Psychological impact of congenital adrenal hyperplasia on adolescent and young girls in Saudi Arabia.

Nasir A. M. Al- Jurayyan1*, Amal A. Al-Hakami2
1Department of Paediatrics, College of Medicine, King Saud University, Saudi Arabia
2College of Medicine, Princess Nourah Bint Abdulrahman University, Riyadh, Saudi Arabia

Abstract

Background: Congenital adrenal hyperplasia (CAH) consists of a family of defects in the synthesis of steroid hormone in the adrenal cortex which results in high androgen levels. The androgen excess is present from early embryogenesis and results in varying degrees of virilization of the external genitalia. It can lead to major medical and psycho-social consequences.

Design and setting: Eight Saudi adolescent and young women whose age ranged between 15 to 25 y with variable severity of CAH were studied to determine the psychosocial impact at the endocrine service, King Khalid University Hospital (KKUH) Riyadh, Saudi Arabia in the period July 2017 to June 2018.

Objective: The aim of this study was to determine the psychological impact of CAH on adolescent and young women who were diagnosis and appropriately reared in the early life.

Methods: Eight Saudi adolescent and young women with similar clinical characteristics where chosen from a cohort of sixty four patients with CAH. Psychosocial information were gathering during a clinic visits from both participant and parents. Participants were interviewed by the primary treating endocrinologist (principal author NJ) to facilitate trust and confidence, utilizing the child behaviour checklist (CBCL) questioner.

Results: There were significant differences between females with CAH diagnosed early in life and reared as females with higher incidence of anxiety, depression (25%), withdrawn/depressed behaviour (25%) and somatic complaints (75%), which depend on the variation of the severity.

Conclusion: Psychological adjustment is variable in females with CAH who were diagnosed and properly reared in early life. Adjustment depends on the severity of the disease. Future multicentre studies involving CAH patients in Saudi Arabia are needed to ensure large sample size. In addition, further researchers should concentrate on various aspects of psychosocial issues of CAH.

Keywords: Psychological impact, Congenital adrenal hyperplasia, Adolescent, Young women, Saudi Arabia.

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Introduction

Congenital adrenal hyperplasia (CAH) consists of family of defects in the synthesis of steroid hormones in the adrenal cortex. It is caused by a defect in 21-alpha-hydroxylase gene (CYP21) in more than 90% of the cases. However, the rest were due to 11-beta-hydroxylase enzyme deficiency and 3-beta-hydroxysteroid dehydrogenase deficiency. The enzyme deficiency results in impaired synthesis of cortisol and aldosterone. The low cortisol levels result in increased production of adrenocorticotropic hormone (ACTH) by the pituitary, which cause increase synthesis of steroid precursors resulting in high androgen levels. The androgen excess is present from early embryogenesis and results in varying degrees of virilization of the external genitalia in girls. CAH is the most common reason for ambiguous genitalia at birth. It requires lifelong medications, frequent hospital visits, and genital surgery [1-6]. In Saudi Arabia where the consanguineous matting is high, there are no precise data on the prevalence of the disease; however, there are impression fostered by clinical experience and local reports of congenital adrenal hyperplasia that this is not an uncommon disease. A calculated incidence had been postulated, 1 in 5000, from the number of patients diagnosed with the disease among the deliveries at King Khalid University Hospital (KKUH) Riyadh Saudi Arabia. Presently, the limited local new-born screening program supported such data [3].

Previous studies had shown that CAH affects all aspects of health-related quality of life, and that adolescent girls and young adult women have an increased risk of psychiatric morbidity. However, there is still a lack of knowledge about quality of life determinants and optimal psychiatric adaptation [7,8].

Although advances in care and medical knowledge have been significantly important, the prognosis for patients living with
CAH still need to comply with life restrictions required to manage the disorder, i.e., lifelong medications [9,10].

Further behaviour studies will provide valuable information that would help in determining whether any intervention can be instituted to minimize the long-term effects.

