone-binding globulin concentrations can be increased. The testosterone production rate in the testes is increased because of an increase in serum LH. This is because of resistance to the negative feedback of androgens on LH secretion at hypothalamic-pituitary level. Serum FSH levels are usually normal. Oestrogen production is increased by up to 70% higher than that of normal men. This oestrogen originates from excess testosterone which is peripherally aromatised to oestrogen, as well as from the testicular secretion of oestrogen.

Diagnosis

Diagnosis is confirmed by finding a 46,XY karyotype in the presence of the above clinical findings, with no female internal genitalia detected on imaging, and a serum testosterone level in the adult male range. The coding region of the gene that encodes the androgen receptor can be sequenced to find the specific mutation.

Management

Gender assignment is not an issue in CAIS, even if diagnosed early. These patients should be raised as females. Full disclosure now forms part of the management plan. In the case of a minor, the parents should be fully informed and counselled, preferably by a psychologist who is familiar with this condition as disclosure of the diagnosis can lead to strong emotional reactions in the parents.

The child may receive information based on age and understanding, but full disclosure should occur before adulthood. Extensive counselling and psychological support may also be required. The damage caused by non-disclosure and finding out the diagnosis inadvertently, much later, can cause more psychological distress. Female family members should also be offered testing. However, there are ethical and legal implications, especially with regard to patient confidentiality.

Dilator therapy to increase vaginal depth and size can be started after puberty before sexual activity is contemplated. This may be sufficient, but occasionally vaginoplasty may be necessary. In a recent study, the majority of patients reported significant problems with desire, arousal and dyspareunia. Long-term sexual problems can occur, especially if surgical results are suboptimal. Sexual problems can occur if the vagina remains small after dilator therapy or vaginoplasty.

The risk of malignancy is increased in patients with cryptorchid testes, which is why these three sisters were referred for gonadectomies. Germ cell tumours or gonadoblastomas can occur in the testes, especially if located intra-abdominally, and can become malignant. A recent review of adults with CAIS estimated the risk of a gonadal malignancy developing of approximately 14% (0-22%). Older figures have estimated the risk as being greater than 30% by 50 years of age, but these included patients with PAIS who, in general, have a much higher risk than patients with CAIS.

Because the development of these tumours in the testes before puberty is very rare, it appears to be
safe to defer gonadectomy until after puberty.18,19,26,30 This will optimise normal breast development and may reaffirm the patient’s gender identity. Surgery may be deemed necessary earlier in the case of an inguinal hernia requiring surgical repair, if the gonads are uncomfortable or painful, or if there is doubt about the completeness of the androgen resistance. In the last scenario, incomplete resistance may lead to virilisation during puberty.24,30

Lifelong hormone replacement therapy is necessary post-gonadectomy in adults, or from the time of puberty if surgery was performed earlier, especially to optimise the already lower-than-normal bone mineral density.14,18,19 Whereas biological offspring has been impossible in the past, fertility is potentially becoming more feasible. If the testes can be preserved after gonadectomy, it may possibly for these patients to have biological children of their own in the future, with intracytoplasmic injection of primitive germ cells or spermatocytes (which are often present) into a donor ovum, and with the assistance of a surrogate.29,31

**Histology**

The testes of these patients are of a similar appearance to those of cryptorchid testes in boys. The number of Leydig cells can be normal or increased. Usually, there is no spermatogenesis.29,32 Sertoli cell nodules, as found in our index case, or Sertoli cell adenomas, are common, but are not premalignant. The epididymis and vas deferens are usually absent, but can be histologically identified adjacent to the testes in up to 36% of patients.32 Carcinoma in situ or germ cell malignancies are rarely identified, but usually only occur in postpubertal patients.32

**Acknowledgements**

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**References**