Cerebral salt wasting syndrome

Harshal Dholke, Ann Campos, C Naresh K. Reddy, Manas K. Panigrahi

Abstract

Traumatic brain injury (TBI) is on the rise, especially in today’s fast-paced world. TBI requires not only neurosurgical expertise but also neurointensivist involvement for a better outcome. Disturbances of sodium balance are common in patients with brain injury, as the central nervous system plays a major role in sodium regulation. Hyponatraemia, defined as serum sodium <135 meq/L is commonly seen and is especially deleterious as it can contribute to cerebral oedema in these patients. Syndrome of inappropriate antidiuretic hormone secretion (SIADH), is the most well-known cause of hyponatraemia in this subset of patients. Cerebral Salt Wasting Syndrome (CSWS), leading to renal sodium loss is an important cause of hyponatraemia in patients with TBI. Although incompletely studied, decreased renal sympathetic responses and cerebral natriuretic factors play a role in the pathogenesis of CSWS. Maintaining a positive sodium balance and adequate hydration can help in the treatment. It is important to differentiate between SIADH and CSWS when trying to ascertain a case for patients with acute brain injury, as the treatment of the two are diametrically opposite.

Key words: Brain natriuretic peptide, cerebral salt wasting, conivaptan, fludrocortisone, hypertonic saline, hyponatraemia

INTRODUCTION

Sodium is the most important osmotically active solute in the extracellular fluid. It is the major determinant of serum osmolality, which in turn plays a major role in the regulation of body water. Increased serum osmolality, triggers the release of anti-diuretic hormone (ADH) from the posterior pituitary. Hypovolaemia and hypotension results in baroreceptor stimulation which reflexly causes ADH release.[1,2]

Sodium balance in the body is maintained via regulation of its renal excretion which is affected by:

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Glomerular filtration rate

About 70% of filtered sodium is reabsorbed in the proximal tubules with <3% being excreted. Any fall in glomerular filtration rate (GFR) would, therefore, mean less filtration and excretion of sodium and vice versa.

Renin-angiotensin-aldosterone system

Sympathetic stimulation decreases in mean arterial pressure or decreases in distal tubular sodium levels, all activate renin-angiotensin-aldosterone system (RAAS) which results in sodium reabsorption secondary to aldosterone release.[1]

Natriuretic peptides (atrial natriuretic peptide and brain natriuretic peptide)

These are produced in the atria and brain and cause reduction of sympathetic outflow from the brainstem as well as induce natriuresis by increasing the GFR, and by inhibiting renin and aldosterone release.[2]
In traumatic brain injury (TBI), hyponatraemia can occur due to a variety of causes, such as syndrome of inappropriate antidiuretic hormone secretion (SIADH), CSWS, Anterior hypopituitarism, and drugs such as oxcabazepine used for seizure prophylaxis. The incidence of hyponatraemia in head injury patients commonly linked to CSWS and SIADH is 5–10%.[4,5] While SIADH is the cause of hyponatraemia in the majority of cases, in a small subset of patients, the diagnosis of CSWS is often missed as it appears to be very similar to SIADH and is therefore often confused with it. This can have disastrous consequences as the treatment of the two conditions is diametrically opposite. In SIADH, despite a low serum osmolality, there is an inappropriate secretion of ADH in response to hyponatraemia, leading to water retention and hypervolaemia.[3] CSWS mimics SIADH except for the fact that salt wasting is the primary defect causing volume depletion.[6,7]

First described by Peters et al. in 1950, cerebral salt wasting (CSW) is a clinical condition characterised by renal loss of sodium causing dehydration and hyponatraemia in patients with intracranial neurological disorders.[6] In 1953, Leaf et al.[7] demonstrated that exogenous administration of the ADH (vasopressin) resulted in hyponatraemia, water retention and weight gain. The increase in the intravascular volume resulted in a decrease in sodium and chloride levels. This was not ‘salt wasting’, but was a physiologic response to an expanded intravascular volume. Four years later, Schwartz et al.[8] published their landmark paper on SIADH. A subsequent paper from the group at Yale attributed hyponatraemia in neurologic disease to SIADH.[7] For over 20 years, the term CSWS virtually vanished from literature. In 1981, Nelson et al.[9] studied hyponatraemia in neurosurgical patients, primarily subarachnoid haemorrhage, and found that isotopically measured blood volumes were contracted; he attributed this finding to CSWS [Figures 1 and 2].