This article reports on our experience at King Khalid University Hospital (KKUH), endocrine service, Riyadh, Saudi Arabia of the psychological impact of CAH on adolescent and young women.

Material and Methods

This is a qualitative descriptive study conducted at paediatric endocrine service, King Khalid University Hospital (KKUH) of King Saud University during the period between July 2017 to June 2018.

The aim of this study is to assess the psychological adjustment of adolescent girls and young women with CAH. Eight Saudi Arabian adolescent girls and young women whose age ranged from 15 to 25 y where chosen from a cohort of sixty four patients with variable causes and similar clinical characteristics (Table 1). Prader classification was used to describe masculinization of external genitalia at diagnosis [11]. Genital surgery was done early in life, so nothing remembers about this. Psychosocial information were gathering during a clinic visits from both participant and parents. Participant was interviewed by the primary treating endocrinologist (principal author NJ) to facilitate trust and confidence, utilizing the child behaviour checklist (CBCL) questioner [12]. Statistical package for social science (SPSS) version 21, (Chicago, IL, USA) was used for the statistical analysis of the data.

Table 1. The clinical characteristics of patients with congenital adrenal hyperplasia (CAH).

<table>
<thead>
<tr>
<th>Serial number</th>
<th>Age at study</th>
<th>Diagnosis</th>
<th>Clinical presentation</th>
<th>Number of surgeries</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>18 y</td>
<td>21-OH-ase deficiency (4)</td>
<td>(+)</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>16 y</td>
<td>Congenital adrenal hyperplastic (CAH)</td>
<td>(+)</td>
<td>3</td>
<td>Sister of “1”</td>
</tr>
<tr>
<td>3</td>
<td>22 y</td>
<td>21-OH-ase deficiency (5)</td>
<td>(+)</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>21 y</td>
<td>Congenital adrenal hyperplastic (CAH)</td>
<td>(+)</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>15 y</td>
<td>11-OH-ase deficiency (5)</td>
<td>(+)</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>25 y</td>
<td>21-OH-ase deficiency (2)</td>
<td>(+)</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>18 y</td>
<td>11-OH-ase deficiency (5)</td>
<td>(+)</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>14 y</td>
<td>21-OH-ase deficiency (4)</td>
<td>(+)</td>
<td>2</td>
<td>Sister of “7”</td>
</tr>
</tbody>
</table>

A.G: Ambiguous Genitalia; SW: Salt-Wasting; SV: Simple Villilizing; Prader: Prader Classification; y: year; 21-OH-ase: 21-hydroxylase; 11-OH-ase: 11-hydroxylase.

Results

Eight adolescents’ girls and young women with variable CAH were included in this study. Their demographic features were stated in (Table 1). Two sitters with 11b-hydroxylase deficiency and six with 21 hydroxylase deficiency. Five salt-wasting (SW) and one simple virilization (SV). All reported good relations with parents and the experience of having to take responsibility. Parents had similar level of education as well above average income. All had at least one genital surgery (clitoroplasty or and vaginoplasty) with a mean age of one year (range; 3 to 18 months). All had at least an undergraduate education with an average and median 15 y.

Our overall results revealed that higher incidence of anxiety, depression (25%) and withdrawn/depressed behaviour (25%) and somatic complaints (75%). However, female with CAH felt within the normal range of other parameters.

Only one young female (12.5%) with 21-hydroxylase deficiency CAH, simple virilizing married and had one child, she reported that she is happy in her life. No participants reported suicidal thoughts, or suicidal attempts.

The participants were from a conservative Muslim community who believe that sexuality issues are private and personal, therefor they decided to express their feelings during interview.

Discussion

Congenital adrenal hyperplasia (CAH) consists of a family of defects in the synthesis of steroid hormones in the adrenal cortex. It is an inborn condition following an autosomal recessive inheritance. More than 90% of disorders is caused by a deficiency in the enzyme 21 hydroxylase (21OHD), while other forms such 11 b hydroxylase deficiency (11OHD), and 3- beta-hydroxysteroid dehydrogenase deficiency (3 B HSD) compromised the rest CAH has long term health problems affecting several body systems [1-6].

