Application of established pathophysiologic processes brings greater clarity to diagnosis and treatment of hyponatremia

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Abstract

Hyponatremia, serum sodium < 135 mEq/L, is the most common electrolyte abnormality and is in a state of flux. Hyponatremic patients are symptomatic and should be treated but our inability to consistently determine the causes of hyponatremia has hampered the delivery of appropriate therapy. This is especially applicable to differentiating syndrome of inappropriate antidiuresis (SIAD) from cerebral salt wasting (CSW) or more appropriately, renal salt wasting (RSW), because of divergent therapeutic goals, to water-restrict in SIAD and administer salt and water in RSW. Differentiating SIAD from RSW is extremely difficult because of identical clinical parameters that define both syndromes and the mindset that CSW occurs rarely. It is thus insufficient to make the diagnosis of SIAD simply because it meets the defined characteristics. We review the pathophysiology of SIAD and RSW, the evolution of an algorithm that is based on determinations of fractional excretion of urate and distinctive responses to saline infusions to differentiate SIAD from RSW. This algorithm also simplifies the diagnosis of hyponatremic patients due to Addison’s disease, reset osmostat and prerenal states. It is a common perception that we cannot accurately assess the volume status of a patient by clinical criteria. Our algorithm eliminates the need to determine the volume status with the realization that too many factors affect plasma renin, aldosterone, atrial/brain natriuretic peptide or urine sodium concentration to be useful. Reports and increasing recognition of RSW occurring in patients without evidence of cerebral disease should thus elicit the need to consider RSW in a broader group of patients and to question any diagnosis of SIAD. Based on the accumulation of supporting data, we make the clinically important proposal to change CSW to RSW, to eliminate reset osmostat as type C SIAD and stress the need for a new definition of SIAD.

Key words: Hyponatremia; Cerebral-renal salt wasting; Fractional excretion of urate

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INTRODUCTION

Hyponatremia, defined as a serum sodium < 135 mEq/L, is the most common electrolyte abnormality that is undergoing changes in methodology with the potential of bringing greater clarity to diagnosis and improved therapeutic outcomes. The present approach initiates the work up of a hyponatremic patient by assessing the status of their extracellular volume despite a unanimous agreement that we cannot accurately estimate the volume status of patients by usual clinical criteria[1-3]. Nevertheless, we continue to attempt to determine whether the patient is euvolemic, hypovolemic or hypervolemic while considering other nuances such as urine osmolality (UNa), serum osmolality and hormones such as renin, aldosterone and atrial/brain natriuretic peptide. We decided to abandon the very tenuous volume approach to hyponatremia and constructed a new algorithm that will hopefully bring greater clarity to the diagnosis and treatment of hyponatremia in addition to encouraging others to explore other parameters that might resolve many of the controversies that exist today.

In this review, we will discuss the evolution of a new approach that will hopefully create changes that are based on credible and reproducible data. We will stress the complexity and importance of differentiating syndrome of inappropriate antidiuresis (SIAD) from cerebral salt wasting (CSW) or more appropriately renal salt wasting (RSW). The importance of this differentiation can be appreciated by our realization that virtually all patients with hyponatremia are symptomatic and should, therefore, be treated[4-9]. Hyponatremia has created a perfect storm for complications by increasing osteoporosis, especially in an elderly population that is already undergoing bone demineralization, and inducing a fourfold increase in falls and fractures[6,7,10]. Textbooks and review articles in medicine consider CSW to be a rare clinical entity as compared to neurosurgeons and critical care physicians who consider CSW/RSW to be common. There is thus an urgency to differentiate CSW/RSW from SIAD because of differences in therapeutic goals, to water-restrict in SIAD and administer salt and water in CSW/RSW. Differentiating CSW/RSW from SIAD has been further complicated by having identical key parameters that identify both syndromes (Table 1). Both syndromes present with hyponatremia, hypouricemia, increased fractional excretion of urate (FEurate), concentrated urine, urine sodium usually > 20 mEq/L with normal renal, adrenal and thyroid function (Table 1). The diagnosis of SIAD cannot, therefore, be made merely because it fulfills the criteria used to define SIAD[11]. A major difference between both syndromes is the volume status (euvolemic/hypervolemic in SIAD and hypovolemic in CSW/RSW), the assessment of which is clinically in both syndromes and how the presence of edema can be found in CSW/RSW. SIADH: Syndrome of inappropriate anti-diuretic hormone; RSW: Renal salt wasting; FE: Fractional excretion; CSW: Cerebral salt wasting.

Table 1  Listing the clinical features that are found in syndrome of inappropriate anti-diuretic hormone and cerebral salt wasting/renal salt wasting

| Findings common to both SIADH and RSW |
|---------------------------------------|
| Association with intracranial disease |
| Hyponatremia                           |
| Concentrated urine                     |
| Urine sodium [Na] usually > 20 mEq/L   |
| Non-euvolemic                          |
| Hypouricemia, with increased fractional excretion urate [FEurate] |
| Only difference between SIADH and RSW |
| Volume state: Normal/high in SIADH, low in RSW |

Note: How the overlapping features are the most commonly encountered clinically in both syndromes and how the presence of edema can be found in CSW/RSW. SIADH: Syndrome of inappropriate anti-diuretic hormone; RSW: Renal salt wasting; FE: Fractional excretion; CSW: Cerebral salt wasting.

EVOLUTION OF A NEW ALGORITHM

The report of hypouricemia, serum urate < 4 mg/dL, with increased FEurate of > 10%, coexisting with hyponatremia in SIAD by Beck in 1979 concluded that the coexistence of hypouricemia and hyponatremia differentiated SIAD from most other causes of hypona-
Hyponatremia

FEurate

< 4%
Volume depletion
Addison’s disease
Edematous states
CHF
Cirrhosis
Nephrosis

4%-11%
Psychogenic polydipsia
Reset osmostat

11%
Normal

FEurate < 11%
SIADH
HCTZ

RSW

FEurate > 11%

Figure 1 Proposed algorithm based on determinations of FEurate to evaluate hyponatremic patients without the need to assess the volume status of the patient or determinations of UNa, plasma renin or aldosterone levels. The dotted line connecting normonatremia and RSW with FEurate > 11% needs further verification. Modified from Ref. [28]. SIADH: Syndrome of inappropriate anti-diuretic hormone; RSW: Renal salt wasting; HCTZ: Hydrochlorothiazide; CHF: Congestive heart failure.

