Primary Adrenal Insufficiency (PAI): A Major Teaching Hospital Experience, Riyadh, Saudi Arabia

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Abstract

**Background:** Primary adrenal insufficiency (PAI) in children is an uncommon, but potentially fatal. The current symptoms include weakness, fatigue, anorexia, abdominal pain, weight loss, orthostatic hypotension, salt craving and characterized by hyperpigmentation.

**Material and Methods:** This is a retrospective, hospital based-study, conducted at King Khalid University Hospital (KKUH), during the period January 1989 and December 2014. Review of medical record of patient diagnosed with primary adrenal insufficiency. The diagnosis was based on medical history, physical examination and low levels of glucocorticoids and raised adrenocorticotropic hormone (ACTH). Appropriate laboratory and radiological investigations were also reviewed.

**Results:** During the period under review, January 1989 and December 2014, a total of 125 patients with the diagnosis of primary adrenal insufficiency were seen. Inherited disorders like congenital adrenal hyperplasia and hypoplasia were common, 85.5%. However, variable autoimmune mediated etiologic diagnosis accounted for, 13%, were also seen. The appropriate various laboratory and radiological investigations should be planned.

**Conclusion:** Although, congenital adrenal hyperplasia was the commonest etiology, however, congenital adrenal hypoplasia should not be over looked. The diagnosis of PAI can be challenging in some patients, and therefore appropriate serological and radiological investigations should be done.

**Keywords:** Etiology; Primary Adrenal Insufficiency; Saudi Arabia.

Introduction

Primary adrenal insufficiency (PAI) in children is an uncommon, but potentially fatal and life-threatening condition. The clinical symptoms of adrenal insufficiency include weakness, fatigue, anorexia, abdominal pain, weight loss, orthostatic hypotension, salt craving and characterized by hyperpigmentation. It comprises a heterogeneous group of both congenital and acquired disorders [1-6]. Congenital adrenal hyperplasia, is a group of autosomal recessive disorders resulting from the deficiency of one of the enzymes required to synthesis cortisol, was the commonest [7, 8]. Auto-immune induced adrenal insufficiency either isolated of as a part of autoimmune polyendocrine syndromes have been described, but the frequency of these occurrence in children with PAI has not been determined. Various mutations in genes, and autoantibodies to gland-specific target antigens such as 21-hydroxylase have been well described [9, 10].

The objective of this study was therefore, to define the etiology of PAI in a large referral, teaching hospital in Riyadh, Saudi Arabia.

Materials and Methods

All patients followed by the author at the Endocrine Unit at the King Khalid University Hospital between January 1989 and December 2014, with the diagnosis of primary adrenal insufficiency (PAI) were reviewed. The diagnostic categories retained were as follows: congenital adrenal hyperplasia (CAH), congenital adrenal hypoplasia, Addison's disease, autoimmune polyendocrine syndromes, triple A syndrome (Allgrove syndrome), adrenoleukodystrophy and bilateral adrenal haemorrhage. Data reviewed included patient's age, sex, clinical presentation, laboratory and radiological investigations. Congenital adrenal hyperplasia (CAH) was diagnosed as recommended [7].
The presence of anti-adrenal antibodies in serum was evaluated by indirect immunofluorescence. Mutational analysis of the AIRE gene was sequenced as described [9].

**Results**

During the period under review January 1989 and December 2014, a total of 125 patients were seen at the Endocrine Unit, King Khalid University Hospital, Riyadh, Saudi Arabia, with the diagnosis of primary adrenal insufficiency Table 1. There were 103 patients, 46 males and 57 females, with CAH aged between newborns to 13 years of age. Of these, 84 (81.1%) patients, 36 male and 48 females, were having CAH due to 21-α-hydroxylase deficiency, with salt-wasting in 77 (91.7%) patients, 15 (14.6%) patients, 7 males and 8 females, patients with 11-β-hydroxylase deficiency which only 4 (3.9%) patients were due to 3-β-hydroxysteroid hydrogen deficiency, all were salt-waster. Table 2.

Congenital adrenal hypoplasia was also diagnosed in 4 patients, from 2 families having history of neonatal deaths. Adrenal haemorrhage is relatively an uncommon condition with a variable and non-specific presentation seen in two patients. Figure 1, showing abdominal computed tomography (CT) revealing adrenal haemorrhage.

Adrenal insufficiency related to autoimmune disorders was the second most common, accounting to approximately 11%. Addison's disease, as an isolated adrenal insufficiency in 7 patients, while in 6 patients it was associated with autoimmune polyendocrine syndrome Type 1 (APS-1).

