Case report

A novel SCNN1A variation in a patient with autosomal-recessive pseudohypoaldosteronism type 1

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What is already known on this topic ?
Autosomal-recessive pseudohypoaldosteronism type 1 (PHA1) is a rare genetic disorder caused by different variations in the ENaC subunit genes. Most of these variations appear in SCNN1A mainly in exon eight, which encodes for the alpha subunit of the epithelial sodium channel ENaC. Variations are nonsense, single-base deletions or insertions, or splice site variations, leading to mRNA and proteins of abnormal length. In addition, a few new missense variations have been reported.

What this study adds ?
We report a novel mutation [ c.729_730delAG (p.Val245Glyfs*65) ] in the exon 4 of the SCNN1A gene in case of autosomal recessive pseudohypoaldosteronism type 1. Patient with PHA1 requires early recognition, proper treatment, and close follow-up. Parents are advised to seek genetic counseling and plan future pregnancies.

Abstract
Pseudohypoaldosteronism type 1 (PHA1) is an autosomal-recessive disorder characterized by defective regulation of body sodium levels. The abnormality results from mutations in the gene-encoding subunits of the epithelial sodium channel (ENaC). Patients with PHA1 present in infancy as being in adrenal crisis. A 41-day-old female who presented with recurrent adrenal crisis did not adequately respond to hydrocortisone and required mineralocorticoid therapy.

The patient's demographic data and clinical features were recorded. Blood samples were collected and tested for endocrine and metabolic characteristics and for use in genetic studies. Bidirectional Sanger sequencing of SCNN1A was conducted. The entire coding region of 12 exons and 20 bp of flanking intron were sequenced. Genetic analyses revealed a new mutation—c.729_730delAG (p.Val245Glyfs*65)—in SCNN1A exon four.

Adrenal crisis during the neonatal period highlights the importance of early screening for PHA1. Genetic testing could help to anticipate the prognosis, severity, onset of the disease, and the mode of inheritance, especially with its extensive phenotype.

Keywords: Pseudohypoaldosteronism, hyperkalemia, hyponatremia, adrenal crisis, congenital adrenal hyperplasia.

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Background
Pseudohypoaldosteronism type 1 (PHA1) is a rare disorder of mineralocorticoid resistance that is characterized by defective regulation of body sodium levels because of the inability of aldosterone to exert its effect on target tissues (i.e. aldosterone resistance). PHA1 is a life-threatening disease that presents during the neonatal period with dehydration, severe salt wasting, and failure to thrive accompanied by hyponatremia, hyperkalemia, metabolic acidosis, and increased plasma renin, with an elevated aldosterone level that is consistent with aldosterone resistance despite normal renal and adrenal functions (1, 2). PHA1 can be divided into renal (PHA1A, OMIM #600983 ) and systemic (PHA1B, OMIM #264350) types based on its physiologic and genetic characteristics. Renal type PHA1A is an autosomal-dominant disorder with heterogeneous inactivating mutations in the NR3C2 gene that codes for the aldosterone receptor (3, 4). The renal autosomal-dominant type is characterized by sodium loss that is restricted to the kidneys. It is usually less severe with gradual clinical improvement during the first several years of life, thus allowing sodium supplementation to be terminated at some point. In contrast, the systemic PHA1 type is a pseudohypoaldosteronism-recessive inheritance pattern caused by inactivating mutations in any subunits of the epithelial sodium channel (ENaC) encoded by SCNN1A, -1B, -1G (5). The systemic autosomal-recessive type typically presents shortly after birth. It is characterized by sodium loss from the kidneys and other mineralocorticoid target tissues, such as the colon, lungs, salivary glands, and sweat glands, which increases the frequency of lower respiratory tract infections. The disorder is more common and severe in infancy, persists into adulthood, and requires lifelong therapy using sodium supplementation (6, 7).
There are no reports on the frequency of PHA1 in Arab populations or the Middle East; however, Al-Shaikh reported two Omani children who presented with PHA1 during their first week of life, both of whom came from a consanguineous family. The first child (a male) had a severe clinical course with skin manifestations, recurrent episodes of severe chest infections, and electrolyte imbalances. Feeding difficulty and treating the electrolyte imbalances were challenges for the treating physician. A genetic analysis was not conducted on this case. The second case (a female) had a milder clinical course with fewer episodes of respiratory infections. Her genetic analysis revealed a new mutation in the SCNN1A ENaC subunit (8).

