Multiple Causes of Hyponatremia: A Case Report

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### Introduction

Hyponatremia, with a serum sodium level <135 nmol/L, is the most common electrolyte disorder, reported in approximately 1% of the general population, in up to 15–20% of all patients in urgent care facilities, and in up to 30% of all patients treated in intensive care units \cite{1}. There are several forms of hyponatremia, i.e., hypervolemic, hypovolemic, euvoletic, that manifest as redistributive and pseudo-hyponatremia. Hyponatremia can be acute or chronic and manifest with mild or more severe symptoms such as confusion, nausea, vomiting, cardiore-
spiratory distress, sleep disorders, seizures, and coma due to several factors, like diuretics, adrenal disorders, inappropriate antidiuresis, etc. [1, 2]. Clinicians frequently encounter such cases in their practice but rarely are they caused by multiple factors simultaneously [1]. Hence, we report a case with 4 causes of hyponatremia that occurred simultaneously.

**Case Report**

A 27-year-old man, conscious and oriented, was admitted to the Regional General Hospital complaining of symptoms such as nausea, vomiting, diarrhea, a swollen face and swollen extremities, and dark colored urine, which had developed over the previous 2 days. Laboratory tests performed upon admission showed leukocytosis (21.7 × 10^9/L) with a predominance of granulocytes, increased levels of urea and creatinine, and the presence of hypokalemia (Table 1). Biochemical urine tests showed microscopic hematuria and positive proteins, and analysis of arterial blood showed hyponatremia with a sodium level below 100 mmol/L (Table 1). Throat swabs and urine cultures were done and therapy was started, including fluid restriction and penicillin at 1.6 million units every 12 h, along with a proton pump inhibitor and metoclopramide.

The patient developed an acute symptomatic seizure and coma (Glasgow coma score 3) and was transported to the Emergency Center of Clinical Centre of Vojvodina, Novi Sad, Serbia. Upon admission to the Emergency Center, the patient was comatose (Glasgow coma score 3) and normotensive (140/70 mm Hg), with a respiratory rate of 30 breaths/min and a heart rate of 90 beats/min. A computed tomography scan of the abdomen, abdominal ultrasonography, and a chest X-ray, performed upon admission, did not show pathological changes. Sodium levels still indicated hyponatremia (103 mmol/L) and a low plasma osmolality (i.e., 247 mosm/kg; normal range 270–300 mosm/kg), while the urine osmolality was relatively high (i.e., 286 mosm/kg; normal range 30–1,300 mosm/kg). There were no signs of hypervolemia at the time of admission, and hence therapy with hypertonic sodium, fluid restriction, correction of potassium, and antibiotics was introduced. Sodium levels in blood were gradually normalized (by not more than 0.5 mmol/L/h) to avoid central pontine myelinosis and the patient gradually regained consciousness over the following 3 days (Table 1). Hypertension was treated with angiotensin-converting enzyme inhibitors given on several occasions. Initially high levels of urea and creatinine also normalized and the patient entered the recovery phase with diuresis of 4,500 mL/day (Table 1).

On the fourth day, the patient was moved to the Department of Endocrinology, Diabetes and Metabolic Disorders of for additional investigations regarding to the etiology of hyponatremia. The patient regained consciousness and a thorough examination showed that he had had a sore throat and fatigue 10 days before admission to the hospital. Although he was feeling unwell, he had gone to a party with his friends at the end of that week where he drank 5,000 mL of beer, 400–500 mL of liquor, and 2,000 mL of energy drink (containing 25 mL/100 mL of caffeine and taurine, vitamins, sugar, citric acid, and caramel) during a 6-h period. The morning after the party, he noticed that his entire face, neck, and extremities were swollen. Afterwards, he experienced nausea, repeated vomiting, diarrhea, and dark colored urine along with excessive weakness and fatigue which continued to progress. Two days after the party, the patient contacted his doctor and was hos-

<table>
<thead>
<tr>
<th></th>
<th>General hospital</th>
<th>Emergency center</th>
<th>Day 1</th>
<th>Day 2</th>
<th>Day 3</th>
<th>Day 4</th>
<th>Day 5</th>
<th>Day 6</th>
<th>Day 11</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium, mmol/L (135–148)</td>
<td>&lt;100</td>
<td>103</td>
<td>109</td>
<td>122</td>
<td>138</td>
<td>140</td>
<td>140</td>
<td>145</td>
<td></td>
</tr>
<tr>
<td>Potassium, mmol/L (3.5–5.5)</td>
<td>2.8</td>
<td>4.1</td>
<td>4.2</td>
<td>4.2</td>
<td>3.6</td>
<td>4.0</td>
<td>4.3</td>
<td>4.3</td>
<td></td>
</tr>
<tr>
<td>pH (7.35–7.45)</td>
<td>7.37</td>
<td>7.51</td>
<td>/</td>
<td>/</td>
<td>7.42</td>
<td>/</td>
<td>/</td>
<td>/</td>
<td></td>
</tr>
<tr>
<td>Lactate, mmol/L (0.5–1.6)</td>
<td>11.4</td>
<td>0.99</td>
<td>1.88</td>
<td>1.04</td>
<td>/</td>
<td>/</td>
<td>/</td>
<td>/</td>
<td></td>
</tr>
<tr>
<td>C-reactive protein, mg/L (0.0–5.0)</td>
<td>/</td>
<td>15.1</td>
<td>/</td>
<td>/</td>
<td>12.4</td>
<td>/</td>
<td>/</td>
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<td></td>
</tr>
<tr>
<td>BUN, mmol/L (2.5–7.5)</td>
<td>14.5</td>
<td>14.8</td>
<td>14.0</td>
<td>10.9</td>
<td>6.8</td>
<td>2.9</td>
<td>3.2</td>
<td>3.8</td>
<td></td>
</tr>
<tr>
<td>Creatinine, μmol/L (50–115)</td>
<td>286</td>
<td>224</td>
<td>177</td>
<td>130</td>
<td>106</td>
<td>90</td>
<td>88</td>
<td>96</td>
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<tr>
<td>Urine output, mL/24 h</td>
<td>/</td>
<td>200 mL/6 h</td>
<td>2,500</td>
<td>4,500</td>
<td>4,000</td>
<td>3,000</td>
<td>2,500</td>
<td>1,850</td>
<td></td>
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<tr>
<td>Red blood cells in urine</td>
<td>mass</td>
<td>mass</td>
<td>/</td>
<td>mass</td>
<td>/</td>
<td>mass</td>
<td>/</td>
<td>/</td>
<td></td>
</tr>
<tr>
<td>Proteins in urine</td>
<td>++</td>
<td>++</td>
<td>/</td>
<td>++</td>
<td>/</td>
<td>+</td>
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Normal ranges are in parentheses. BUN, blood urea nitrogen.
pitalized at the General Hospital Sremska Mitrovica. A throat swab culture test that was done at the Regional General Hospital revealed *Streptococcus beta haemolyticus* of group A and a negative anti-streptolysin O titer. A repeated throat swab was negative and a seroconversion of a previously negative anti-streptolysin O titer (anti-streptolysin O titer 400 IU) was seen together with a decrease in C3 complement components levels. Urea and creatinine levels and the daily urine volume started to normalize but proteinuria (with a sodium level of 1,800 mg/day) with microscopic hematuria remained (Table 1). The estimated glomerular filtration rate was 102 mL/min/1.73 m². Mild normocytic and normochromic anemia was also observed.