Interestingly, the increased FEurate in SIAD returned to normal when the hyponatremia was corrected by water restriction[15]. Others not only demonstrated this unique relationship between hypouricemia and hyponatremia in SIAD but also noted normalization of a previously increased FEurate with correction of hyponatremia by water restriction[16-20]. We encountered a patient with bronchogenic carcinoma and hypouricemia and increased FEurate, concentrated urine, UNa of 42 mEq/L and normal renal, adrenal and thyroid function, who presented with postural hypotension and reflex tachycardia that was consistent with volume depletion. He responded well to saline infusions with a rapid rise in serum sodium but in view of the coexistence of hyponatremia, hypouricemia and increased FEurate a diagnosis of SIAD was made. Despite the absence of cerebral disease and negative CT scan of brain a diagnosis of salt wasting was made because of postural hypotension with reflex tachycardia and a rapid increase in serum sodium. We corrected the hyponatremia by water restriction to determine whether FEurate would normalize or remain increased as postulated. Correcting the hyponatremia by water restriction and salt supplementation resulted in symptoms of hypovolemia, including return of postural hypotension with reflex tachycardia, postural dizziness, slurred speech, somnolence and staggered gait. The serum sodium finally increased to 138 mEq/L while FEurate remained increased at 14.7%. The clinical course of the patient was highly consistent with volume depletion due to RSW as the persistently increased FEurate after correction of hyponatremia was pathophysiologically different from SIAD. We instead postulated it might be a common feature of RSW[21]. We proceeded to report FEurate to be persistently increased after correction of hyponatremia by water restriction in a patient with metastatic pancreatic carcinoma with ascites, edema and serum albumin of 1.1 mg/dL, bronchogenic carcinoma metastatic to brain, disseminated cryptococcosus with meningitis and uncomplicated Hodgkin’s disease[21]. Absence of cerebral disease in 3 of the 5 patients suggested at this time that RSW can occur without clinical evidence of cerebral disease but we waited until we had stronger evidence for RSW to make such a proposal. We reported hypouricemia, hyponatremia, increased FEurate (many with normonatremia and some after correction of hyponatremia) and cerebral atrophy in patients with various intracranial diseases and in AIDS, 10 of whom had postural hypotension and CVP of 0 cm H2O that were consistent with RSW[22-24]. As a result of these studies, we felt we had enough data to propose differentiating SIAD from RSW by correcting the hyponatremia and observing whether there was normalization of a previously increased FEurate as in SIAD or was persistently increased as in RSW (Figures 1 and 2)[14,25-28]. This unique relationship between FEurate and serum sodium appears to have identified two pathophysiologically different groups of patients.

Reset osmostat

While attempting to sort out the relationship between FEurate and serum sodium, we encountered hyponatremic patients who met the criteria for SIAD and RSW but had normal FEurates[29]. Of 14 consecutive hyponatremic patients with normal FEurate, 6 had spontaneously excreted dilute urines which was diagnostic of a reset osmostat (RO). The remaining 8 patients had a normal water-loading test to prove the diagnosis of RO, 6 of whom had undetectable plasma antiuretic hormone (ADH) levels at a time when the urine was dilute to prove further the diagnosis of RO (Figure 3)[30]. As noted in Figure 3, type C SIAD represents patients with RO, which was found to make up about 30% of the patients studied[26,30]. Our experience is that many hyponatremic patients admitted to the hospital have intercurrent illnesses that reset their osmostat and normalizes after resolution of their intercurrent illness. This was exemplified by a patient with a renal transplant who presented with mild hyponatremia of 133 mEq/L on a routine outpatient visit. He later developed a fever for 10 d and was admitted to the hospital with a pneumocystis pneumoniae infection after falling and being confused with a serum sodium of 119 mEq/L. A diagnosis of RO was made by noting spontaneously excreted dilute urine of only 92 mosm/kg and normal FEurate of 7% and 8%. His serum sodium returned to normal one month after successful treatment of his pneumonia. This case illustrates how a slowly evolving pneumonia had...
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gradually lowered his osmostat and serum sodium[31]. RO, however, can also exist chronically for over 10 years and the persistently normal FErrate is not consistent with the proposal that chronic hyponatremia is responsible for the high FErrate in SIAD[29,32]. While a normal FErrate has effectively identified patients with RO, patients with psychogenic polydipsia also have a normal FErrate, but differentiating RO from psychogenic polydipsia can be readily made by the large volumes of water ingested and polyuria with dilute urines in psychogenic polydipsia (Figure 1)[33]. Not only did this study simplify the diagnosis of RO but it provided additional data to support our proposal that FErrate was superior to hypouricemia when evaluating patients with hyponatremia. We have noted normal or increased FErrate with hypouricemia and increased FErrate with normonatremia[22,24,29]. It also provided important pathophysiologic data to propose eliminating RO as a subtype of SIAD by virtue of a normal FErrate and predictable inhibition of ADH secretion by water loading[30].

ADDITIONAL CONTRIBUTIONS OF DETERMINING FErrATE

Determinations of FErrate have also been extremely useful in identifying the cause of hyponatremia in other clinical conditions. In a recent publication, we report a hyponatremic patient with a bronchogenic carcinoma who met the criteria for SIAD and RSW (Table 1). His unresponsiveness to liberal amounts of saline was most consistent with SIAD so to prove the diagnosis of SIAD we administered 1.5% hypertonic saline to increase serum sodium to 138 mEq/L while observing a gradual decrease in FErrate from 26.2% at baseline, to 11% and 8% (Figure 4)[34]. This maneuver not only provided a potentially convenient way to differentiate SIAD from RSW by correcting the hyponatremia to see if FErrate normalizes as in SIAD or remains persistently increased as in RSW (Figures 1 and 2). It also confirmed our long-held contention that contrary to popular opinion, saline has only a meager effect on FErrate (Table 2)[26,34-36].