**PATHOPHYSIOLOGY OF CEREBRAL SALT WASTING SYNDROME**

In TBI, there can be disruption of hypothalamo-renal pathways,[3] imbalance of sympathetic output with decreased renal sympathetic activity,[10] and also possibly direct injury to the anterior and posterior pituitary, all of which can play a role in the pathogenesis of hyponatraemia in these patients. This can disrupt the cerebral influence on renal salt and water balance, and therefore, disturb the kidneys ability to handle sodium properly.[6]

It is now believed that natriuretic factors such as an atrial natriuretic peptide, brain natriuretic peptide (BNP), C-type natriuretic peptide, and possibly dendroaspis natriuretic peptide are secreted by the injured brain and may play a role in CSWS. Of all these factors BNP might be the main factor in CSWS.[11] These peptides have potent effects on cardiovascular homeostasis by dampening the sympathetic response thereby altering the vascular tone and causing dilatation of arteries and veins.[11] Natriuretic peptides also induce sodium loss (natriuresis) by inhibiting renin release from the renal juxtaglomerular cells and preventing aldosterone release from the adrenals thus antagonising the RAAS.[12] This effect on the afferent tubules of nephrons leads to, dilatation of the afferent arteriole resulting in increased filtration of water and sodium through the glomerulus. These molecules also have renal natriuretic and diuretic effect by inhibiting the angiotensin-induced sodium reabsorption from collecting ducts and antagonising the action of vasopressin at the collecting duct, respectively.[12]

Local production of natriuretic peptides within the adrenal medulla has been demonstrated, which, might have paracrine inhibitory effects on mineralocorticoid synthesis.[13] This paracrine mechanism might explain why, in patients with CSWS, aldosterone and renin levels fail to rise despite the presence of hypovolaemia.

Other mechanisms suggest that downregulation of renal sodium transporters due to extracellular volume expansion and the adrenergic surge that occurs in the early phase of brain injury might cause pressure natriuresis.[14,15]

**CLINICAL FEATURES AND DIAGNOSIS**

In clinical practice, it is important to distinguish CSWS from SIADH as they share several diagnostic criteria. The following laboratory studies may be indicated in patients with cerebral salt-wasting syndrome:[13]

**Serum sodium concentration**

Patients with untreated CSWS are often hyponatremic and signs and symptoms may vary according to the severity as shown in Table 1.[16] As the decline in serum sodium concentration reduces serum osmolality, a tonicity gradient develops across the blood-brain barrier that causes cerebral oedema. Symptoms include lethargy, agitation, headache, altered consciousness, seizures and coma.[17,18]

**Serum osmolality**

Normal serum osmolality is 285–295 mosmosm/L. This is found to be decreased in SIADH but is either normal or decreased in CSWS. If the measured serum osmolality exceeds twice the serum sodium concentration and azotaemia is not present, hyperglycaemia or mannitol should be suspected as the cause of hyponatraemia.[15,18]
Urinary output
Urine appears relatively dilute and the flow rate is often high in CSWS. It is concentrated with a low flow rate in SIADH. However, in CSWS, despite its apparently diluted appearance, urine osmolality is high due to increased sodium loss.

Fluid balance
The differentiation of SIADH from CSWS depends on an accurate estimation of intravascular volume. Unfortunately, no single physical finding can accurately and reproducibly measure effective circulating volume. Commonly used signs of hypovolaemia include orthostatic tachycardia or hypotension, increased capillary refill time, increased skin turgor, dry mucous membranes and a sunken anterior fontanel. These signs usually appear only when the degree of dehydration is moderate to severe. Central venous pressure may be an unreliable determinant of extracellular volume.\textsuperscript{[15]}

In SIADH, there is generally euvoalaemia or hypervolaemia. As opposed to this the important finding in CSWS is volume depletion. The daily sodium excretion is also more than the intake, and the overall sodium balance is negative in CSWS. Therefore, a daily clinical examination for signs of hypovolaemia as well as a daily intake and output charting should be done, which will reveal an overall negative balance.\textsuperscript{[15]} Sometimes, hypovolaemia has been identified in patients who fulfill all other diagnostic criteria for SIADH. This occurs because the volume depletion of CSWS causes a secondary rise in ADH. However, under such conditions, the correct diagnosis is CSWS rather than SIADH.\textsuperscript{[19]}