Based on these data, the diagnosis of acute poststreptococcal glomerulonephritis was established. The severe acute hyponatremia followed by coma was caused by excessive simultaneous consumption of fluids, alcohol and caffeine. The normal levels of glycemia, thyroid hormones (free T3, free T4), thyroid-stimulating hormone, cortisol and adrenocorticotropic hormone led to exclusion of the possibility that hypocortisolism and hypothyroidism caused hyponatremia. After 14 days the patient was discharged from the hospital and advised to take care of his hygiene and diet and to abstain from alcohol and energy drinks. One month later, laboratory findings still indicated the presence of proteinuria (sodium level 864 mg/day).

**Discussion**

Adequate treatment of hyponatremia in our patient led to a complete recovery without consequences. The literature indicates that severe hyponatremia can lead to different outcomes, from complete recovery to permanent brain damage and death [1]. Gradually increasing the concentration of sodium, as was done in this case, results in a gradual but complete recovery without any complications due to too rapid an increase in serum sodium [1]. The first cause of hyponatremia in this case was acute poststreptococcal glomerulonephritis and the patient’s health was endangered by 3 other potential causes of hyponatremia, including consumption of a large amount of liquids poor in sodium (5,000 mL of beer), intake of caffeine, and acute alcohol intoxication followed by subsequent vomiting. The so-called, “beer potomania” is known to be the cause of hyponatremia [3]. Acute alcohol intoxication can also be associated with pronounced hyponatremia [4, 5]. Considering the facts that the patient’s weight was about 95 kg and that he had been drinking alcohol for 6 hours, the modified Widmark formula estimates that the average alcohol level in blood was about 0.3544 g/dL [6]. Apart from the immediate effect, this amount of alcohol caused the patient’s frequent vomiting and worsened his hyponatremic state, compelling him fortunately to contact his doctor. [5] Consumption of energy drinks (2,000 mL) led to an excessive caffeine intake. The caffeine content declared on the labels of energy drinks, (i.e., 25 mg/100 mL) confirms that this drink is relatively safe when consumed in small amounts. By drinking 2,000 mL of this drink, the patient consumed 500 mg of caffeine. The diuretic and natriuretic effects of caffeine at the dose consumed by the patient could solely lead to hyponatremia. The European Food Safety Authority considers consumption of 200 mg of caffeine to be safe and has stated that certain side effects, including hyponatremia, can be caused by higher doses [7]. Pronounced hyponatremia is rarely expected in acute post-streptococcal glomerulonephritis although there are reports in the literature that acute poststreptococcal glomerulonephritis can lead to hyponatremia [8]. Although a simultaneous occurrence of 4 causes of hyponatremia in 1 patient could be considered very unusual, the fact that this patient with the stated sodium levels (Na⁺ <100 mmol/L) was conscious, oriented, communicative, and without neurological disorders should be taken as an exception. Coma and acute symptomatic seizure occurred a few hours after the level of sodium was confirmed to be below 100 mmol/L. A possible explanation for the maintained state of consciousness at these sodium levels over a long period of time is chronic hyponatremia, though this would not be option in a previously healthy person as in this case. The amount of taurine that the patient ingested by drinking energy drinks might be an alternative explanation. Taurine is a cerebral membrane stabilizer and the transport of taurine to brain cells depends on sodium and chloride [9]. It has been shown that there is a sodium/taurine symporter in the guinea pig heart which increases the efflux of both substances to the heart cells when the concentration of any of these 2 substances is increased. If a similar mechanism exists in humans, it is possible that large amounts of taurine led to maintenance of the level of sodium in the cells of the patient, thus delaying for some time the development of severe symptoms of hyponatremia, such as convulsions and coma [10].

**Conclusion**

This patient had 4 simultaneous causes of hyponatremia. The late onset of coma could have been due to the large amount of taurine, which resulted in higher levels of intracellular sodium.

**Disclosure Statement**

None.
References