Acute Intermittent Porphyria with SIADH and Fluctuating Dysautonomia

Nabin A,1 LJ Thapa,2 Paudel R,3 Rana PVS4

INTRODUCTION

Porphyria is a rare metabolic disorder having autosomal dominant inheritance with incomplete penetrance with some cases having autosomal recessive and complex inheritance.1,2,3 The gene coding for enzymes has been characterized and several mutation in porphyria genes are reported.1 Though genetically heterogenous, different porphyrrias have a uniform clinical phenotype of the three hepatic porphyrias having acute presentation, acute intermittent porphyria (AIP) is commonest and usually manifest after puberty predominantly affecting young female population.1,4 It present with acute abdominal crisis and neuropsychiatric manifestations. When inadvertent administration of precipitating factors continues in misdiagnosed cases, a rapidly progressing sensorimotor neuropathy develops with respiratory and bulbar paralysis and dysautonomia. Advancement in ICU care, avoidance of precipitating factors and prompt treatment with haematin, has reduced the mortality from about 30% to less than 10%.5,6 In addition these patients need constant follow up for early detection of hepatoma, hypertension and renal impairment.7,8 and modifications of treatment of subsequent systemic diseases developing in them. Three cases of intermittent porphyria, diagnosed and treated are reported. Importance of recognition of syndrome of inappropriate antidiuretic hormone (SIADH) and potentially fatal dysautonomia is stressed and literature is reviewed.

CASE REPORT

Case 1: MS (HN 17023), a 20 years old lady, was admitted with severe generalized abdominal pain and vomiting. She gave history of similar episode, but of lesser severity, in the past. At admission she was afebrile with pulse 96/min, Respiration-20/min and BP 1140/90 mms of Hg. Per abdominal examination revealed diffuse tenderness. No lump, rigidity, guarding or free fluid detected. Rest systems were normal. Investigations i.e. complete hemogram, urinalysis, serum electrolytes, renal and metabolic parameters, ECG, USG abdomen were normal. She was...
Case 1: AG (HN: 214954), a 25 years old lady presented with acute nephrotic syndrome. Hypertension was noted on admission and was severe. She was afebrile with pulse 104/min, BP 160/110 mm Hg. She gave h/o of abdominal pain in the past lasting for five days duration. It was not associated with vomiting, bowel disturbances or genitourinary symptoms. However, she gave h/o of abdominal pain in the past lasting for variable period. At admission she was in agony and was afebrile with pulse 68/min, BP 160/130 mm of Hg. Abdominal examination revealed soft abdomen with diffuse tenderness. Bowel sounds were sluggish. No organomegaly, rigidity, free fluid detected. Rest systems were normal. She was initially diagnosed and treated as acute pancreatitis. Investigation revealed blood urea 56 mg/dl and serum creatinine 1.7 mg/dl. Other investigations i.e. including hemogram, urine analysis, blood sugar, serum electrolytes, liver function tests, serum amylase, ECG, X-ray chest & abdomen, USG abdomen, upper GI endoscopy and urinary catecholamine assay were normal. Subsequently, in view of accelerated hypertension, tachycardia possibility of porphyria was suspected and diagnosis confirmed by qualitative test for porphobilogen. She was treated with high parenteral carbohydrate intake, antihypertensive agent, chlorpromazine, pethidine and antispasmodics. She made gradual recovery with normalization of BP and marked reduction of abdominal symptoms. However, on day 8th of her admission she became drowsy, incontinent and lapsed into deep coma. In view of normalization of blood pressure, passing of normal colored urine, absence of localizing neurological deficits and normal pupils, possibility of SIADH suspected and confirmed. With continuation of high parenteral dextrose and restriction of fluid intake, she recovered gradually and was discharged on 17th day with advice to avoid precipitating factors.

Case 2: Gradual Clearing of urine with recovery. 

Figure 1(b). Case-2: Gradual Clearing of urine with recovery. 

Figure 1(a). Case-2:Change of urine color on exposure to sun (Case 1).

noting change of color of urine on exposure to sunlight (fig-1(b)) and qualitative test for porphobilogen.