Case Presentation
A 41-day-old female was referred to our hospital as a case of possible congenital adrenal hyperplasia. She was a product of a consanguineous marriage and an uneventful pregnancy. The patient was born at a gestational age of 37 weeks by normal spontaneous vaginal delivery with a birth weight of 3 kg without complications. At the age of 6 d, she began to have difficulties feeding, lethargy, and frequent vomiting. She was taken to the emergency room with severe dehydration and acidosis (arterial pH = 7.20; serum bicarbonate = 11 mmol/L), high potassium (K) (10 mmol/L), and low sodium (Na) (119 mmol/L). Cardiopulmonary resuscitation was initiated and the patient was intubated. The treating team considered the possibility of an adrenal crisis and conducted initial hormonal studies (Table 1). The patient was treated with a stress dose of hydrocortisone along with a fluid bolus and sodium bicarbonate (NaHCO3). Hyperkalemia was managed with calcium gluconate, insulin, and glucose. The patient was admitted to the neonatal intensive care unit and tested for respiratory and metabolic issues, both of which were negative. She was later discharged from the hospital and treated with hydrocortisone and fludrocortisone with no sodium chloride (NaCl).

This patient had an elder sibling and a cousin who died with similar presentations. At the age of 20 d, the newborn was readmitted to the hospital with episodes of vomiting, poor feeding, and lethargy and was found to have high K and low Na levels. The conditions were managed using a stress dose of hydrocortisone, a fluid bolus, and NaHCO3. Hyperkalemia was managed using calcium gluconate, insulin, glucose, and emergency peritoneal dialysis to lower her K levels. After the patient was stabilized, she was referred to our hospital for further investigation and management. On examination, the patient was conscious, not dysmorphic, and well hydrated. She had normal female genitalia and showed no systemic abnormalities. Our initial lab results revealed 138 mmol/L Na, 5 mmol/L K, and 23 mmol/L bicarbonate (HCO–3). The urinalysis and urine cultures were normal. A fluorescence in situ hybridization study was sent from the primary hospital, which was negative for the Sex-Determining Region gene, and chromosomal analysis revealed an XX female karyotype. The results of abdominal and pelvic ultrasound examinations were normal. The results of hormonal estimations are provided in Table 1.

Initially, our patient’s symptoms were managed with hydrocortisone and fludrocortisone treatment for possible congenital adrenal hyperplasia. However, because the initial lab results for hormone levels (Table 1) were not consistent with congenital adrenal hyperplasia, and the patient had presented with recurrent adrenal crisis since the neonatal period, she did not adequately respond to hydrocortisone and required mineralocorticoid therapy. In addition, follow-up examinations revealed papular skin rashes and recurrent chest findings, which suggested a diagnosis of pseudohypoaldosteronism. A sweat-chloride test showed elevated levels (128 mmol/L; normal range, 0–39 mmol/L for patients > 4 months).

Material and Methods
The present study was approved by the research and ethics committee of King Fahad Medical City (KFMC), Riyadh, Kingdom of Saudi Arabia (IRB log number 18-609). The patient’s demographic data and clinical symptoms were recorded. Blood samples were collected and tested for endocrine and metabolic characteristics and for use in genetic studies. Molecular genetic testing was conducted initially for NR3C2 sequencing for autosomal-dominant pseudohypoaldosteronism. Bidirectional Sanger sequencing of SCNN1A was then conducted. The entire coding region of 12 exons and 20 bp of flanking intron were sequenced. Informed consent for all tests was obtained from the patient’s parents.

Results
The results of the hormone tests on the patient are presented in Table 1. Molecular genetic testing for NR3C2 sequencing for autosomal-dominant pseudohypoaldosteronism. The results did not show any sequence variant that may have caused the disease. Based on the majority of reported mutations Bidirectional Sanger sequencing of SCNN1A was then conducted. The results revealed that the patient was homozygous for a novel mutation at c.729_730delAG in the exon 4 of SCNN1A (Fig. 1), which was predicted to result in a frameshift and premature protein termination (p. Val245Glyfs*65). Using in silico prediction tools, such as SIFT-indel and Mutation Taster, this variant is predicted to be deleterious (Table 2). Moreover, the amino acid valine in position 245 is highly conserved among different species (Fig. 2). This variant is not found in humans (PM2).

