Clinical Characteristics of 46,XX Males with Congenital Adrenal Hyperplasia

**Short Running Title:** 46,XX Adrenal Hyperplasia Raised as Male

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**Abstract**

**Objective:** We aimed to retrospectively evaluate the follow-up data in patients with 46,XX congenital adrenal hyperplasia who were raised male.

**Methods:** A national database was created. The data of patients were asked to be recorded in the data form.

**Results:** The median age of diagnosis was 3 (0.1-18.3) years in 44 patients. Twenty nine cases were diagnosed after age of two years. Ninety five point four percent of cases were stage 4-5 virilized. Hysterectomy and bilateral salpingoopherectomy at the median age of 7.25 (2.4-25.3) were performed in 35 cases. Testicular prosthesis was placed in 11 (25%) cases. Median age of testicular prosthesis replacement was 11.2 (2.8-17) years. The final height was mean 149.2 cm (132.8-172) in 38 patients (simple virilizing (n=18), salt-wasting (n=6), 11 beta hydroxylase (n=12)). Of the 16 patients above the age of eighteen, university education was completed in 25% of cases.

**Conclusion:** It was seen that most of the 46,XX CAH cases raised male (2/3) were diagnosed after 2 years of age. In these cases, hysterectomy and bilateral salpingoopherectomy, genital corrective surgeries and testicular prosthesis operations were performed in a very wide age rage.

**Keywords:** 46,XX, congenital adrenal hyperplasia, final height

**Introduction**

According to consensus guidelines published in 2006, severe virilization during the neonatal period should not be considered a criterion for male sex determination, and 46,XX congenital adrenal hyperplasia (CAH) cases should be raised as females (1). This recommendation is supported by the fact that most of these patients identify as females; moreover, they retain their sexual and reproductive function (2). However, the age at diagnosis, age of initiation of glucocorticoid treatment, and degree of virilization were not considered. Some studies that considered these factors proposed that severely virilized patients with 46,XX CAH can be raised as males (3-5). Gender determination at birth and appropriate family and social support may be important in shaping sexual identity (5,6). Lee et al. suggested that the social and cultural environment is important to the gender identity of severely virilized 46,XX CAH patients (4). Gender dissatisfaction in adolescence and adulthood is high among patients with brain virilization caused by perinatal androgen exposure, particularly in girls with a delayed CAH diagnosis (7). Daae et. al. performed a systematic review of the literature, investigating sexual orientation in individuals with CAH. This recently published review included 30 studies investigating sexual orientation in patients with CAH assigned female at birth (46,XX) (n=927) or assigned male at birth (46,XY and 46,XX) (n=274). A majority of assigned females with CAH self-identified (defined themselves) as heterosexual, but figures varied widely across studies (40-100%). Results indicate that fewer assigned females reported homosexual (3-20%) or bisexual orientation (3.4-37%). The rates of non-heterosexual orientation were higher in assigned females with CAH than controls, whereas no individuals with CAH assigned male (46,XY or 46,XX) expressed any non-heterosexual orientation (8). Patients with CAH are often diagnosed late in countries without a newborn screening program for the disorder. Such patients tend to be raised as males by their families. Even virilized CAH patients diagnosed early may be raised as males in male-dominant societies. Follow-up data on 46,XX CAH patients raised as males are insufficient. Experiences in 46,XX CAH patients raised as males are mostly in the form of case reports. In this study, we planned to evaluate these cases followed up...
with this diagnosis in our country. Therefore, this study aimed to create a national database of 46,XX CAH patients including retrospectively collected diagnostic and follow-up data.