She was treated with high parenteral carbohydrate intake, antihypertensive agent, chlorpromazine, pethidine and antispasmodics. She made gradual recovery with normalization of BP and marked reduction of abdominal

Case 2: AG (HN: 214954), a 25 years old lady presented with severe and continuous upper abdominal pain of five days duration. It was not associated with vomiting, bowel disturbances or genitourinary symptoms. However, she gave h/o of abdominal pain in the past lasting for variable period. At admission she was in agony and was afebrile with pulse 68/min, BP 160/130 mm of Hg. Abdominal examination revealed soft abdomen with diffuse tenderness. Bowel sounds were sluggish. No organomegaly, rigidity, free fluid detected. Rest systems were normal. She was initially diagnosed and treated as acute pancreatic. Investigation revealed blood urea 56 mg/dl and serum creatinine 1.7 mg/dl. Other investigations i.e. including hemogram, urine analysis, blood sugar, serum electrolytes, liver function tests, serum amylase, ECG, X-ray chest & abdomen, USG abdomen, upper GI endoscopy and urinary catecholamine assay were normal. Subsequently, in view of accelerated hypertension, tachycardia possibility of porphyria was suspected and diagnosis confirmed by qualitative test for porphobilogen. She was treated with glucose administration, antispasmodics and chlorpromazine and made rapid and full recovery.

Case 2: HT (HN:215841), a 17 years old girl was admitted with acute neurological illness manifesting with fever, higher function changes and recurrent seizures. At admission she was febrile (Temp-99.6°F) with pulse-102/min and BP 160/110 mm Hg. She was responding to painful stimuli by localized. No lateralizing neurological deficits detected. Deep tendon reflexes were symmetrical with planter response mute on both sides. Pupils were equal and reacting to light normally. Fundus examination was normal. No neck rigidity was detected. Rest systems were normal. She was diagnosed as a case of meningocencephalitis of probable viral etiology with herpes simplex encephalitis as the likely possibility. Investigations done i.e. Blood: Hb 14.8 gm/dl, TLC 14800/cmm, DLC P 89%, L19%; Serum electrolytes (mmol/L) : Na+ 104 , K+ 3.6; CT Scan revealed small pinched ventriciles with a hydropodense non enhancing lesion in right temporal region; CSF examination revealed normal biochemistry. Cell: WBC 6/cmm and RBC 1200/cmm. Blood sugar, renal parameters, Liver function tests, ECG and X-Ray chest were normal. EEG showed no periodic discharges. Her seizures were controlled with IV diazepam and diphenyl hydantoin (DPH). She was treated with acyclovir, IV antibiotics, antihypertensive, antipyretics, IV alimentation and sodium supplementation. Maintenance dose of DPH continued. On day 7th of her admission, she was afebrile, had normal orientation and was taking fluid orally. She was extubated, transferred out of ICU and ambulation started. A day later, she was noted to be passing high colored urine which changed color on exposure to sun suggesting porphyria (confirmed by qualitative test for porphobilinogen). DPH was withdrawn and replaced with gabapantine and high dextrose alimentation started.

However, on day 11th she developed abnormal behavior, tachycardia, and tachypnoea with spO2 falling to 70% and rapidly progressing areflexic quadriparesis, bulbar and respiratory paralysis. She was put on ventilator and DVT prophylaxis started. Other treatment continued. On day 15th, fluctuating BP (150-160/90-110 to 70-80/45-50 mms of Hg) and tachycardia, requiring frequent modification of antihypertensive drugs doses. Her BP stabilized but tachycardia persisted. There was no improvement in neuropathy. Two days later, she suffered a generalized seizure (controlled with diazepam) and developed acute abdominal pain and features of paralytic ileus. QTC prolongation was noted on day 18th. She continued to have frequent cardiac arrhythmia. Inspite of continued intensive care, her illness terminated fatally when she had cardiac...
arrest following ventricular tachycardia.

DISCUSSION

Due to varied symptomatology AIP is often misdiagnosed as is evident by the cases referred above. While case 1 and 2 had the referral diagnosis of acute pancreatitis, the third case developed symptoms of acute porphyria while recovering from encephalitic illness. In addition in case -2 an alternative diagnosis s of pheochromocytoma and polyarteritis nodosa was considered and excluded. Clinical presentation in the cases presented was dominated by dysautonomia i.e. persistent tachycardia and mild hypertension (Case-1), accelerated hypertension and tachycardia (Case-2) and persistent tachycardia and hypertension followed by episodes of hyper and hypotension, tachycardia and bradycardia and cardiac arrhythmias respectively.