RSW with edema

Determining FErrate was pivotal in a patient with advanced Hodgkin’s disease who presented with a serum sodium of 127 mEq/L, Uosm of 308 mosm/kg, a 20 pound weight gain over one month with increasing edema of both lower extremities, postural hypotension and reflex tachycardia, ascites, pleural effusion, urine sodium of only 10 mEq/L, decreased cardiac output and normal renal, thyroid and adrenal function[29]. The nephrology attending made a diagnosis of RSW based on the combination of a high FErrate of 17.2% and postural hypotension and reflex tachycardia that was consistent with a volume depleted state. The lower extremity edema was postulated to be due to obstruction of the inferior vena, a finding that was later confirmed by CT scan of the abdomen. Isotonic saline was administered and within 10 h after initiation of saline therapy, the Uosm decreased to 140 mosm/kg when a plasma ADH was undetectable. The dilution of urine and undetectable plasma ADH level illustrated how saline eliminated the more potent volume stimulus for ADH secretion and allowed the coexisting hyposmolality to inhibit ADH secretion (Figure 5)[35]. Because of the increase in free water excretion his serum sodium increased by 6 mEq/L after 12 h so DSW was administered to prevent serum sodium from increasing beyond 6 mEq/L per day to prevent osmotic demyelination[29]. On the second day, the medical team, using the unreliable volume approach, made an alternate diagnosis of congestive heart failure that was based on the edema, ascites, pleural effusion, UNa of 10 mEq/L, concentrated urine and decreased cardiac output (actually due to volume depletion) and administered

Table 2: Summary of extracellular volume expansion with isotonic, hypotonic and hypertonic saline on fractional excretion of sodium [FESodium] and urate [FErrate] at control and experimental periods after saline administration. Note the meager changes in FErrate despite very high FENa.

|       | Control | Exp | Control | Exp | Ref. |
|-------|---------|-----|---------|-----|------|
| Isotonic | 1.04  | 4.43 | 7.98  | 9.76 | [36] |
| Hypertonic | 1.6  | 8.2  | 5.0   | 5.8  | [35] |
| Hypotonic | 2.9  | 18.6 | 5.4   | 12.1 | [33] |

Printed with permission Ref. [26]. Exp: Experimental.
furosemide. Because the proximal tubule is the main site of natriuretic activity in RSW, see below, the large sodium load that would ordinarily be transported by the distal nephron was inhibited by furosemide resulting in a markedly increased urine output and hemodynamic instability that required large volumes of saline to attain hemodynamic stability. The initial finding of a high FEurate was pivotal in arriving at the diagnosis of RSW because a patient in heart failure and normal kidneys would decrease FEurate to < 4% (Figure 1)\[31\]. The UNa of only 10 mEq/L was consistent with a prerenal state in which proximal tubule solute reabsorption increases to give rise to low UNa, decreased FEurate of < 4% and the hallmark increase in BUN to creatinine ratio\[39\]. The low UNa was actually indicative of a decreased appetite and low salt intake, which can occur in RSW and SIAD.

Addison’s disease
Addison’s disease is a well-known cause of RSW, but in contrast to other causes of RSW, mineralocorticoid deficiency induces a defect in sodium transport in the distal nephron to create a volume depleted state with an intact proximal tubule where urate is exclusively transported\[40\]. The volume depletion in Addison’s disease due to a defect in sodium transport in the distal tubule leads to the classic prerenal state in which an intact proximal tubule increases solute transport to give rise to the oft-used increase in the BUN to creatinine ratio to characterize a prerenal state\[39\]. So the low FEurate of 1.4% was a simple means of identifying Addison’s disease as the cause of hyponatremia\[31\]. The hyperkalemia, high UNa of 140 mEq/L, high ACTH and low cortisol levels accompanied the low FEurate and the
dilution of urine after initiation of saline led to a rapid increase in serum sodium that required D5W infusions to retard the rate of increase in serum sodium to avoid osmotic demyelination. These features collectively strengthened the diagnosis of volume depletion due to RSW of distal tubular origin. A low FEurate of 1.4% in a hyponatremic patient should thus include Addison’s disease (Figure 1).

We have accumulated sufficient data to rely on the value of determining FEurate in hyponatremic conditions. While the algorithm can effectively diagnose conditions such as RO, psychogenic polydipsia, Addison’s disease and prerenal causes of hyponatremia, there is still the formidable task of differentiating SIAD from CSW/RSW. The diagnosis of SIAD cannot be made simply by satisfying the definition of SIAD, which is virtually identical to CSW/RSW (Table 1). We will review the salient pathophysiological features of both syndromes, highlight some important pathophysiological differences, critically review the literature and present a suggested approach to differentiate SIAD from RSW.

PATHOPHYSIOLOGY OF SIAD
The syndrome of inappropriate secretion of antidiuretic hormone, first described in 1957 by Schwartz et al. and known simply as SIAD, rightfully deserves to be regarded as a syndrome that was derived by pure reason, recognizing patients who had identical clinical characteristics as healthy subjects receiving daily injections of ADH without being able to determine ADH levels in plasma. It is now well-known that ADH can be ectopically produced without being responsive to volume and osmolar stimuli, hence the term inappropriate secretion of ADH, that is, ADH levels being increased in the presence of euvoLemia and hypo-osmolality of plasma. They initially go into negative sodium balance and then quickly reach a steady state where sodium input equals sodium output. Because of persistently increased ADH levels, renal water reabsorption decreases serum sodium levels. They eventually escape the effects of ADH by increasing urine volume in part by markedly decreasing renal aquaporin-2 protein. Patients with SIAD typically present with hyponatremia and hypochloremia, hypouricemia with increased FEurate, concentrated urine, UNa usually exceeding 20 mEq/L, absence of edema and normal renal, thyroid and adrenal function (Table 1). They are considered to be euvoLemic according to textbook descriptions of the syndrome but are actually hypervolemic as determined by sulfate space and the gold standard radio isotope dilution methods, using chromium-labeled red blood cells and/or radioiodinated serum albumin. Plasma renin and aldosterone levels are normally suppressed and because of the volume expansion, ANP levels are increased. Euvolemia is a necessary prerequisite to the diagnosis of SIAD because hypovolemia is a normal and potent stimulus for ADH secretion and takes precedence over the inhibitory effect of plasma hypo-osmolality (Figure 5). Unfortunately, extracellular volume cannot be accurately assessed by usual clinical criteria. SIAD is frequently associated with a myriad of comorbid conditions that often involve the pulmonary and central nervous systems. Because ADH increases renal water retention, treatment is to water restrict these patients to < 1000 mL/d with some salt supplementation or inhibit the V2 receptor for ADH, the vaptans.