Fractional excretion of uric acid
This is defined as the percentage of urate filtered by glomeruli that is excreted in urine. It is calculated by dividing the product of (urinary uric acid [mg/mL] × serum creatinine [mg/mL]) by the product of (serum uric acid [mg/mL] × urinary creatinine [mg/mL]) and multiplying the result by 100. Normal values are <10\%.\textsuperscript{[20]}

Patients with either CSWS or SIADH can have hypouricaemia and elevated fractional excretion of uric acid (FEUA). However, after correction of hyponatraemia, the hypouricaemia and elevated FEUA may normalise in SIADH but persist in CSWS (renal salt wasting).\textsuperscript{[18]}

Fractional excretion of phosphate
This should be determined when evaluating patients with hyponatraemia and hypouricaemia. Elevated fractional excretion of phosphate >20% suggests cerebral salt-wasting syndrome as opposed to SIADH where it is <10\% [Table 2].\textsuperscript{[20,21]}

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**Table 1: Clinical presentation of hyponatraemia**

<table>
<thead>
<tr>
<th>Plasma concentration of Na in (mmol/L)</th>
<th>Signs/symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;125</td>
<td>Asymptomatic</td>
</tr>
<tr>
<td>120-125</td>
<td>Nausea, malaise, vomiting</td>
</tr>
<tr>
<td>120-110</td>
<td>Muscle cramps, weakness, confusion, agitation, delirium, lethargy and seizures</td>
</tr>
<tr>
<td>&lt;110</td>
<td>Seizures, coma, permanent brain damage, respiratory arrest</td>
</tr>
</tbody>
</table>

**Table 2: Clinical and biochemical features of the syndrome of inappropriate antidiuretic hormone secretion and the cerebral salt wasting syndrome**

<table>
<thead>
<tr>
<th>Biochemical marker</th>
<th>SIADH</th>
<th>CSWS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intravascular volume</td>
<td>Normal to high</td>
<td>Low</td>
</tr>
<tr>
<td>Serum sodium</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>Urinary sodium level</td>
<td>High</td>
<td>Very high</td>
</tr>
<tr>
<td>Vasopressin level</td>
<td>High</td>
<td>Low</td>
</tr>
<tr>
<td>Urine output</td>
<td>Normal to low</td>
<td>High</td>
</tr>
<tr>
<td>Serum uric acid level</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>Initial fractional excretion of urate</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Fractional excretion of urate after correction of hyponatraemia</td>
<td>Normal</td>
<td>High</td>
</tr>
<tr>
<td>Urinary osmolality</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Serum osmolality</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td>BUN/creatinine level</td>
<td>Low to normal</td>
<td>High</td>
</tr>
<tr>
<td>Serum potassium level</td>
<td>Normal</td>
<td>Normal to high</td>
</tr>
<tr>
<td>Central venous pressure</td>
<td>Normal to high</td>
<td>Low</td>
</tr>
<tr>
<td>Pulmonary capillary wedge pressure</td>
<td>Normal to high</td>
<td>Low</td>
</tr>
<tr>
<td>Brain natriuretic peptide level</td>
<td>Normal</td>
<td>High</td>
</tr>
<tr>
<td>Fractional phosphate excretion (%)</td>
<td>&lt;10</td>
<td>&gt;20</td>
</tr>
</tbody>
</table>

BUN=Blood urea nitrogen, SIADH=Syndrome of inappropriate antidiuretic hormone secretion, CSWS=Cerebral salt wasting syndrome

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**TREATMENT**

The main-stay of management of CSWS is the replacement of water and sodium which is lost due to diuresis and natriuresis, whereas, in SIADH, free water has to be restricted.\textsuperscript{[22]}
Once a diagnosis of CSWS is made, efforts should be made to address hypovolaemia first. This can be done with the use of crystalloids like 0.9% normal saline. This treatment holds true for those patients with mild hyponatraemia. By doing this, both hypovolaemia and hyponatraemia can be addressed.[23,24] When the patient is severely hyponatraemic and hypovolemic he will require aggressive resuscitation to first become euvoletic, followed by the correction of hyponatraemia, with the use of 3% saline which should be administered via a central line. With the use of 3% saline, the sodium correction should not exceed 12 meq/L for every 24 h. This is necessary, to avoid complications such as central pontine myelinolysis, metabolic acidosis, volume overload and pulmonary oedema.[25,26]