**Materials Methods**

This study included 46,XX patients with CAH raised as males (21 hydroxylase and 11 beta hydroxylase-deficient CAH patients; analyzed as a single group given their similar clinical characteristics). The diagnosis of CAH was made based on medical history, physical examination, and cytogenetic and hormonal analyses. The diagnosis of salt-wasting 21 hydroxylase deficiency was made on the basis of findings of salt-wasting, ambiguous genitalia and elevated 17-hydroxyprogesterone levels. The diagnosis of simple virilizing 21 hydroxylase deficiency was made on the basis of findings ambiguous genitalia and elevated 17-hydroxyprogesterone levels without salt wasting. The diagnosis of 11 beta hydroxylase deficiency was made on the basis of findings of ambiguous genitalia and elevated 11-deoxycortisol levels. Cytogenetic studies confirmed the karyotype to be 46,XX. Patients with a history of additional chronic systemic disease or chronic drug use for reasons other than CAH were excluded from the study.

National-scale projects in Turkey, such as the current study, are supported by the Pediatric Endocrinology and Diabetes Association. After receiving approval from the Pediatric Endocrinology and Diabetes Association the details of the project, including the start and end dates for data collection (July 11, 2018 and January 31, 2019, respectively), were sent to all participating pediatric endocrinology clinics via e-mail. The clinics were also provided with access to online data entry forms (cedd.saglik-network.org site). Adult endocrinology clinics were not invited to take part in the study. The data form is provided in the supplemental material. Diagnostic and follow-up data were obtained retrospectively from patient records.

The genital virilization levels of the patients were evaluated by pediatric endocrinologists at the time of diagnosis and staged according to the Prader classification (9).

The Hospital Clinical Research Ethics Committee approved this study (No: 15/2018).

**Results**

**Total Number and Distribution of Patients**

Data for 44 patients from seven Pediatric Endocrinology Centers were analyzed. In seven pediatric endocrinology centers, the total number of patients with 46,XX CAH was 439. Of these 439, 44 were raised as male. Of the 44 CAH cases, 30 (68.1%) were CYP21A2-deficient CAH and 14 (31.8%) were CYP11B1-deficient. Of the CYP21A2 deficient CAH cases, 10 (33.3%) were of the salt-wasting type and the remainder were of the simple virilizing type. The characteristics and clinical findings of the patients at diagnosis and the final follow-up are given in Table 1.

**Age and Virilization Stage in Diagnosis**

The median age at diagnosis was 3 years (range: 0.1–18.3 years). Fifteen patients (34%) were diagnosed at < 2 years of age, while 29 (65.9%) were diagnosed at ≥ 2 years of age. Eight patients were diagnosed during the neonatal period. Only two cases (4.6%) were Prader stage 3 (one case early diagnosis 11 beta hydroxylase deficiency, one case late diagnosis simple virilizing 21 hydroxylase deficiency); the remaining 42 cases (95.4%) were stage 4 or 5 (13 and 29 cases, respectively).

**Surgeries**

Hysterectomy and bilateral salpingoopherectomy were performed in 34 cases, at a median age of 7.25 years (range: 2.4–25.3 years) (early diagnosis cases 7.8 years (3.5-12), late diagnosed cases 7.8 (2.4-25.4)); in 6 cases (17%), these procedures were performed at ≤ 3.5 years of age.

Genital reconstructive surgery was performed in 12 (27.2%) patients; 2 of these patients were Prader stage 3, and 10 were Prader stage 4. Genital reconstructive surgery was performed once in four cases, twice in six cases, and four times in two cases. The first genital corrective surgery was performed at the median age of 8.5 years (range: 2–14.5 years).

A testicular prosthesis was placed in 11 (25%) cases. The testicular prosthesis was replaced at a median age of 11.2 years (range: 2.8–17 years) (early diagnosis cases 15.5 years (14–17), late diagnosed cases 9.8 (2.8-14)). Breast development occurred in 12 cases with bilateral oophorectomy, 4 of whom underwent a mastectomy. Some of the patients did not consent to surgery.

**Treatment**

All patients were started on glucocorticoid replacement treatment at the time of diagnosis (hydrocortisone and hydrocortisone-equivalent steroid treatment in 30 and 14 patients, respectively). However, steroid treatment compliance was low, at 56.8%.