Need for reclassification and redefinition of SIAD
SIAD is currently classified by 4 different types of responses to serum hypo-osmolality (Figure 3), type A with ADH levels being increased throughout a wide spectrum of serum osmolalities, Type B where ADH levels remain slightly increased as serum osmolality decreases, type C a reset osmostat with ADH being inhibited at a lower serum osmolality than normal and type D where ADH levels are suppressed throughout the
spectrum of serum osmolalities due to a gain in function of the V2 receptor that is referred to as nephrogenic syndrome of inappropriate antidiuresis\(^{30}\). Because the patients selected for this study were considered to be euvoicmep by unreliable clinical methods, it is probable that type B SIAD may be renal salt wasters because the patients in this study were given hypertonic saline to increase serum osmolality to normal and then given a water load to determine ADH levels as serum osmolality decreased. The assumption is that these patients had RSW but remained volume depleted to maintain increased levels of ADH despite the hypertonic saline\(^{30}\). The classification of type C SIAD has also come under scrutiny because patients with RO have pathophysiologic differences from SIAD by a normal FEurate as compared to being increased in SIAD and noting how lowering serum osmolality appropriately and predictably inhibits ADH secretion\(^{39}\). RO should thus be considered a separate clinical syndrome and eliminated as type C SIAD. Elimination of RO as a subtype of SIAD should thus lead to a readjustment of SIAD by eliminating the Uosm of < 100 mosm/kg, since excretion of dilute urines, Uosm < Posm, is extremely unusual in SIAD, vide infra\(^{11,30}\).

**CEREBRAL/RENAL SALT WASTING**

CSW, as first described by Peters et al\(^{49}\) in 1950, had fallen into disrepute for many years in part because, as previously reviewed, they did not prove RSW in their report. SIAD became the preferred and dominant syndrome. RSW is now accepted as a legitimate syndrome that exists and is being recognized with greater frequency. It appears that a natriuretic factor, as discussed below, initiates RSW by increasing renal salt and water excretion to a point where the patient becomes volume depleted, the extent being determined by the combination of the severity of the inhibition on sodium transport and salt and water intake. The defect in sodium transport appears to reside in the proximal tubule because urate, which is transported exclusively in the proximal tubule by reabsorbing and secreting transporters, is also reduced\(^{40}\). An increase in BUN to creatinine ratio as previously proposed in RSW has not been confirmed by recent reports addressing this problem\(^{12,13}\). In RSW, similarities in the BUN to creatinine ratio in SIAD and RSW can be explained by the reduced sodium and water reabsorption in the proximal tubule that cannot establish a urea concentration gradient that is necessary for passive urea reabsorption to occur\(^{49}\). The patient initially goes into a state of negative sodium balance, but like SIAD eventually reaches a new steady state where input of salt matches output by activation of humoral, hemodynamic and neuronal factors, but reaches a steady state at a lower extracellular volume. UNa can thus be low if sodium intake is low as can be noted in both SIAD and RSW and is, thus, not a consistently reliable marker of both syndromes. It is extremely unlikely that atrial or brain natriuretic peptides are responsible for RSW as proposed since their main meager action is in the distal tubule and have been found to be elevated in disparate conditions such as SIAD, subarachnoid hemorrhage (SAH) and salt retaining states such as congestive heart failure and low normal in an unequivocal case of RSW\(^{42}\).

In contrast to SIAD, ADH levels in RSW are appropriately increased because the more potent volume stimulus perpetuates the hyponatremia despite the increasing plasma hypo osmolality (Figure 5). As we demonstrated in 4 previous unequivocal cases of RSW, saline eliminates the volume stimulus for ADH secretion to reduce ADH to undetectable levels and allowing the coexistent hypo-osmolality to inhibit ADH secretion, which induces free water excretion by diluting the urine and promptly increasing serum sodium (Figure 6)\(^{12,13,31}\). In contrast, saline did not dilute the urine or significantly increase serum sodium in two documented cases of SIAD in whom the diagnosis of SIAD was made by an increased FEurate and hypervolemia as determined by gold standard radioisotope dilution methods by \(^{51}\)chromium-labeled red blood cells and radioiodinated serum albumin (Figure 7)\(^{13}\). This same maneuver has been used by others to propose a volume-depleted hyponatremic state if isotonic saline increases serum sodium increases by more than 5 mEq/L and at the same time commenting on our inability to asses accurately the volume status of patients by clinical criteria\(^{41}\).

Our case of a hyponatremic patient with a hip fracture and no clinical evidence of cerebral disease illustrates the essential features of RSW and proves unequivocally the existence of RSW\(^{12}\). When seen in consultation, she was hyponatremic and hyperchloremic with normal renal, adrenal and thyroid function, concentrated urine, Uosm 321 mosm/kg and a UNa of only 6 mEq/L. Her hyponatremia was initially thought to be a due to a volume-depleted state with normal renal function that would give rise to a prerenal state where proximal tubule solute reabsorption would increase to give rise to a low FEurate of usually < 4%\(^{39}\). A low serum urate of 3.4 mg/dL led to a determination of FEurate which was elevated at 29.6% and was thus included in our IRB-approved protocol. Her baseline plasma renin, aldosterone and ADH levels were increased, ANP was low normal at 35 pg/mL and blood volume as determined by \(^{51}\)chromium-labeled red blood cells and radio-iodinated serum albumin was reduced by 7.1%\(^{12}\). She received isotonic saline and as can be seen in Figure 6, Uosm progressively decreased to dilute levels 13 h after initiation of isotonic saline infusion when plasma ADH was undetectable as would be expected in a hypovolemic, hyponatremic patient with RSW. The excretion of free water promptly increased serum sodium to normal within 48 h when FEurate remained persistently increased (Figure 6)\(^{12}\). The patient reported feeling much better and was hungry upon arising 18 h after initiation of isotonic saline infusion, which reflected an improvement of...
her volume depleted state by an erroneous diagnosis of SIAD and fluid restricted for 10 d prior to our intervention. This patient satisfies all of the important features of RSW by the persistence of an elevated FEurate after correction of hyponatremia (Figures 1 and 2), and illustrated the appropriate increase in plasma ADH, elimination of the volume stimulus by saline to permit the coexistent hypo-osmolality to inhibit ADH secretion, excretion of dilute urines and prompt correction of hyponatremia. It also illustrates how RSW can occur without clinical evidence of cerebral disease, how the low ANP is a very unlikely cause of RSW, how fluid restriction of a patient with RSW for an erroneous diagnosis of SIAD can increase morbidity and how a low salt intake can result in low UNa even in RSW (and in SIAD as well). A similar case of RSW associated with an uncomplicated pneumonia and no evidence of cerebral disease responded to isotonic saline infusions by diluting the urine and correcting the hyponatremia within 48 h after initiation of isotonic saline infusions[13]. This case provided us with enough credible data to activate our earlier instincts by proposing to change CSW to RSW[14].