Targeted sequencing for that variant was conducted on both parents, who were found to be carriers for the sequence variant c.729_730delAG (p. Val245Glyfs*65) in SCNN1A in heterozygous form.

Discussion
“Adrenal crisis” was the initial impression, which included adrenal hypoplasia and congenital adrenal hyperplasia, and was based on the patient’s clinical presentation and a history of poor feeding, lethargy, frequent vomiting, consanguinity, family history of sudden infant death, hyperkalemia, hyponatremia, and metabolic acidosis. The patient was accordingly treated with hydrocortisone. She had normal female genitalia, hyponatremia, and hyperkalemia with normal androgen precursor levels, but a high level of plasma aldosterone, renin activity, and cortisol, all of which were more consistent with PHA1 than congenital adrenal hyperplasia.

Other differential diagnoses based on the results of the initial electrolyte levels were hyperkalemia and hyponatremia, including secondary PHA, which is a transient form of aldosterone resistance secondary to a urinary tract infection. A urinary tract malformation was also a possibility but was unlikely because the results of the urine tests and ultrasound were normal. Renal tubular acidosis type 1 is characterized by low levels of ammonia in the urine and is associated with hyperkalemia, mild hyponatremia, and metabolic acidosis with hyperaldosteronism or aldosterone insensitivity.
A poor response of the patient to corticosteroids indicated aldosterone resistance and either renal or systemic PHA1. Patients with PHA1 supposedly present in the neonatal period with hyponatremia, hyperkalemia, and metabolic acidosis, and the condition can be diagnosed by significantly elevated plasma aldosterone levels and renin activity (9-11).

The clinical manifestations of renal PHA1 may vary in asymptomatic patients and can only be diagnosed using presentations of hyperaldosteronism and elevated renin levels compared with patients presenting with a salt-wasting crisis. The clinical characteristics and severity of this condition can vary widely. Severe phenotype has been reported due to mutations in both alleles in the NR3C2 gene (29). Overall, PHA1 has a mild clinical course followed by remission over time (12-16). The systemic form of PHA1 leads to salt loss from organs that express ENaC, such as the kidneys, salivary and sweat glands, and the colon. Children typically present shortly after birth with electrolyte disturbances mimicking an adrenal crisis (4). Acute initial presentation, frequency of respiratory symptoms, and a positive sweat-chloride test led to the diagnosis of systemic PHA1 in our patient. In vitro screening of several mutant genes has led to advances in understanding the physiology of the mineralocorticoid receptors and ENaC; therefore, the underlying molecular pathology of renal and systemic PHA1 has been associated with mutations in the mineralocorticoid receptor and ENaC subunit genes SCNN1A, SCNN1B, and SCNN1G, respectively (11). Aldosterone plays a central role in electrolyte homeostasis and maintenance of fluid in the distal nephron. Loss-of-function mutations in two key components of the aldosterone response—the mineralocorticoid receptor and ENaC—lead to PHA1, a rare genetic disease of aldosterone resistance characterized by salt loss, dehydration, failure to thrive, hyperkalemia, and metabolic acidosis (13).

The prickly heat-like skin rash in our patient, which was typically aggravated by salt deprivation, is a characteristic feature of systemic PHA1. As a consequence of ENaC expression, this manifestation is consistent with recent studies that have reported that skin manifestations are common in systemic PHA1 patients as are other phenotypic features, such as cholelithiasis, polyhydramnios, and hypercalciuria leading to hypocalcemia. (17-23).

A sweat-chloride test is positive in both systemic PHA1 and cystic fibrosis because both disorders lead to salt loss from the sweat glands. The gold standard for a diagnosis of cystic fibrosis is based on a clinical diagnosis plus an abnormal sweat-chloride test and the determination of a mutation in the cystic fibrosis transmembrane conductance regulator gene CFTR; therefore, based on our patient’s initial clinical presentation and her symptom improvement after salt supplementation and potassium exchange resins, cystic fibrosis was a less-likely diagnosis and a systemic PHA1 diagnosis was not necessary in this case (24).