Testosterone treatment was started in 21 cases. The mean age at initiation of testosterone treatment was 14 ± 1.85 years (range: 10-17 years) and the mean duration of the treatment was 4.76 ± 2.63 years. Testosterone depot forms were started at a dose of 50 mg/4 weeks. The dose was increased by 50 mg over 6 months, and the full dose of 250 mg was reached in two years. Only three cases (14.2%) were unsuitable for testosterone treatment. Testosterone treatment was started in 13 patients at the age of ≥14 years. The age of initiation of testosterone treatment in eight patients was between the ages of 10-14 years. Testosterone treatment was not started in 8 cases despite an age > 14 years.

**Duration of Follow-up and Final Height**

The median age at the last examination was 15.3 years and the mean follow-up duration was 10.7 years (range: 0.1–24 years). A total of 28 (63.6%) patients were aged > 14 years. Thirty-eight patients (18 patients with simple virilizing 21 hydroxylase deficiency, 6 patients with salt wasting 21 hydroxylase deficiency, 14 patients with 11 beta hydroxylase deficiency) reached their final height during the follow-up. The mean final height was 149.2 cm (range: 132.8–172 cm).

**Education and Job**

Of the 16 patients aged > 18 years, two were primary school graduates, four were high school students, six were high school graduates, two were university students, and two were university graduates. University education was completed by 25% of the patients.

Regarding employment type, one patient was a shepherd, one had an assembly job, one was an office apprentice, one was an “asphalt worker” and one was a chemical engineer. Only 5 of the 10 adult patients (2 primary school graduates, 6 high school graduates, and 2 university graduates) had a job with medical insurance.
Discussion

Our study is the largest case series of 46,XX CAH patients raised as males reported to date. Although it has been recommended by physicians that patients diagnosed as newborns or during early infancy be raised as girls, families may desire to raise their child as a boy in male-dominated societies. In our study, the majority of the 46,XX CAH patients raised as males were diagnosed after the age of 2 years, but approximately one-third were diagnosed at an earlier stage. The upbringing of 46,XX CAH patients diagnosed early and identified as male may depend on the culture of the country of birth. In the literature, the majority of 46,XX CAH patients raised as males were diagnosed late (4,10-13). Few patients are diagnosed before the age of 2 years (3). In our cohort, it was recommended that female patients diagnosed at < 2 years of age be raised as females, particularly those diagnosed during the neonatal period. However, some families insisted on raising their child as a male. Although family preferences and sociocultural factors are important, virilization also plays a role in the decision to raise these patients as males. In studies from countries with different sociocultural contexts, most patients with 46,XX CAH raised as males are Prader stage 4 or 5 (3,10,13,14). In our study, 95.5% of the patients were Prader stage 4 or 5; only two (4.5%) were Prader stage 3. According to the literature, few Prader stage 2–3 patients are raised as males, similar to our study (12,15,16).

According to the current consensus, surgeries leading to irreversible infertility should not be performed in patients with sex development disorders until their sexual identity is clear (17). However, hysterectomy and bilateral salpingoophorectomy are almost always performed during childhood in 46,XX CAH patients (3,13-15). In our series, the median age of patients who underwent hysterectomy and bilateral salpingoophorectomy was 7.25 years (range: 2.4–25.3 years); the age range was wide. In eight (22.8%) patients, hysterectomy and bilateral salpingoophorectomy were performed at the age of < 2.5 years. These surgeries were performed at the age of 25.5 years in a patient diagnosed with 46,XX CAH when he was 18 years and 4 months old. This patient had not presented to the endocrine department prior to being diagnosed with 46,XX CAH. He was diagnosed with an undescended testicle at the ages of 5 and 8 years, but no further examinations were performed. In 33 other patients, hysterectomy and bilateral salpingoophorectomy were performed before the age of 15 years. About 68% of the cohort were aged < 10 years, which is the average age of onset of puberty, and the surgery was performed without a full assessment of sexual identity. Hysterectomy and bilateral salpingoophorectomy operations should be performed only after the sexual identity of the child becomes clear, i.e., when they can adequately express their desire in that respect. If the sexual identity is not clear, puberty can be delayed until the oophorectomy is performed.