**COMMENTS ON NATRIURESIS IN RSW AND RELATIONSHIP BETWEEN NATREMIA AND URATE TRANSPORT**

Natriuresis in RSW. The syndrome of RSW has been clearly established from a clinical perspective but the mechanism by which this occurs has not been resolved. The consistency with which we were encountering urate transport abnormalities in RSW suggested that a circulating natriuretic factor, if present, would have its major effect on proximal tubule sodium transport. But because of the ability of the distal tubule to increase sodium transport, we reasoned that sodium excretion induced by a factor affecting proximal sodium transport might be minimized or nullified. To this end we elected to study lithium transport in rats exposed to plasma from neurosurgical patients who have been reported to have RSW and high FEurates[24,45,46,50]. Without entering into the controversy whether or not lithium is exclusively transported in the proximal tubule, lithium is known to be transported on a 1:1 basis with sodium almost exclusively in the proximal tubule under ambient conditions[51]. We injected the plasma from 21 neurosurgical patients, 14 of whom had FEurates exceeding 10%, and 14 age and gender-matched controls into rats and demonstrated a significant increase in FENa from 0.29% to 0.59% and FElithium from 24% to 36.6%, in controls and neurosurgical patients, respectively[52].

We extended our interest in patients with increased FEurate with normonatremia by investigating patients with Alzheimer’s disease (AD) who have been reported to be hypouricemic[53]. We studied 18 patients with advanced AD, mini-mental examination scores of < 10, and compared them to 6 patients with multi-infarct or vascular dementia (MID) and 11 normal age and gender-matched controls. All patients were normonatremic except for one hyponatremic patient with AD, FEurate being significantly higher and serum urate lower in AD as compared to the other groups. Infusion of the sera from all groups of patients resulted in FENa of 0.33%, 0.38% and 0.63% and FElithium 27.2%, 31.2% and 41.7% in control, MID and AD, respectively. FENa and FElithium were significantly increased in AD as compared to control and MID. In both the neurosurgical and AD rat studies, the distal sodium load of 36.6% and 41.7% as determined by the FElithium, had a significant sodium uptake by the distal nephron to account for the FENa of only 0.59% and 0.63% in the final urine. The RSW patient with B cell lymphoma obstructing the inferior vena cava had a similarly high distal sodium load that was inhibited by furosemide to generate an exaggerated diuresis that resulted in profound hemodynamic instability[31]. Interestingly, there were no differences in blood pressure or inulin clearances throughout both studies.

![Figure 6: Changes in urine osmolality and serum sodium concentration during isotonic infusion in a hyponatremic patient with a hip fracture without clinical evidence of cerebral disease. The low Uosm at baseline reflected a low sodium intake when UNa was only 6 mEq/L and a weakened medullary solute concentration, which increased rapidly with a marked increase in Uosm after 4 h of saline infusion. Plasma antidiuretic hormone (ADH) was increased with increased plasma renin and aldosterone levels at baseline and ADH level was undetectable when the urine was dilute 13 h with increase in serum sodium to 138 mEq/L within 48 h after initiation of isotonic saline infusion. Printed with permission, Ref. [12].](image-url)
In a third group of studies, proteins were purified from ammonium sulfate precipitates of urine proteins and placed in transwells to determine transport of radioactive sodium, \( ^{22}\text{Na} \), across cultured porcine proximal tubule cells, LLC-PK1 cells, grown to confluence on a semipermeable membrane. Urine proteins from normonatremic neurosurgical patients with increased FEurate inhibited \( ^{22}\text{Na} \) transport across cultured LC-PK1 cells as compared to those with normal FEurate\(^{54}\).

These studies demonstrate the presence of a natriuretic factor in plasma and urine of normonatremic neurosurgical patients with increased FEurate and in sera of normonatremic AD patients with increased FEurate. The natriuretic factor has its major effect on proximal tubule sodium transport to support our proposal that a high FEurate in the presence of normonatremia might be a marker of RSW without going through a phase of hyponatremia (Figure 1). Future studies must address this interesting possibility. Atrial or brain natriuretic peptides are extremely unlikely as the natriuretic factor in RSW because their main site of action is in the inner medullary collecting duct\(^{55}\).

**Relationship between natremia and FEurate**

The intriguing relationship between serum sodium and FEurate has been uniquely coupled in many hyponatremic and non hyponatremic conditions such as RSW. Because the natriuretic factor present in the plasma of patients with RSW affect sodium transport mainly in the proximal tubule, it would be interesting to speculate that the natriuretic factor might also affect reabsorbing and/or secretory transporters or anion exchangers for urate in the proximal tubule\(^{50}\). This circulating factor can have effects on the sodium and urate transporters regardless of whether the patient is hyponatremic or normonatremic.

The relationship between sodium and urate in SIAD continues to elude any rational explanations. It can be readily understood why FEurate remains persistently increased in the presence of normonatremia in RSW, but in SIAD, normalization of FEurate after correction of hyponatremia has not been fully explained. Some have implicated the V1 receptor activity of pitressin to explain the increase in FEurate in SIAD but others were able to induce SIAD with increased FEurate in healthy volunteers by DDAVP which lacks any V1 activity. In addition, the V1 activity of pitressin is an unlikely cause of the increase in FEurate in SIAD because pitressin levels are still increased when FEurate normalizes after correction of hyponatremia\(^{56}\). The same group has made commendable efforts to explain the increase in FEurate in SIAD by implicating chronic hyponatremia, but the normal FEurate reported in psychogenic polydipsia and reset osmostat where hyponatremia has been documented for up to 10 years do not support such an hypothesis\(^{52}\). At the present time, the relationship between serum sodium and FEurate remains unexplained in SIAD.