CFTR encodes a membrane protein and chloride channel, which play important roles in transepithelial salt absorption and secretion in some epithelial tissues, such as the intestine, airways, and sweat glands, and reproductive system. Studies have shown that CFTR modulates the function of epithelial Na(+) channels (ENaC) in salt absorption, but the mechanism is still controversial. CFTR mutations lead to an inadequacy in the CFTR–ENaC relationship, which may lead to increased Na+ absorption in the airways, resulting in hypersecretion of mucus, decreased mucociliary clearance, and bacterial colonization (25-27).

PHA1 was first reported by Cheek and Perry in 1958 as a sporadic occurrence with a severe salt-wasting syndrome; however, the number of patients with PHA1 has increased over the past decade. Since 1958, the literature has mainly consisted of case reports or case series. Based on HGMD, more than 71 NR3C2, 47 SCNN1A, 47 SCNN1B, and 20 SCNN1G mutations have been reported in the literature. Most of the reported mutations were single-base deletions/insertions, or splice-site mutations with severe courses; missense mutations usually follow a mild course. Several new mutations have been reported with phenotypic variations (22, 24). In 2017, Nur (23) reported one case and reviewed 27 others from clinical presentation to follow-up and outcome. In 2018, Cayir (25) summarized the diagnosis and current management of PHA1.

To the best of our knowledge, this is the first recognized case of PHA1 with a new mutation that results in a frameshift and premature protein termination (p.Val245Glyfs*65), which leads to a sequence variant designated as c.729-730delAG in SCNN1A exon four. This finding confirmed the hypothesis that an autosomal-recessive or a sporadic PHA1 systemic type is a genetically heterogeneous disease involving other unidentified genes (14).

However, null variant (frameshift), in SCNN1A gene is a known mechanism of disease, single nucleotide deletion in the same coding region has been classified as pathogenic SCNN1A:c.729delA which leads to different frameshift (p.Val304TrpfsTer4). This reported mutation is associated with lung symptoms(recurrent lower respiratory tract infections) without development of chronic lung disease, that is a similar finding to our case (28).

Renal type 1 PHA1, an autosomal-dominant disease resulting from heterogeneous mutations in the NR3C2 gene that encodes for the mineralocorticoid receptor, results in a condition of aldosterone not being able to bind to its receptor. Severe phenotype has been reported due to mutations in both alleles in the NR3C2 gene. Approximately 50 distinct mutations have been reported, but there have been no reports on genotype/phenotype correlations (11). In contrast, systemic type 1 PHA1 is an autosomal-recessive disorder that typically presents during the neonatal period with severe symptoms, such as hyponatremia, hyperkalemia, and metabolic acidosis. This condition can be diagnosed by significantly elevated plasma aldosterone levels and renin activity with the symptoms persisting into adulthood (14, 16, 17).

Treatment of PHA1 should be considered a challenging emergency, and life-saving measures are needed to rectify dehydration, replace sodium loss, and correct hyperkalemia and acidosis during the acute phase (4).

Renal PHA1 can be managed with salt supplementation, which is often decreased over the lifetime of the patient. Although systemic PHA1 requires aggressive management, including salt supplementation, intensive fluid administration, potassium exchange resins, and dietary manipulations to reduce hyperkalemia, the use of indomethacin may be beneficial in some patients and when treating systemic complications of the disease. The course of the disease is often interrupted by recurrent episodes of salt-loss crises with hyperkalemia associated with nausea, vomiting, and difficulty feeding, which might require gastrostomy/jejunostomy feeding, as well as a chronic pulmonary syndrome with or without associated chest infections (17, 18).

Sodium supplementation may be given in the form of NaHCO₃, NaCl (1 g contains ~17 mEq Na), or sodium citrate. Up to 110 mEq/kg/d can be given. Generally, it is difficult for patients to tolerate a high dose of salt orally; therefore, the dose
intake can be divided and mixed with baby formula and food. It may need to be administered by nasogastric tube/jejunostomy in poor tolerance situations (17, 25).

