Acute intermittent porphyria caused by novel mutation in HMBS gene, misdiagnosed as cholecystitis

Majid Alfadhel1,3
Neam Saleh2
Helal Alenazi2
Henry Baffoe-Bonnie2
1Division of Genetics, Department of Pediatrics, 2Division of General Medicine, Department of Medicine, King Abdulaziz Medical City, Riyadh, Kingdom of Saudi Arabia; 3College of Medicine, King Saud Bin Abdulaziz University for Health Sciences, Riyadh, Kingdom of Saudi Arabia

Background: Acute intermittent porphyria (AIP) is an autosomal dominant neurovisceral inherited disorder due to a defect in the heme biosynthesis pathway. Misdiagnosis of the porphyrias is not uncommon.

Case report: We present a case of a 26-year-old female with suspected acute cholecystitis, mental status changes, and seizures. Biochemical and molecular investigations confirmed the diagnosis of AIP by findings of elevated urinary porphobilinogen, 5-aminolevulinic acid, and total porphyrins. DNA molecular testing showed a novel heterozygous mutation (c.760delC p.L254X) in the exon11 of the HMBS gene. To the best of our knowledge, this is the first report of a misdiagnosis of AIP presenting with acute cholecystitis.

Conclusion: Clinicians are alerted to consider the possibility of AIP in an adult presenting with an acute abdomen, features of cholecystitis, and neuropsychiatric manifestations.

Keywords: porphyria, cholecystitis, acute abdomen, acute intermittent porphyria, HMBS gene, hydroxymethylbilane synthase

Introduction

Acute intermittent porphyria (AIP) (MIM#176000) is an autosomal dominant inborn disorder of heme biosynthesis, that is caused by a mutation in the gene encoding hydroxymethylbilane synthase (HMBS) or porphobilinogen deaminase.1 Currently, >386 mutations are known in HMBS gene, mostly, missense, nonsense, splicing site mutations, deletions, insertions, or duplications.2 Because AIP is rare and its clinical presentation is heterogeneous, it can be easily misdiagnosed. However, a diagnosis of AIP masquerading as acute cholecystitis has not been reported yet in the literature.

We describe a 26-year-old woman with AIP, misdiagnosed initially as cholecystitis and found to have a novel heterozygous mutation in the HMBS gene.

Case report

The Saudi female patient in the UK developed recurrent bouts of abdominal pain, nausea, and vomiting during first pregnancy, diagnosed as hyperemesis gravidarum, and was treated with IV fluids and antiemetics. There was improvement in the condition and the patient delivered a healthy child (in Saudi). After 2 months, the patient presented with a bout of severe abdominal pain (diagnosed as cholecystitis), followed by repeated seizures, hyponatremia, elevated and uncontrolled blood pressure, sinus tachycardia, delirium, lower limb weakness, hyporeflexia, and psychological abnormalities. She was intubated for 3 days. Initial postextubation, physical examination
result was unremarkable except for unexplained behavioral disturbances and lower limb weakness. Initial instigations showed sodium levels ranged between 118–125 mmol/L (135–144 mmol/L), mildly elevated transaminases (aspartate aminotransferase [AST]/alanine transaminase [ALT]), and slight elevation of anti-neutrophil antibodies of the perinuclear type (P-ANCA). Alkaline phosphatase, bilirubin, complete blood count (CBC), thyroid-stimulating hormone (TSH), vitamin B12 level – markers for connective tissue diseases, viral titer values, and cerebrospinal fluid (CSF) analysis findings were all within normal range. The abdominal imaging result was normal, except for contracted gallbladder and stones.

Urine became dark red on exposure to light. Total porphyrins in urine were elevated: 29,100 nmol/24 hours (normal: ≤214), with main elevation in uroporphyrin level, 24,200 nmol/24 hours (normal: ≤30), and mild elevation in coproporphyrin level, 118 nmol/24 hours (normal: ≤38). DNA molecular testing showed a novel heterozygous mutation in the exon 11 of the HMBS gene (c. 760delC p.L254X). The patient was treated with intravenous heme arginate (4 mg/kg/day) for 5 days, that was followed by an improvement in the chemical and clinical parameters.

Discussion
AIP manifests later in life; only 10% to 15% of those exhibiting carrier types develop the clinical syndrome. Typically, patients suffer intermittent bouts of neurovisceral attacks for up to several days, recurring over several weeks, manifesting as severe abdominal pain (>95% of cases) and neurological and/or psychological symptoms. Abdominal pain is associated with nausea, vomiting, and constipation. Psychological symptoms vary, including mood swings, depression, anxiety, and occasionally, auditory and visual hallucinations, disorientation, and mental deterioration. Neurological features can affect the central/peripheral nervous system, leading in some cases to seizure, severe myalgia, ascending flaccid paralysis and in catastrophic cases to respiratory arrest or irreversible quadriplegia. Other commonly encountered findings include difficult to treat hypertension, tachycardia and SIADH induced hyponatremia. External triggers may be porphyrinogenic drugs, alcohol, infections, calorie-restricted diet, and stress; internal triggers are hormonal (pregnancy and menstrual cycle).

Diagnosis of AIP is biochemical/molecular dependent. Biochemical testing, usually, shows increased concentrations of porphobilinogen, 5-aminolevulinic acid, and total porphyrins in the urine (mainly uroporphyrins), with decreased enzyme activity of porphobilinogen deaminase (≤50%) in erythrocytes. Fecal porphyrin levels are normal or increased. Diagnosis of AIP is confirmed by the detection of heterozygous mutation in the HMBS gene. In the presented case, the pathogenicity of this novel mutation is supported by creation of premature stop codon, which is very likely to result in truncated protein or loss of protein production.

AIP can be misdiagnosed as psychosis, Guillain–Barré syndrome, and chronic hepatitis (Table 1). The presented patient was misdiagnosed as having cholecystitis, as she presented with severe abdominal pain and gallbladder stones; however, the neuropsychiatric manifestations made the treating physicians rethink about the diagnosis and treatment. To avoid such misdiagnosis in the future, the clinicians should consider the possibility of AIP in any individual presenting with an acute abdomen and neuropsychiatric manifestations.

Attacks of AIP are typically triggered by stress; symptoms start during pregnancy. Perhaps, supportive care obfuscated manifestations of porphyrin attacks. Nonetheless, increased estrogen and progesterone levels in pregnancy may precipitate...
porphyric attacks. The currently recommended and effective treatments in the moderate to severe cases, are the early administration of IV human hemin along with glucose loading, discontinuation of triggering porphinogenic drugs, and symptomatic treatments of the concomitant hypertension, tachycardia, pain and neurological symptoms.

The cause of neuropsychiatric manifestations in AIP is not well understood. Several investigators suggested that increased levels of hepatic 5-aminolevulinic acid and porphobilinogen may play an essential role in the occurrence of such complications, and the supportive evidence of such theory is the finding that several patients with severe AIP became asymptomatic after liver transplantation. Other postulated mechanisms include focal demyelination, vacuolization of the neurons, and multifocal ischemia resulting from a vascular narrowing and/or glial proliferation. Treatment of such complications consists of prompt intervention with the aforementioned supportive measures; in addition, gabapentin proves its effectiveness in reducing neuropathic pain and seizure episodes.

**Conclusion**

AIP could be misdiagnosed as cholecystitis. Our report adds a novel nonsense mutation to the previously reported AIP mutations.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**