In AIP, neurotoxic d-aminolaevulinic acid and porphobilinogen accumulates due to induction of haem synthesis and their subsequent excretion from liver in large amount. The areas lacking barrier protection i.e. peripheral and autonomic nervous system and hypothalamic and limbic areas of brain are affected. A high incidence of symptoms attributable to autonomic neuropathy also suggest high vulnerability of autonomic nerves to abnormal accumulation of porphyrin metabolites. Peripheral neuropathy in porphyria is mainly an axonal neuropathy which is followed by “dying back” degeneration. Tachycardia is attributable to unopposed sympathetic activity due to vagal damage and hypertension is the result of deafferentation of baroreceptors due to damage to vagus and glossopharyngeal nerves, Rana et al., (1981) studied five cases of porphyria when they were recovering. In addition to acute abdominal pain, the common combination was of hypertension, tachycardia and abnormal sweating patterns and diarrhea was noted 3 cases. Postural hypotension was noted in 2 patients with one having postural symptoms. Non invasive tests for ANS functions revealed abnormal valsalva ratio (1.01+/-0.05) and E: I ratio (1.00 +/- 0.16) as compared to 1.41+/-0.35 and 1.25+/-0.30 in control groups. Cold immersion test and response to atropine and propranolol was abnormal in 4, 4 and 3 patients respectively. Other studies also noted dysautonomia using similar noninvasive tests. Focal vascular damage with reversible cerebral edema has been reported in patients presenting as encephalopathy. These patients also show high catecholamines in urine due to inhibition of its uptake by platelets by raised d-ALA and porphobilinogen that may be confused with pheochromocytoma. However, urine catecholamines were normal in Case -2 were this diagnosis was considered.

Motor neuropathy and dysautonomia of porphyria resemble AIDP. Loss of reflexes corresponding to muscle weakness, severe abdominal pain, neuropsychiatric symptoms and occurrence of seizures differentiate porphyria from AIDP. Early proximal muscles involvement differentiates this from other toxins induced dying back neuropathy. Also, combination of focal CNS involvement and peripheral neuropathy is unusual for other neuropathies. Heavy metal poisoning (i.e. arsenic & lead and thallium) presents with abdominal pain, seizures and encephalopathy and simulates AIP making screening for heavy metals a mandatory requirement.

To avoid misdiagnosis and potential fatality in acute porphyria, suspected patients should be investigated to confirm the diagnosis. Where qualitative and quantitative estimation of urinary PBG is not possible a simple test of observing urine color after exposure to sunlight is enough as was done in present cases. AIP is confirmed by marked rise in urinary porphobilinogen (uroporphyrin 1 & III more than coproporphyrin 1 & III), fecal coproporphyrins normal or slightly raised and coproporphyrin III/I ratio less than 2. If all three are normal porphyria is excluded. If only erythrocyte PBG deaminase activity is abnormal, analysis of HMBS gene is needed. If positive it suggests, AIP in remission or in latent stage when family history is positive. Specific treatment of AIP administration of haem preparation (i.e. Hematin argenate or hematin) It should be administered at an earlist.The dose is 3 mg/kg/day for 3-4 consecutive days Hematin was not available for treatment. Treatment in case 1 and 2 case consisted of parenteral high carbohydrate intake, antispasmodics, diazepam and atenolol. While in first case complete remission was obtained, in case-2 abdominal pain was relieved on day 5th day but she developed altered sensorium leading to deep coma. SIADH was suspected and confirmed as the cause of coma. It responded to fluid restriction. In Case-3, case diagnosis of encephalitis was entertained as fever is not a feature of acute porphyria. It was supported by improvement in neurological status inspite of administration of DPH. Later, she developed severe degree of dysautonomia with changing profile. It is postulated that she had AIP in latent stage when admitted which manifested with continued administration of DPH.

Complete recovery in second case inspite of prolong coma emphasize the importance of early recognition and prompt treatment of SIADH, an adverse prognostic factor. Development of acute neuropathy with bulbar and respiratory paralysis requiring mechanical ventilation and severe fluctuating dysautonomia are associated with high mortality. Dysautonomia manifesting with fluctuating BP, episodic bradycardia followed by tachycardia and, ventricular fibrillation was the cause of fatal termination in third case.

As development of complication can not be predicted, all AIP cases should be treated in ICU for constant monitoring for dysautonomia and for institution of appropriate remedial measures similar to that applied for AIDP cases, where similar dysautonomia is frequently noted. Dysautonomia persist even in latent phase suggesting...
Case Note

subclinical neuropathy in porphyria. Continued monitoring of autonomic functions should be continued even when patients have recovered from acute episode.

REFERENCES


17. Rana PVS, Suri ML, Sahai B. A study of autonomic functions in neurological disorders. AFRC Project No 1030/78.


