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Beer Potomania: A Challenging Case of Hyponatremia

Hind Rafeia, Raza Yunus, Parvinder Khurana

Abstract

Beer potomania is a syndrome of hyponatremia associated with excessive beer drinking. Little or no salt content of beer results in marked reduction in the solute load to the kidney. This leads to impaired water clearance and dilutional hyponatremia. A 66-year-old man with history of alcoholism and alcoholic cardiomyopathy presented to the emergency room with tremors of his upper and lower extremities. He had a significant history of alcohol consumption, usually drinking 4 - 5 cans of beer per night for the past 34 years. In addition, he had consumed a fifth of a vodka bottle the day before presentation. He had a pattern of often skipping meals though was compliant with both his diuretics medications: furosemide 40 mg once daily and spironolactone 25 mg daily. On physical exam, he was euvoelic. Neurological exam revealed resting tremors of both his hands. Labs were remarkable for plasma sodium of 122, brain natriuretic peptide of 474, serum osmolality of 268, urine osmolality of 223, and urine sodium of 20. Patient was assessed to have moderate euvoelic hypotonic hyponatremia. The combination of euvoelic hyponatremia with history of excessive beer drinking made beer potomania very likely. His urine osmolality and urine sodium, however, were higher than expected in beer potomania. These could be explained by the two diuretics that the patient was taking. Patient was managed with fluid restriction, appropriate nutritional and sodium intake and withholding of his diuretics. Plasma sodium slowly corrected to 130 over the course of 3 days. This case illustrates the condition beer potomania, an infrequent cause of hyponatremia. Findings in hyponatremia do not always point in one direction, especially with the concomitant use of diuretics. Recognition of beer potomania is critical as it is associated with serious neurologic sequelae that should be part of counseling against alcohol abuse.

Keywords: Beer potomania; Hyponatremia; Diuretics

Case Report

A 66-year-old man with history of alcoholism and alcoholic cardiomyopathy presented to the emergency room with tremors of his upper and lower extremities. History dated back to 6 days prior to presentation when he developed tremors of all his extremities and was unable to walk. He tripped at home but denied loss of consciousness or head trauma. He had a significant history of alcohol consumption, usually drinking 4 - 5 cans of beer per night for the past 34 years. In addition, he had consumed a fifth of a vodka bottle the day before presentation. He had a pattern of often skipping meals though was compliant with both his diuretics medications: furosemide 40 mg once daily and spironolactone 25 mg once daily. He was not on any antipsychotic or antidepressant drugs. His history was negative for seizures, confusion or somnolence. He denied diarrhea, vomiting, cold intolerance, shortness of breath, swelling, or excessive water intake. He denied cough, chest pain, hemoptysis, weight loss, fevers, or night sweats. On physical exam, he was afebrile, hypertensive to 165/75, with a heart rate of 67, respiratory rate of 18, and an oxygen saturation of 100%. He was awake, cooperative and in no acute distress. He was euvoelic with no jugular venous distention, lower extremity edema or pulmonary crackles. Neurological exam revealed resting tremors of both his hands with no asterixis.

His laboratory workup included a complete blood count that revealed a white blood count of 4.8 × 10^3/µL, hemoglobin of 10.8 g/dL, hematocrit of 32.1%, and a platelet count of 156 × 10^3/µL. His basic chemistries on admission showed hyponatremia with a sodium level down to 122 mmol/L, a potassium level of 5 mmol/L, chloride of 88 mmol/L, bicarbonate of 28 mmol/L, blood urea nitrogen of 8 mg/dL and a creatinine of 1 mg/dL. Blood glucose on admission was 114 mg/dL. Mineral panel showed calcium of 8.6 mg/dL, phosphate of 3.3 mg/dL...
and magnesium of 1.7 mg/dL. Liver function tests were positive for 2:1 ratio of AST to ALT with AST of 61 IU/L and ALT of 30 IU/L. Otherwise, total protein was 5.7 g/dL, albumin was 3.6 g/dL, alkaline phosphatase was 79 mU/mL, total bilirubin was 1.4 mg/dL, direct bilirubin was 0.2 mg/dL and INR was 1. Brain natriuretic peptide (BNP) was 474 pg/mL compared to 4,000 pg/mL on a previous admission for congestive heart failure exacerbation. Thyroid stimulating hormone (TSH) was 1.12 µIU/mL. Serum osmolality was 268 mOsm/kg H$_2$O while urine osmolality was 223 mOsm/kg H$_2$O with urine sodium of 20 mmol/L. A chest X-ray was also done and was normal.

Patient was admitted to the hospital for management of hyponatremia. Based on the history of excessive beer drinking with recent binge drinking as well as the physical exam and laboratory findings suggestive of hypotonic euvolumeic hyponatremia, patient was diagnosed with beer potomania especially that other known causes of hyponatremia were excluded. Management consisted of fluid restriction, adequate nutritional supplementation and withholding of the two diuretics. Intravenous fluids were not administered, as patient was not acutely symptomatic. Plasma sodium was frequently monitored. It slowly corrected up to 130 over the course of 3 days.