Some clinicians have found it useful to the use of mineralocorticoids in CSWS. Fludrocortisone is one such drug, which promotes the increased reabsorption of sodium and the loss of potassium by the renal distal tubules. Secondary effects such as hypokalaemia, pulmonary oedema and hypertension may occur with prolonged use. Apart from this, the steroid base can cause hyperglycaemia warranting the periodic monitoring of serum potassium and blood sugars. Hence, it’s use is only indicated when salt and fluid replacement are unable to correct the hyponatraemia.[27,28]

Diuretics and fluid restriction are the main-stays of treatment in SIADH. However, one of the newer drugs recently approved by the United States-Food and Drug Administration is worth a mention, Conivaptan, is a non-selective antagonist at V1 and V2 vasopressin receptors. It antagonises the action of vasopressin at the collecting ducts causing electrolyte-free water excretion, thereby raising serum sodium in patients with SIADH.[29,30] Most recently, Ghali et al. published results from their randomised, double-blind, placebo-controlled trial conducted across 21 cities in the United States, Canada and Israel, involving the efficacy of oral conivaptan in the treatment of patients with euvoletic and hypervolaemic hyponatraemia.[31] Based on currently available studies, conivaptan appears to be effective in inducing aquareesis to correct hyponatraemia in both euvoletic and hypervolaemic hyponatreemic patients. Although conivaptan has been shown to be an effective aquaretic with short-term use, this is not without limitations. Adverse effects reported with short-term use are typically minimal, but may include serious effects such as hypokalaemia, orthostatic hypotension, and unexpectedly rapid serum sodium correction. Careful patient selection, avoidance of combined use with conventional diuretics, and close monitoring may reduce complication rates with the use of conivaptan.[31,32] When used in CSWS, conivaptan can cause a negative fluid balance and further worsen the situation. This novel treatment for SIADH also backs the need for accurate differentiation between CSWS and SIADH before the start of hyponatraemia correction in cerebral injuries.[33,34] Conivaptan should not be administered to patients in whom CSWS or a high likelihood of cerebral vasospasm is suspected.[34,35]

PREVENTION OF HYPONATRAEMIA IN TRAUMATIC BRAIN INJURY

In TBI, the maintenance of intracranial pressure (ICP) is pivotal and we need to strongly address the changes in serum osmolality and serum sodium levels. There are ample data which suggests that maintenance of slight hypernatraemia is associated with a reduced increase in ICP.[6] This can be very well achieved with the continuous infusion of hypertonic saline (3% NaCl) and is found to be well tolerated in these patients.[26]
A recent study at the University of California, Los Angeles used an aggressive sodium correction treatment regimen as a component of their TBI protocol. The mean target goal was a serum sodium level of 138 mmol/L or higher if ICP was 15–20 mm Hg. This was achieved with the use of 3% NaCl to control serum sodium levels and prevent hyponatraemia. Using this protocol, there was a 12% incidence of serum sodium levels of <137 mmol/L and a 1% incidence rate of serum sodium levels of <132 mmol/L. There were no documented cases of central pontine myelinolysis using this protocol.[9]

Treatment protocol suggested for sodium regulation in TBI[8]
• Measure serum sodium level twice daily for initial 96 h after TBI
• Establish serum sodium goal of ≥138 mmol/L
• Start continuous infusion of 3% NaCl at 25 mL/h for Na of ≤138 mmol/L
• Maintain 3% NaCl at 15–25 mL/h if ICP is 15–20 mm Hg
• Add fludrocortisone 0.1 mg bid (oral) if Na is ≤138 mmol/L
• Increase 3% NaCl to around 50 mL/h to achieve Na of >145–155 mmol/L if ICP is >20 mm Hg despite the use of cerebrospinal fluid drainage using ventriculostomy.

CONCLUSION
Hyponatraemia can complicate the clinical outcome in TBI. CSWS is a syndrome of hypovolemic hyponatraemia caused by renal natriuresis and diuresis. Brain natriuretic peptide, secreted by the injured brain plays a crucial role in the pathogenesis of CSW. Making a distinction between SIADH and CSWS is important due to different treatment required for the two conditions. The maintenance of high normal levels of serum sodium in patient with TBI may help limit increases in ICP as well as avoid the detrimental effects of hyponatraemia due to CSWS or SIADH in these patients.

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Conflicts of interest
There are no conflicts of interest.

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