X-linked adreno-leukodystrophy (ALD), an inherited disorder, due to the deficiency of ligoro-ceroyl-coA ligase (synthetase) enzyme, required for synthesis of long-chain-fatty acid was present in one patient. Figure 2 of magnetic resonance imaging (MRI) with characteristic changes. Allgrove syndrome, a rare autosomal disorder which is characterized by symptoms of alacrima, achalasia, and adrenal insufficiency.

**Discussion**

Adrenal insufficiency is the clinical manifestation of deficient production or action of glucocorticoids, with or without deficiency also in mineralocorticoids and adrenal androgen. It is a life-threatening disorder that can result from variable aetiologies, of which, primary adrenal failure was the commonest [1-6]. Congenital adrenal hyperplasia (CAH), is the common (82.3%) disorder, encountered in our study. It is an autosomal recessive disorder caused by reduced or complete absence of the enzymatic activities of steroid biosynthesis pathway [7]. In Saudi Arabia, this is not a usual finding in a country with increased rate of consanguinity [11-13]. It should be differentiated from congenital adrenal hypoplasia. In contrast, congenital adrenal hypoplasia, is relatively a rare X-linked recessive disorder characterized by glucocorticoid and mineralocorticoid deficiency with low levels of androgen and normal external genitalia. Some rare cases develop adrenal failure only in adulthood [14, 15]. Bilateral adrenal haemorrhage is one of other causes of adrenal insufficiency presented in two infants, in this series. It is rare potentially life-threatening condition. The diagnosis is often complicated by its non-specific presentation and its tendency to intervene in stressful critical illness. Due to many coagulation disorders, haemorrhage is a major cause of morbidity and mortality. It can be diagnosed by ultrasonography, computed tomography (CT) and magnetic resonance imaging (MRI). The exact incidence of the condition is unknown. It is classically associated with meningococcemia (Waterhouse-Friderichsen syndrome), but may occur with any septic process. Traumatic events, burns, antiphospholipid syndrome, heparin associated thrombocytopenia, thrombophilic syndrome, anti-coagulation therapy, and abdominal surgery are some of the other causes of adrenal bleeding. The pathophysiology is still unclear. However, some particular anatomic features of the glands, such as a rich arterial blood supply that feeds into a dense and delicate sub-capsular capillary network [16, 17].

Autoimmune adrenal failure was the second leading cause of PAI, Table 1. Etiology of primary adrenal insufficiency (PAI) in 125 patients.

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>No.</th>
<th>%</th>
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<tbody>
<tr>
<td>Genetic disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>- congenital adrenal hyperplasia</td>
<td>103</td>
<td>82.3%</td>
</tr>
<tr>
<td>- congenital adrenal hypoplasia</td>
<td>4</td>
<td>3.2%</td>
</tr>
<tr>
<td>Bilateral adrenal haemorrhage</td>
<td>2</td>
<td>1.6%</td>
</tr>
<tr>
<td>Autoimmune-mediated</td>
<td>14</td>
<td>11.3%</td>
</tr>
<tr>
<td>- Addison disease</td>
<td>7</td>
<td>-</td>
</tr>
<tr>
<td>- Autoimmune polyendocrine syndrome type 1</td>
<td>6</td>
<td>-</td>
</tr>
<tr>
<td>- Adreno-leukodystrophy</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>Triple A (Allgrove) syndrome</td>
<td>2</td>
<td>1.6%</td>
</tr>
</tbody>
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Table 2. Distribution of 103 patients with congenital adrenal hyperplasia (CAH) and enzyme deficiency.

<table>
<thead>
<tr>
<th>Enzyme deficient</th>
<th>No. of patients</th>
<th>%</th>
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<tbody>
<tr>
<td>21-α-hydroxylase</td>
<td>84</td>
<td>81.60%</td>
</tr>
<tr>
<td>11-β-hydroxylase</td>
<td>15</td>
<td>14.60%</td>
</tr>
<tr>
<td>3-β-hydroxysteroid dehydrogenase</td>
<td>4</td>
<td>3.90%</td>
</tr>
</tbody>
</table>

in our series, accounting for 11% of cases. It can present as an isolated condition or can be associated with other autoimmune disorders (autoimmune polyendocrine syndrome). Patients with autoimmune polyendocrine syndrome usually developed primary adrenal insufficiency earlier and had a history of chronic candidiasis and/or hypoparathyroidism. All patients with adrenal failure due to autoimmune syndrome had positive auto-adrenal antibodies [9, 18]. Adrenal failure was the common presentation in X-linked adrenoleukodystrophy (ALD) [19, 20].

Conclusion

The diagnosis of primary adrenal insufficiency (PAI) can be challenging. Congenital adrenal hyperplasia was the commonest 82.3% diagnosis, with autoimmune disorders being the second found in 11%.

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References