The diagnosis must be based on history and physical examination supported by the accumulation of ACTH-stimulated steroid precursors above the enzymatic block, with
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the exception of lipid hyperplasia (P450 scc deficiency) in which almost no steroids are produced. Also, an experienced paediatric radiologist, as part of a multidisciplinary team, plays an important role as the sonographic appearance of the adrenal gland, the cerebriform appearance, is characteristic, and the internal genitalia can be determined. This can be supported by other modalities such as genitography, computed tomography (CT) scan, and magnetic resonance imaging (MRI) [1-7].

Thus, the complexity of the disorder (CAH) which can lead to various health problems emphasizes the need for care providers to establish a multidiscipline team approach to minimize both medical and psychological consequences. The team should consist of paediatric endocrinologist, genetics, gynaecologist, urologist, paediatric surgeon, psychologist or psychiatrist, plastic surgeon and specialist nurse. Other discipline should be included according to clinical needs [5,10,13].

Although various studies documented the medical aspect of CAH, yet to our knowledge this is the first psychological study in Saudi Arabia. The aim of this study was to assess the psychological impact of CAH on Saudi Adolescent and young girls, and bring to the attention of the care provider of such need. There were multiple studies worldwide of psychological assessment of adjustment of women with CAH, depend on nature of the methodology used [14-27]. Our patients exhibit high rate of psychopathology disorders. Among the disorders, depression and anxiety [28] were the most prevalent. The relevance of androgens in the development of psychopathology has been suggested based on sex related differences in the prevalence, course and treatment response of several psychiatric disorders [29].

Although, the aetiology of anxiety disorders is not fully understood, evidence from both preclinical and clinical studies support the contribution of corticotropin-releasing hormone (CRH) in the expression of anxiety related disorders, mainly through CRH neuro circuits connecting the amygdala and locus coeruleus. In the CAH patients increased CRH secretion is expected in the untreated or undertreated state. This increased CRH secretion reflects a block in cortisol production leading to the production of CRH and subsequent increased CRH secretion [30]. Shepard et al. [31] found decreased amygdala volume in CAH patients suggesting that alteration in the hypothalamic-pituitary-adrenal axis affect the growth and development of amygdala.

The finding of psychological well-adjusted girls with CAH may be attributed to successful genital surgery, adequate medical treatment, and appropriate social support [7]. Other studies suggested that chronic illness and other difficult life experiences have only temporary effect on adjustment [32,33].

Several studies indicated reduce fertility in women with CAH. Pregnancy rates in patients with salt wasting (SW) CAH and simple virilizing (SV) forms were similar [34]. There are multiple causes for reduced fertility in females with CAH, including improper treatment.

Finally, this study is not without any limitations. Firstly, patients were diagnosed infancy with ambiguous genitalia or salt wasting and reared as females. Therefore, it was not possible to highlights the effects of the wrong sex assignments. Secondly, all participants were selected into our study; these individuals may be relatively healthy and interested in the therapies measures provided. Lastly but not the least, the smaller size of the study group (only eight patients) and the limited answers will not allow appropriate and adequate statistical power. However, this article hopefully, will stimulate researchers to design new studies that attempt to model some degree of complexity of lives of children, adolescent, and young adults’ women with the disease CAH.

In conclusion, psychological adjustment is variable in females with CAH who were diagnosed and appropriately reared in early neonatal life. Adjustment depends on the severity of the disease rather than the patient and parent level of education and income.

Future multicentre studies involving CAH patients in Saudi Arabia is needed to ensure large sample size in addition, further researchers should concentrate on various aspects of psycho-social outcome of CAH in Saudi children.

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References


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*Correspondence to
Nasir A. M. Al-Jurayyan
Department of Paediatrics
College of Medicine
King Saud University
Saudi Arabia