**Discussion**

Based on the low serum osmolality of 268 mOsm/kg H$_2$O, hyponatremia was categorized as hypotonic hyponatremia [4]. It was further considered moderate due to a level between 120 and 130 mmol/L [5]. The fact that he presented 6 days after the onset of symptoms and that he lacked major neurologic symptoms such as seizures, confusion, somnolence, etc. made chronic hyponatremia (> 48 h) more likely [6]. Congestive heart failure exacerbation as the etiology of hyponatremia was excluded in this case due to many factors. Patient lacked attributable signs and symptoms such as shortness of breath, orthopnea, volume overload, cardiopulmonary decompensation on chest X-ray, elevated BNP, etc. Furthermore, TSH was normal which excluded hypothyroidism. Psychogenic polydipsia was unlikely as patient denied excessive water intake and urine osmolality was not less than 100 mOsm/kg H$_2$O. History, physical exam and workup made SIADH also an unlikely explanation. Chest X-ray was negative for a pulmonary process. Malignancy was not a concern at that point as patient denied systemic symptoms of weight loss, fevers, night sweats and was up-to-date with age-specific screening. History was negative as well for intake of antipsychotics, antidepressants or other drugs that can lead to SIADH and no intracranial process was evident from history or physical exam.

The history of excessive beer drinking with recent binge drinking made beer potomania a very likely explanation of patient’s euvolumeic hyponatremia. The criteria used to diagnose beer potomania syndrome included a history of binge beer drinking and poor dietary intake along with decreased serum sodium levels in the absence of other known causes of hyponatremia [7, 8]. While beer potomania is usually characterized by
low urine sodium and urine osmolality [9], the urine osmolality and urine sodium were slightly higher than expected. These findings could be explained by the two diuretics patient was taking. Both furosemide and spironolactone can cause excretion of sodiu in urine, which leads to a higher urine sodium and osmolality [10]. In a review of 22 cases of beer potomania, Sanghvi et al showed that hyposthenuria on admission was not a consistent finding in beer drinkers with hyponatremia. They concluded that recognizing this finding prevents the erroneous dismissal of beer potomania [11]. Figure 1 illustrates the pathophysiolo of hyponatremia in this particular patient. Patient’s presentation could not be explained by excessive diuresis from furosemide and spironolactone alone due to many factors. Patient lacked symptoms of hypovolemia. Furthermore, urine sodium in diuretic-induced hyponatremia is usually higher than in our patient (usually above 40 mmol/L) with associated higher urine osmolality (above 450 mOsm/kg H\textsubscript{2}O) [12].

Beer potomania is a well-known cause of hyponatremia. It is mainly explained by water intake that exceeds excretory capacity [11]. Free water clearance is dependent on solute excretion and urinary diluting capacity [13]. The osmolar load to the kidney is caused both by urea generation from proteins as well as dietary sodium and potassium intake. Under normal conditions, typical osmole excretion is approximately 600 - 900 mOsm/day. While the maximum urinary dilution capability is 50 mOsm/L, a large amount of water (> 20 L) must be ingested before overwhelming the urinary dilution capacity, as seen is psychogenic polydipsia. The low solute content of alcohol as well as prevention of protein breakdown (urea generation) by the carbohydrate (high calorie) content of beer in conjunction with poor dietary supplementation of sodium and potassium intake as seen in many alcoholics, all together, leads to a decrease in solute excretion. Consequently, the ability to excrete free water becomes limited. Because the obligatory solute loss in a day is approximately 250 mOsm in these patients, with a urinary dilution capability of 50 mOsm/L, water intake greater than 5 L (or 14 cans of beer) results in hyponatremia [7]. As a result, antidiuretic hormone (ADH) levels are expected to be suppressed in patients with beer potomania due to excess of water [14]. The low ADH levels limit free-water reuptake in the collecting tubules of the kidney and lead to brisk diuresis when solute is presented. Water diuresis can produce large increases in serum sodium levels in a short period, which in turn predisposes to the negative consequences of rapid correction of hyponatremia. Sequelae include in extreme cases pontine and extrapontine myelinolysis known as osmotic demyelination syndrome (ODS) due to the osmotic stress on oligodendrocytes that occurs with rapid correction of hyponatremia [15]. Thus, management of hyponatremia in beer potomania might be difficult.

In their review of beer potomania cases, Sanghvi et al [11] recommended that treatment goals for patients presenting with beer potomania should be limited to a serum sodium increase of less than 10 mmol/L in the first 24 h and less than 18 mmol/L in the first 48 h. If the patient is asymptomatic, fluid restriction and patient’s monitoring is the safest approach. If the desired goal is exceeded, D5W infusion might be started to match urine output. The rate of infusion can be adjusted based on sodium levels monitored every 2 h.

Conclusion

Beer potomania is an infrequent cause of hyponatremia caused by excessive alcohol intake and poor diet. Findings in hyponatremia do not always point in one direction, especially with the concomitant use of diuretics. Thus, understanding of pathophysiology can be useful in interpreting lab results. Recognition of beer potomania as a separate entity of hyponatremia is critical, as management should be tailored to the specific pathophysiological state of this condition. If unrecognized, beer potomania can lead to serious neurologic sequelae. Patient education and alcohol-related counseling are essential to prevent recurrence.

Disclosure

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References