**Prevalence of RSW**

The prevalence of RSW in hyponatremic patients is presently unknown because of multiple factors: (1) the term CSW confined RSW to patients with cerebral disease; (2) textbooks and review articles in internal medicine consistently consider CSW to be a rare entity or is not included as a clinical entity of hyponatremia; (3) utilizing the ineffective volume approach to hyponatremia; and (4) lack of application of the pathophysiologic characteristics in the evaluations of SIAD and RSW. As noted in Table 1, overlapping of key clinical parameters that characterize SIAD and RSW makes it extremely difficult to differentiate one syndrome from the other. One key difference is to determine the volume status of the patient by credible means at a time when they are hyponatremic. There are 3 studies in hyponatremic and normonatremic neurosurgical patients that determined intravascular volume by gold standard radio isotope dilution methods, using \( ^{51}\text{Cr} \)chromium-labeled red blood cells and radio iodinated serum albumin. As can be seen in Table 3, 83% and 94% of patient with UNa exceeding 40 mEq/L and a variety of neurosurgical diseases had decreased blood volumes that were
consistent with CSW/RSW. By comparison only 3 of the 39 patients studied had increased blood volumes that were consistent with SIAD. Interestingly, 67% of normonatremic patients with SAH had decreased blood volumes that were consistent with RSW while only 33% had increased volume consistent with SIAD. RSW occurring in normonatremic patients is consistent with the popular view of CSW/RSW but we would like further clarification of whether normonatremia in the presence of a high FEurate is indicative of RSW (Figure 1). Fluid restricting patients with SAH was harmful by increasing morbidity and mortality so it is customary to administer saline to patients with SAH because of the prevalence of RSW in patients with SAH.

These volume studies in neurosurgical patients overcame the most important deficiency when trying to differentiate SIAD from RSW, but are rarely considered when discussing CSW. Instead, a retrospective study of patients with SAH where CSW was found in only 6.5% of patients with hyponatremia, the majority being SIAD, is cited to prove the rarity of CSW/RSW. In two retrospective studies they determined whether their method of analysis that 4.8% and 2.7% of patients had combined CSW and SIAD, a highly incongruous combination that might occur, but would be extremely difficult to prove, especially in a retrospective study. We critically reviewed many flawed articles on CSW/RSW in a previous publication and feel it is important to comment on a more recent prospective study of hyponatremia in patients with SAH by the same group which concluded that the hyponatremia was again due to SIAD. The diagnosis of SIAD was made because they fulfilled the definition of SIAD without commenting on the same definition being applicable to RSW. The accompanying editorial to this manuscript agreed with the findings in this large study and alluded to the rarity of CSW. A critical review of this manuscript reveals that the authors followed the recommendation to administer saline rather than fluid-restrict hyponatremic patients with SAH. Virtually every patient received isotonic saline from the time of admission to the hospital, none were fluid restricted and none received hypertonic saline or a vaptan class of drugs at any time.

It is intriguing to note that 36 of 49 patients developed hyponatremia within 3 d after SAH. This unaddressed outcome raises questions as to whether the patients received free water or desalinated when UNa exceeded 150 mEq/L while receiving isotonic saline. Every patient corrected their hyponatremia during the hospitalization with the median time of correction being 3 d. Because of the correction of hyponatremia, they concluded that the "syndrome" was short lived. It is well known that patients with SIAD do not respond to saline infusions to correct their hyponatremia; otherwise there would be no need for fluid restriction or treatment with the vaptan class of drugs. Instead, a more likely diagnosis is RSW, which is consistent with the volume studies performed in patients with SAH (Table 3).

Isotonic saline eliminated the volume stimulus for ADH secretion and permitted the coexisting hypo-osmolality to inhibit ADH secretion, excrete dilute urines and predictably correct the hyponatremia, which does not happen in SIAD. ADH levels would have been undetectable at the time of excretion of dilute urines as we demonstrated in cases of RSW within 24-48 h after initiation of isotonic saline therapy. We contend that these patients would have had high FEurates exceeding 11% and would have remained increased when serum sodium returned to normal (Figures 1 and 2). Correction of hyponatremia would not mean there was resolution of the underlying RSW. The reader is encouraged to read a more thorough critical review of controversial papers in the literature.

**Differentiating SIAD from RSW**

As noted in Table 1 and Figures 1 and 2, SIAD and RSW share common clinical parameters including an increased FEurate > 11%. An increased FEurate most often involves RSW and SIAD with hydrochlorothiazide (HCTZ) and possibly selective serotonin reuptake inhibitors (SSRIs) drugs coming into the picture. Discontinuation of HCTZ and SSRIs should result in correction of the hyponatremia but persistence of the hyponatremia and increased FEurate would indicate that SIAD must be differentiated from RSW in order to arrive at the proper therapeutic strategy of water-restricting or administering V2 receptor blockers for SIAD or administering saline for RSW. We have now accumulated enough data to utilize two strategies to differentiate SIAD from RSW. The first is to correct the hyponatremia by either water-restriction or use of hypertonic saline to determine whether FEurate normalizes as in SIAD or remains increased as in RSW (Figures 1, 2 and 4). It is important to note that contrary to common perceptions, saline has a very meager effect on FEurate as noted in Figure 4, Table 2. This is illustrated by Figure 4 when large volumes of saline, including hypertonic saline, gradually decreased FEurate from 26.2% to 8% as serum sodium increased to normal to confirm the diagnosis of SIAD.

The other alternative is to administer saline soon after FEurate has been shown to be increased, > 11%. As discussed above, the urine will usually become dilute

| Table 3  Summary of volume studies by gold standard radio-isotope dilution methods in hyponatremic and normonatremic neurosurgical patients |
|---------------------|-----------------|-----------------|-----------------|
| Ref.               | n of patients  | Low blood volume RSW | Increased blood volume SIADH | Urine Na mEq/L |
| Nelson et al[45]   | HN             | 12              | 10 (83%)       | 2               | 41-203       |
| Wijdicks et al[46] | HN             | 9               | 8 (89%)        | 1               | --           |
| NN                 | 12             | 8 (67%)        | 1               | 4               |
| Sivakumar et al[47] | HN         | 18             | 17 (94%)       | 4               | 43-210       |

RSW much more common than SIAD. Printed with permission, Ref. [26]. SIADH: Syndrome of inappropriate anti-diuretic hormone; RSW: Renal salt wasting; HN: Hyponatremic; NN: Normonatremic.
within the first 24-48 h after initiation of saline infusion in RSW with a rapid increase in serum sodium which should not increase by more than 4-6 mEq/L per 24 h to prevent osmotic demyelination[38]. The rapid increase can be slowed by infusing DSW or administering DDAVP intranasally realizing that the recommended correction to prevent osmotic demyelination is based on 24 h and not less[38]. In a thorough review of the literature, we have found dilution of urine in SIAD only under two conditions, infusing isotonic saline at a rate of 16 mL/min for 2 h in the first case of SIAD reported in 1957 and extreme salt restriction in experimentally induced SIAD in normal human subjects[42,43]. It would appear therefore that the differentiation of SIAD from RSW can be accomplished by noting normalization of a previously increased FEurate as in SIAD or persistence of increased FEurate in RSW after correction of hyponatremia and response to saline infusions (Figures 1, 2, 6 and 7).