A testicular prosthesis was placed in 11 patients (25%) in our series and the median age of implantation was 11.2 years (range: 2.8–17 years). Similar to the literature, the rate of testicular prosthesis placement was low and the age at placement varied (3,13-15). There is no standard age for testicular prosthesis placement, and no recommendations regarding changing the testicular prosthesis of a patient to one of a different size (3,13-15). Therefore, this is left to the discretion of the clinicians.

Breast development is particularly obvious during puberty in 46,XX CAH patients raised as males. Breast development may also occur in patients who are unsuitable for steroid treatment; breast surgery may be required in these cases (10,13). In our series, breast development occurred in 12 patients; mastectomy was performed in 4 of these cases. Hysterectomy and bilateral salpingoophorectomy were performed in 10 of 12 cases with breast development. This suggests that treatment incompatibility may have an etiological role. It should be emphasized (both to patients and their families) that patients who need a large number of genital corrective operations must comply with treatment to prevent the requirement for an additional mastectomy.

Compliance with testosterone treatment was generally good among our patients. However, steroid compliance was poor and approximately half of the patients were unsuitable for steroid treatment. Without steroid treatment, 46,XX CAH patients may exhibit increased levels of androgens, potentially making them feel better. However, to avoid the negative effects associated with a lack of steroid treatment (short stature, risk of adrenal crisis at any age), the importance of treatment compliance should be impressed upon patients and their families.

Thirty-eight patients reached their final height during the follow-up. The mean final height was 149.2 cm (range: 132.8–172 cm), which is too short for individuals to continue living as men. In all children with CAH, boys and girls, androgen excess causes accelerated bone maturation and growth and reduces adult height. In these children, hydrocortisone replacement therapy is very challenging: overdosing results in growth inhibition and excessive weight gain, whereas underdosing results in accelerated bone maturation and short adult height (18). A meta-analysis reviewed adult height data until 2008 and confirmed height loss: the mean adult height in salt wasting and simple virilizing patients was -1.38 SD-score (-1.56 to -1.2) (19). Brottes P et al. monitored French CAH children from the pre-screening era and found that it was shorter adult height than general population (mean: -1.2 SD (156.7 cm) in girls and -1 SD (168.8 cm) in boys. In comparison with the general French population, short AH (< -2SD score) was seen in 24% of the cohort (22.5% of girls and 26% of boys) and present a dramatic advanced 8-year bone maturation that influences strongly the risk of short adult height (20). Several authors have found a better height outcome in patients who received fludrocortisone (21). A higher pubertal hydrocortisone dose was associated with a slightly higher risk of short adult height. This may be due to well-known negative effect of excessive glucocorticoid doses on growth (21,22). High hydrocortisone doses may reflect a poorly controlled disease either because of disease severity or secondary to poor compliance to the treatment. The adult heights achieved are actually below the average and 20% of adult CAH patients have a short adult height (below -2 SD) (20). The final height achieved in patients with 46,XX CAH is very important, particularly if they are to be raised as males, given its psychosocial effects. Woelfle et al. reported a patient who attempted suicide due to short stature (11). Families of 46,XX CAH patients raised as males should be informed that their children will not be fertile, and that they will be short. In these cases, early and long-term growth hormone treatment and aromatase inhibitors may be beneficial (23). If growth hormone treatment is to be used (which should be for a protracted period), planning should begin at the youngest possible age.

Follow-up of 46,XX CAH cases raised as males is important during adulthood to assess quality of life; further studies on this subject are needed. Only 25% of our 16 cases aged > 18 years received a university education. According to the statistics of
the Statistical Institute in our country, the ratio of faculty or high school graduates to the population was 15.7%. It was observed that the education rate of our patients was not lower than the general population.

All patients were followed up in the pediatric endocrine clinic. Clinics facilitating the transition to adulthood should be established for 46,XX CAH patients aged > 18 years raised as males. A multidisciplinary approach should be taken, with follow-up throughout the life course to assess endocrine function and psychiatric status.