Severe hyperkalemia and acidosis can be managed by the intravenous slow infusion of NaHCO₃ and then oral replacement when the condition stabilizes. A combination of glucose and insulin infused slowly at the usual ratio of 1 unit of insulin to every 5 g dextrose is an effective measure by which to reduce serum potassium. Salbutamol infusion or continuous nebulisation has been added as adjunctive therapy. In severe persistent hyperkalemia, peritoneal dialysis is used, after which calcium chloride is infused to protect the heart from arrhythmias until the serum potassium levels are normalized. Potassium exchange resins, which act as potassium-binding agents, are widely used in hyperkalemia management. The most commonly used potassium-binding agents are sodium polystyrene sulfonate and calcium polystyrene sulfonate; however, the sodium-containing resin is preferred to the calcium-containing resin because it simultaneously corrects both hyponatraemia and hyperkalemia. It is used orally or rectally at approximately 1–2 g/kg/dose every 6 h when high doses administered orally cannot be tolerated. In these cases, rectal administration of the drug is a good option (19, 25).

Genomic testing for a carrier state in the asymptomatic siblings and the parents is highly recommended to detect homozygosity or compound heterozygosity (8).

**Study Limitations**

One limitation to the present study was that we did not measure testosterone levels during the patient’s hospitalization.

**Conclusion**

Adrenal crisis during the neonatal period highlights the importance of early screening for PHA1 to facilitate early recognition, proper treatment, close follow-up, and genetic counseling for the family. Parents should be advised to seek genetic counseling and testing when planning future pregnancies.

**References**

Table 1: Results of hormonal studies on the patient

<table>
<thead>
<tr>
<th>Investigation</th>
<th>Results</th>
<th>Normal range</th>
</tr>
</thead>
<tbody>
<tr>
<td>17-hydroxyprogesterone</td>
<td>72 ng/dL</td>
<td>(5–115)</td>
</tr>
<tr>
<td>Serum renin activity</td>
<td>23. nmol/L/hr</td>
<td>(4–12)</td>
</tr>
<tr>
<td>Serum aldosterone</td>
<td>4244.2 pmol/L</td>
<td>(1000–3800)</td>
</tr>
<tr>
<td>Serum ACTH</td>
<td>5 Pmol/l</td>
<td>1.6–13.9</td>
</tr>
<tr>
<td>Serum cortisol</td>
<td>94.1 µg/dL</td>
<td>(2.3–11.9)</td>
</tr>
<tr>
<td>Dehydroepiandrosterone sulfate (DHEA-s)</td>
<td>13 ng/dL</td>
<td>(5–35)</td>
</tr>
</tbody>
</table>

Table 2: in-silico analysis of the SCN1A gene mutation founded in our patient and ACMG classification

<table>
<thead>
<tr>
<th>Mutation (nucleotide)</th>
<th>Protein change</th>
<th>Mutation Taster</th>
<th>SIFT-indel</th>
<th>gnomAD Exomes MAF</th>
<th>ACMG criteria</th>
<th>Classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>c.729_730delAG</td>
<td>p.Val245Glyfs*6</td>
<td>Disease causing</td>
<td>Damaging</td>
<td>-</td>
<td>PVS1 PM2 PP3</td>
<td>Pathogenic</td>
</tr>
</tbody>
</table>

PVS1 (Very Strong): Null variant (frame-shift), in gene SCN1A, for which loss-of-function is a known mechanism of disease (gene has 8 pathogenic LOF variants and LOF Z-Score = 1.97 is greater than 0.7), associated with Pseudohypoaldosteronism, type I, Liddle syndrome 3 and Bronchiectasis with or without elevated sweat chloride 2. PM2 (Moderate): Variant not found in gnomAD exomes (unable to check gnomAD exomes coverage). PM3: Variant not found in gnomAD genomes (good gnomAD genomes coverage = 30.5). PP3:Variant from controls (at extremely low frequency if recessive) in Exome Sequencing Project, 1000 Genomes Project, or Exome Aggregation Consortium. PP3 (Supporting): Pathogenic computational verdict based on 1 pathogenic prediction from GERP vs no benign predictions.
Figure 1: the electropherogram of the novel variant in SCNN1A: c.729_730delAG

Figure 2: conservation of the amino acid (Valine) in the position 245 of the amino acid sequence among different species