Challenges of Diagnosing Hyponatremic Syndromes in Pulmonary and Extra Pulmonary Tuberculosis


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Abstract

Introduction Pulmonary tuberculosis (PTB) is one of the rare pulmonary infections causing hyponatremia (serum sodium <135 mmol/L) and severe hyponatremia (serum sodium <125 mmol/L). Although the major cause of hyponatremia in TB patients is syndrome of inappropriate antidiuretic hormone (SIADH) secretion, cerebral salt wasting syndrome (CSWS) can occur and requires evidence of inappropriate urinary salt losses and reduced arterial blood volume. Adrenal insufficiency (AI) is rare in TB with scanty literature describing it. The two reported cases highlight three possible causes of severe symptomatic hyponatremia in TB: pleural effusion and disseminated TB, their treatment modalities, and the need to increase the index of suspicion to diagnose TB hyponatremia in children.

Case Report Case 1: a 10-year-old girl with TB pleural effusion who developed recurrent hyponatremia in the first few weeks of anti-TB treatment which was responsive to sodium correction. Case 2: an 8-year-old girl presenting to our facility with presumptive TB. She deteriorated over several months and progressed to disseminated TB with AI.

Discussion Early diagnosis and prompt and correct treatment of TB hyponatremia cannot be overemphasized, as AI, SIADH secretion, and CSWS, each require different therapeutic regimens, most especially AI on its own poses a huge clinical challenge.

Conclusion A high index of suspicion, with intensified case finding at all levels of care, is necessary to identify and manage children with TB hyponatremia because early diagnosis and prompt treatment is lifesaving.

Introduction Pulmonary tuberculosis (PTB) is one of the rare pulmonary infections causing hyponatremia.1,2 It is usually mild to moderate, asymptomatic, self-limiting, and reversible with anti-TB therapy.1,2 Hyponatremia is the reduction in serum sodium <135 mmol/L3,4 and severe hyponatremia occurs when levels are <125 mmol/L.3 It is one of the commonest electrolyte disturbances4,5 with prevalence of 1 to 4% in severe forms1,3 and 15 to 30% in nonsevere forms among hospitalized adults and children.1,4 Hyponatremia occurs in active TB in 11 to 51%1,6 of cases and thus its screening is of paramount importance.5 TB causes hyponatremia by affecting the adrenal glands, hypothalamus, pituitary gland, meninges, or lungs through...
adrenal insufficiency (AI), as well as inappropriate antidiuretic hormone (ADH) secretion. The main causes of hyponatremia are AI, syndrome of inappropriate ADH (SIADH), and cerebral salt wasting syndrome (CSWS).\textsuperscript{1,2} SIADH secretion should be considered in cases of hyponatremia with low serum osmolality, normal acid–base balance, urine osmolality $>100$ mOsm/L, and urine sodium $>40$ meq/L.\textsuperscript{3} It is poorly understood but believed to be due to hypoxia and reduced vascular volume.\textsuperscript{2,7} CSWS involves renal salt loss causing hyponatremia and extracellular fluid volume reduction, while SIADH is physiologically inappropriate secretion of ADH or increased renal sensitivity of ADH causing renal conservation of water and euvolemic hyponatremia.\textsuperscript{9} Conversely, increased serum ADH does not exclude CSWS as it may increase physiologically in response to hypovolemia.\textsuperscript{8} The different pathophysiological mechanisms between SIADH and CSW make early differentiation mandatory for correct treatment since both require diverse therapeutic regimens.\textsuperscript{9} Debates continue to trail CSWS, leaving it as an interesting academic controversy, and until more studies are done, CSWS should be considered a rare cause of hyponatremia compared with SIADH secretion.\textsuperscript{10}