By utilizing both strategies and the algorithm in Table 2 and Figure 3 in a completed study of 52 patients with hyponatremia outside of the neurosurgical intensive care unit, we found about an equal number of patients with SIAD, RO and RSW with the majority of patients with RSW to be free of clinical evidence of cerebral disease. The surprising large number of patients with RSW and absence of cerebral disease in more than 80% of these patients supports our previous proposal to change CSW to RSW[14]. This change has enormous clinical relevance because RSW would not be considered in the absence cerebral disease[64].

**FEphosphate**

We have previously stated that an increased FEphosphate > 20% at baseline is consistent with RSW[21]. We have encountered only one patient with an increased FEphosphate with RSW, but while it might be potentially useful in differentiating SIAD from RSW, there are pitfalls that can alter its value. We have demonstrated in a previous renal micropuncture study that phosphate transport is very sensitive to saline infusions, because parathyroid hormone (PTH) increases rapidly as calcium and possibly magnesium decrease in serum to stimulate PTH release[65]. Since saline is frequently administered to patients with hyponatremia, determinations of FEphosphate must be performed at baseline or analyzed according to whether or not the patient was receiving saline at the time the test was performed.

**CONCLUSION**

Hyponatremia, the most common electrolyte, is undergoing fundamental changes that would benefit from a new mindset of abandoning the volume approach. We present an algorithm based on supportive data where the determination of FEurate has been helpful in arriving at a more accurate diagnosis of the causes of hyponatremia. The realization that hyponatremic patients are symptomatic has led to the recommendation to treat virtually all hyponatremics, thus creating an urgency to differentiate SIAD from RSW because of divergent therapeutic goals. The clinical utility of applying two distinctive pathophysiologic characteristics to distinguish SIAD from RSW, such as: (1) demonstrating normalization of a previously increased FEurate in SIAD and persistent increase in FEurate as in RSW after correction of hyponatremia by water restriction or hypotonic saline or; (2) to note whether infusion of isotonic saline induces excretion of dilute urines with a prompt increase in serum sodium as in RSW or continued excretion of concentrated urines without correction of hyponatremia as in SIAD. Utilization of this pathophysiologic approach has uncovered many with RSW without cerebral disease. So contrary to popular perceptions of the rarity of CSW/RSW, it is probable that the increase in morbidity and mortality associated with hyponatremia may in part be iatrogenic because patients with RSW are being fluid-restricted for an erroneous diagnosis of SIAD. Finally, we feel it is time to abandon the volume approach to patients with hyponatremia and to appreciate the unreliability of determining plasma renin and aldosterone levels because of so many circumstances such as the use of ACE inhibitor, angiotensin II receptor blockers or saline that affect their blood levels, and to be aware of the following comments to which we have supporting data: (1) have less reliance on UNa in RSW and SIAD as sodium excretion is dependent on sodium intake; (2) the misconceptions of having an increased BUN to creatinine ratio in RSW when there is no difference; (3) appreciate the remarkable overlapping of common clinical characteristics between SIAD and RSW (Table 1); and (4) because of the expanding prevalence of RSW occurring in all hyponatremic patients, we must be vigilant in insisting on clearly distinguishing SIAD from RSW in the future and to question previous reports on SIAD, realizing that RSW could be included in significant numbers. We feel we have accumulated enough data to support these final comments and to encourage others to expand our efforts to derive the true prevalence of RSW in a broad population of hyponatremic and normonatremic patients.

**REFERENCES**

1. Chung HM, Kluge R, Schrier RW, Anderson RJ. Clinical assessment of extracellular fluid volume in hyponatremia. *Am J Med* 1987; 83: 905-908 [PMID: 3674097 DOI: 10.1016/0002-9344(87)90649-8]
2. Singh S, Bohn D, Carlotti AP, Cusimano M, Rutka JT, Halpern ML. Cerebral salt wasting: truths, fallacies, theories, and challenges. *Crit Care Med* 2002; 30: 2575-2579 [PMID: 12441772 DOI: 10.1097/0003246-20021100-00028]
3. Oh MS, Carroll HJ. Cerebral salt-wasting syndrome. We need better proof of its existence. *Nephron* 1999; 82: 110-114 [PMID: 10364701 DOI: 10.1159/000045385]
4. Decaux G. Is asymptomatic hyponatremia really asymptomatic? *Am J Med* 2006; 119: 579-582 [PMID: 16843909 DOI: 10.1016/j.amjmed.2006.05.013]
5. Gankam Kengne F, Andres C, Sattar L, Melot C, Decaux G. Mild hyponatremia and risk of fracture in the ambulatory elderly. *QJM* 2008; 101: 583-588 [PMID: 18477645 DOI: 10.1093/qjmed/chn002]

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6 Hoorn EJ, van der Lubbe N, Zietse R. SIADH and hyponatraemia: why does it matter? NDT Plus 2009; 2: iii-iii[11] [PMID: 19881934]

7 Renneboog B, Musch W, Vandenbergel X, Manto MU, Decaux G. Mild chronic hyponatraemia is associated with falls, unsteadiness, and attention deficits. Am J Med 2006; 119: 71.e1-71.e8 [PMID: 16431193 DOI: 10.1016/j.amjmed.2005.09.026]

8 Schrier RW. Does ‘asymptomatic hyponatraemia’ exist? Nat Rev Nephrol 2010; 6: 185 [PMID: 20348927 DOI: 10.1038/nrne ph.2010.21]

9 Arief AI, Llach F, Massry SG. Neurological manifestations and morbidity of hyponatraemia: correlation with brain water and electrolytes. Medicine (Baltimore) 1975; 55: 121-129 [PMID: 1256311 DOI: 10.1097/00005792-197603000-00002]

10 Verbalis JG. Barsony J, Sugimura Y, Tian Y, Adams DJ, Carter EA, Resnick HE. Hyponatremia-induced osteoporosis. J Bone Miner Res 2010; 25: 554-563 [PMID: 19751154 DOI: 10.1359/jbmr.090827]

11 Janicic N, Verbalis JG. Evaluation and management of hypom- osmolality in hospitalized patients. Endocr Rev Clin North Am 2003; 32: 459-481, vii [PMID: 12805041 DOI: 10.1016/S0272-6386(03)00004-5]

12 Maesaka JK, Miyawaki N, Palaia T, Fishbane S, Durham JH. Renal salt wasting without cerebral disease: diagnostic value of urine determinations in hyponatraemia. Kidney Int 2007; 71: 822-826 [PMID: 17311074 DOI: 10.1038/sj.ki.5002093]