The main strengths of our study were the inclusion of a large number of 46,XX CAH patients raised as males, and the availability of follow-up data. It is important to note that this study included a sample drawn from clinics nationwide, such that the outcomes should be generalizable. A limitation of our study was that some of the patient data were obtained retrospectively. Also, comparing 46,XX CAH patients raised as females with those raised as males during the same period would have provided more valuable information.

No guidelines for the care and management of 46,XX CAH patients raised as males are available. The results of our study, which is the large case series of 46,XX CAH patients raised as males conducted to date, can be summarized as follows: The 46,XX CAH patients raised as males were diagnosed late, and included cases of advanced virilization. Surgeries that eliminated the potential for fertility were mostly performed without a full assessment of gender identity. The testicular prosthesis placement rate, and the age at placement, were highly heterogeneous. The onset of testosterone treatment was late, and the rate of treatment was inadequate. Steroid treatment compliance was poor, and the final height of most patients was short for males. We recommend that irreversible surgeries that impair the possibility of fertility should be avoided unless an explicit self-consent is obtained. A consensus should also be sought regarding testicular prosthesis replacements, and the importance of growth hormone treatment compliance to final height should be emphasized to 46,XX CAH patients raised as males. Sexual identity assessments should be performed periodically, and clinics should be established to facilitate the transition to adulthood. Finally, given the importance of sharing experience, adult follow-ups visits should be scheduled.

References


Table 1. Clinical follow-up characteristics of 46,XX CAH patients raised male

<table>
<thead>
<tr>
<th>Clinical Characteristics</th>
<th>Value</th>
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<tbody>
<tr>
<td>Age of diagnosis (mean ±SD year)</td>
<td>4.2 ± 4.2 (0.1-18.3)</td>
</tr>
<tr>
<td>Age of last follow-up (mean ±SD year)</td>
<td>14.9 ± 5.7 (-24.9)</td>
</tr>
<tr>
<td>Duration of follow-up (mean ±SD year)</td>
<td>10 ± 6.1 (0.1-24)</td>
</tr>
<tr>
<td>Distribution of patients by age of diagnosis</td>
<td></td>
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<tr>
<td>New born</td>
<td>(n=8)</td>
</tr>
<tr>
<td>&lt; 2 years</td>
<td>(n=7)</td>
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<tr>
<td>Pre-schooler</td>
<td>(n=18)</td>
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<tr>
<td>School-aged</td>
<td>(n=5)</td>
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<tr>
<td>Adolescents</td>
<td>(n=5)</td>
</tr>
<tr>
<td>Adult (≥ 18 years)</td>
<td>(n=1)</td>
</tr>
<tr>
<td>Virilization Prader stage in diagnosis</td>
<td></td>
</tr>
<tr>
<td>Stage 3</td>
<td>n=2 (4.5%)</td>
</tr>
<tr>
<td>Stage 4</td>
<td>n=13 (29.5%)</td>
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<tr>
<td>Stage 5</td>
<td>n=29 (65.9%)</td>
</tr>
<tr>
<td>Age of hysterectomy and bilateral salpingoopherectomy (median year) (n=34)</td>
<td>7.25 (2.4-25.3)</td>
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<tr>
<td>Number of patients who had more than one operation</td>
<td>n=13 (38.2%)</td>
</tr>
<tr>
<td>Testicular prosthesis placement age (n=11) (median year)</td>
<td>14 (2.8-17)</td>
</tr>
<tr>
<td>Steroid treatment compliance</td>
<td>Good (n=19) (42.2%)</td>
</tr>
<tr>
<td>Poor (n=26) (57.7%)</td>
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<tr>
<td>Testosterone treatment start age (mean ±SD year) (n=21)</td>
<td>14 ± 1.85 (10-17)</td>
</tr>
<tr>
<td>Testosterone treatment compliance</td>
<td>Good (n=18) (85.7%)</td>
</tr>
<tr>
<td>Poor (n=3) (14.2%)</td>
<td></td>
</tr>
<tr>
<td>Final height (cm) (n=38)</td>
<td>149.2 (132.8-172)</td>
</tr>
</tbody>
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