AI is now rare in children\textsuperscript{11} with scanty literature describing it but TB remains its most common cause in the developing countries\textsuperscript{12}; it has been proposed that more than 90% of the adrenal gland should be affected by TB before insufficiency appears.\textsuperscript{13} Hyponatremia in AI is due to sodium wasting with secondary antidiuresis due to aldosterone deficiency\textsuperscript{10} and is associated with hyperkalemia and increased urinary potassium.\textsuperscript{2} The treatment of choice is steroid replacement.\textsuperscript{14,15} This report highlights the occurrence of severe symptomatic hyponatremia in TB pleural effusion, as well as disseminated TB with AI, which may be confused with TB treatment failure. Diagnosing AI is a clinical challenge that requires suspicion in a background of active TB.\textsuperscript{15} A high index of suspicion is therefore needed for both the diagnosis of childhood TB and early diagnosis with prompt treatment of hyponatremia because SIADH, CSWS, and AI, all require different therapeutic approaches.\textsuperscript{2,9}

**Case Reports**

**Case 1:** a 10-year-old girl with 4 months’ history of cough, fever, weight loss, 1 week’s vomiting, and 5 days’ diarrhea. Her father had TB 3 years earlier. Vital signs on admission were temperature, 39.8°C; respiratory rate (RR), 46 cycles/minute; pulse, 148/minute; blood pressure (BP), 110/70 mm/Hg; and SpO$_2$ 91% in room air. Her weight was 23 kg (10th percentile), while her height was 140 cm (>50th percentile). Salient findings were dyspnea, tachypnea, tachycardia, low blood pressure, and decreased respiratory effort. The radiograph showed a large left-sided pleural effusion with bilateral pitting pedal edema up to the vulva. Her weight was 27 kg (>50th percentile). Salient features were reduced tactile and vocal fremitus posteriorly with dull percussion notes upper zones anteriorly and stony dull percussion notes in the lower zones laterally and posteriorly, Pulse, 120/min; BP, 110/70 mm/Hg; disturbed abdomen with moderate ascites; 10-cm hepatomegaly; 4-cm splenomegaly; neck stiffness; and meningeal signs. A diagnosis of disseminated TB involving the lymph nodes, lungs, pleura, abdomen, and probably meninges was made. Chest X-ray showed right apical lobe opacities with pleural effusion. Abdominal USS revealed hepatosplenomegaly with matted mesenteric lymphadenitis, copious intraperitoneal echo-rich fluid collection, suggestive of abdominal TB. Full blood count (FBC) was white blood cell (WBC), 17.5 $\times$ 10$^3$/ul; packed cell volume (PCV), 28.9%; neutrophil, 93.8%; and lymphocyte, 3.1%. Erythrocyte sedimentation rate (ESR), 13 mm/h; red blood cell (RBC), Random blood sugar (RBS), 6.7 mmol/L; and Na, 127 mmol/L. She was to commence anti-TB drugs but developed convulsion and coma after 10 days of therapy with serum sodium of 114 mmol/L. This recurred in subsequent weeks with sodium levels ranging from 114 to 124 mmol/L but each episode responded to sodium correction with without corticosteroids and anti-TB drugs were continued. She stabilized after 10 weeks and was eventually discharged home. Follow-up was satisfactory.

**Case 2:** an 8-year-old girl presented with fever, cough, neck swellings of 10 months, weight loss of 6 months, and abdominal and leg swelling of 3 months’ duration. She lived with her grandmother who had TB. She had lymph node incisional biopsy and antibiotics before presentation, with temporary relief but deteriorated over several months with weight loss and increasing abdominal and leg swelling and eventually presented with generalized body swelling, nausea, vomiting, and weakness 10 months later. She was chronically ill looking, wasted, moderately pale, with bilateral cervical lymphadenopathy in anterior and posterior cervical, submandibular and axillary regions ranging from 2 cm $\times$ 3 cm to 4 cm $\times$ 6 cm and had an 8-cm incisional scar in the anterior part of the neck with bilateral pitting pedal edema up to the vulva. Her weight was 27 kg (>50th percentile). Salient features were reduced tactile and vocal fremitus posteriorly with dull percussion notes upper zones anteriorly and stony dull percussion notes in the lower zones laterally and posteriorly, Pulse, 120/min; BP, 110/70 mm/Hg; disturbed abdomen with moderate ascites; 10-cm hepatomegaly; 4-cm splenomegaly; neck stiffness; and meningeal signs. A diagnosis of disseminated TB involving the lymph nodes, lungs, pleura, abdomen, and probably meninges was made. Chest X-ray showed right apical lobe opacities with pleural effusion. Abdominal USS revealed hepatosplenomegaly with matted mesenteric lymphadenitis, copious intraperitoneal echo-rich fluid collection, suggestive of abdominal TB. Full blood count (FBC) was white blood cell (WBC), 17.5 $\times$ 10$^3$/ul; packed cell volume (PCV), 28.9%; neutrophil, 93.8%; and lymphocyte, 3.1%. Erythrocyte sedimentation rate (ESR), 13 mm/h; red blood cell (RBC), Random blood sugar (RBS), 6.7 mmol/L; and Na, 127 mmol/L. She was to commence anti-TB drugs but developed worsening body weakness, cold extremities, nausea, and vomiting. Radial pulse was not palpable but brachial pulse was 130/min and BP dropped to 90/60 mm Hg. SPO$_2$ was 63% in room air and oxygen, normal saline infusion, intravenous hydrocortisone of 200 mg were given.

![Fig. 1](image_url) Left-sided tuberculous pleural effusion of case 1.
She improved and stabilized for a few hours and then had another episode of persistent vomiting with cold extremities, weak thready pulse, gasping respiration, and a blood pressure of 80/50 mm Hg and couldn’t be resuscitated.

**Discussion**

Severe hyponatremia of 114 mmol/L in case 1, came as a surprise of 10-day anti-TB treatment and was announced by convulsion and coma. Severe hyponatremia occurs less commonly than mild forms, and so this case is a rarity. A rapid reduction of serum sodium to 110 to 120 mmol/L is usually associated with cerebral edema and possible brain herniation. She, however, responded to sodium correction, steroids, and anti-TB medications. Reports have shown treatment modalities of water restriction in SIADH secretion and salt and water infusion in Renal Salt Wasting (RSW), and it is important to differentiate CSWS from SIADH secretion since their treatments differ. Our patient had normal urine sodium, serum osmolarity and urine osmolarity. It is believed that urine sodium is raised in CSWS and reduced in SIADH secretion but a report showed the opposite. She developed tender hepatomegaly and renal parenchymal disease, raising the possibility of adrenal involvement and again, her episodes of coma were always associated with severe hyponatremia. Reports document RSW in 38% of patients in general hospital wards, with 21 of 24 lacking cerebral disease, and hence the proposals for a name change to RSW were made.

The initial presentations of case 2 with fever, cough, neck swellings, and positive TB contact constituted the diagnostic consensus is an initial screening test of early childhood TB is still underdiagnosed with missed opportunities prevalent in health facilities, and thus there is an urgent need to raise the index of suspicion at all levels of care to identify these children. Early diagnosis and prompt treatment of TB hyponatremia is lifesaving because SIADH secretion, CSWS, and AI all require different regimens.

**Conclusion**

Childhood TB is still underdiagnosed with missed opportunities prevalent in health facilities, and thus there is an urgent need to raise the index of suspicion at all levels of care to identify these children. Early diagnosis and prompt treatment of TB hyponatremia is lifesaving because SIADH secretion, CSWS, and AI all require different regimens.

**Specific Disclosure**

Some of the results of these studies have been previously reported in the form of a published conference abstract (ATS 2019 Dallas, Texas).

Authors’ Contributions
Conception and design, or acquisition, or analysis and interpretation of data: A.U.E., E.A.O, and K.O.I.

Drafting the article or revising it critically for important intellectual content: A.U.E.

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Conflicts of Interest
None declared.

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**References**


22 Ewa AU, Ochang EA, Inaku KO, Anachuna KC, Ivoke EJ. Recurrent hyponatremia in tuberculous pleural effusion—who is the enemy within? Am J Respir Crit Care Med 2019;199:A4974.