13 Bitew S, Imbriano LJ, Miyawaki N, Fishbane S, Maesaka JK. More on renal salt wasting without cerebral disease: response to saline infusion. Clin J Am Soc Nephrol 2009; 4: 309-315 [PMID: 19201917 DOI: 10.2215/CJN.07240608]

14 Maesaka JK, Imbriano LJ, Ali NM, Ilamathi E. Is it cerebral or renal salt wasting? Kidney Int 2006; 73: 934-938 [PMID: 19641485 DOI: 10.1038/ki.2009.263]

15 Beck LH. Hypouricemia in the syndrome of inappropriate secretion of antidiuretic hormone. N Engl J Med 1979; 301: 528-530 [PMID: 466006 DOI: 10.1056/NEJM197909033010005]

16 Assadi FK, John EG. Hypouricemia in neonates with syndrome of inappropriate secretion of antidiuretic hormone. Pediatr Res 1985; 19: 424-427 [PMID: 4000678 DOI: 10.1203/00006435-198508000-00003]

17 Dorhout Mees EJ, Blom van Assendelft P, Nieuwenhuis MG. Elevation of uric acid clearance caused by inappropriate antidiuretic hormone secretion. Acta Med Scand 1971; 189: 69-72 [PMID: 5121533 DOI: 10.1111/j.1600-0462.1971.tb04340.x]

18 Sonnenblick M, Rosin A. Increased uric acid clearance in the syndrome of inappropriate secretion of antidiuretic hormone. N Engl J Med 1978; 24: 20-23 [PMID: 3346144]

19 Weinberger A, Santo M, Solomon F, Shalit M, Pinkhas J, Sperling BF, editor. Novel Insights on chronic kidney disease, acute kidney injury and polyuric/polysytic kidney disease. Croatia: Intech; 2012: 41-66

20 Maesaka JK, Imbriano LJ, Ali NM, Miyawaki N, Maesaka JK. Normal fractional urine excretion identifies hyponatremic patients with renal salt wasting? J Am Soc Nephrol 2012; 25: 833-838 [PMID: 22307440 DOI: 10.1159/000360074]

21 Zerbe R, Stropes L, Robertson G. Vasopressin expression in the syndrome of inappropriate antidiuresis. Annu Rev Med 1980; 31: 315-327 [PMID: 677290 DOI: 10.1146/annurev.me.31.020800.01513]

22 Imbriano LJ, Mattana J, Drakakis J, Maesaka JK. Identifying Different Causes of Hyponatremia With Fractional Excretion of Uric Acid. Am J Med Sci 2016; 352: 385-390 [PMID: 27776720 DOI: 10.1016/j.amjms.2016.05.035]

23 Decaux G, Prosper F, Soupert A, Musch W. Evidence that chronicity of hyponatremia contributes to the high urine clearance observed in the syndrome of inappropriate antidiuretic hormone secretion. Am J Kidney Dis 2000; 36: 745-751 [PMID: 11007676 DOI: 10.1053/ajkd.2000.17623]

24 Ali N, Imbriano LJ, Maesaka JK. The Case A 66-year-old male with hyponatremia. Psychogenic polydipsia. Kidney Int 2009; 76: 233-234 [PMID: 19564861 DOI: 10.1038/ki.2009.150]

25 Cannon PJ, Svahn DS, Demartini FE. The influence of hypertonic saline infusions upon the fractional reabsorption of urine and other ions in normal and hypertensive man. Circulation 1970; 41: 97-108 [PMID: 5420637 DOI: 10.1161/01.CIR.41.1.97]

26 Diamond H, Meisel A. Influence of volume expansion, serum sodium, and fractional excretion of sodium on urine excretion. Pfugers Arch 1975; 356: 47-57 [PMID: 1238979 DOI: 10.1007/BF00583520]

27 Steele TH. Evidence for altered renal urine reabsorption during changes in volume of the extracellular fluid. J Lab Clin Med 1969; 74: 288-299 [PMID: 5799512]

28 Stricker EM, Verbalis J. Water intake and body fluids. In: Squire LE, Roberts JL, Spitzer NC, Zigmond MJ, McConnell SK, Bloom BF, editors. Fundamental Neuroscience. 2nd ed. California: Elsevier; 2003: 1011-1029

29 Sterns RH. Disorders of plasma sodium--causes, consequences, and correction. N Engl J Med 2015; 372: 55-65 [PMID: 25551526 DOI: 10.1056/NEJMra1404899]

30 Abuelo JG. Normotensive ischemic acute renal failure. N Engl J Med 2007; 357: 797-805 [PMID: 17715412 DOI: 10.1056/NEJMr064398]

31 Lipkowitz MS. Regulation of uric acid excretion by the kidney. Curr Rheumatol Rep 2012; 14: 179-188 [PMID: 22359229 DOI: 10.1007/s11926-012-0240-z]

32 Schwartz WB, Bennett W, Curelop S, Barter FC. A syndrome of renal sodium loss and hyponatraemia probably resulting from inappropriate secretion of antidiuretic hormone. Am J Med 1975; 53: 529-542 [PMID: 13469824 DOI: 10.1016/0002-9345(75)90224-3]

33 Leaf A, Bartter FC, Santos RF, Wrong O. Evidence in man that urinary electrolyte loss induced by pitressin is a function of water retention. J Clin Invest 1953; 32: 868-878 [PMID: 13084753 DOI: 10.1172/JCI102805]

34 Jaenike JR, Waterhouse C. The renal response to sustained administration of vasopressin and water in man. J Clin Endocrinol Metab 1961; 21: 231-242 [PMID: 13789146 DOI: 10.1210/jcem-21-3-231]

35 Verbalis JG, Murase T, Ecelbarger CA, Nielsen S, Knepper MA.
Studies of renal aquaporin-2 expression during renal escape from vasopressin-induced antidiuresis. *Adv Exp Med Biol* 1998; 449: 395–406 [PMID: 10002683 DOI: 10.1007/978-1-4615-4871-3_51]

**54** Nelson PB, Seif SM, Maroon JC, Robinson AG. Hyponatremia in intracranial disease: perhaps not the syndrome of inappropriate secretion of antidiuretic hormone (SIADH). *J Neuromusi* 1981; 55: 938–941 [PMID: 7299468 DOI: 10.3171/jnms.1981.55.6.0938]

**55** Wijdicks EF, Vermeulen M, ten Haaf JA, Hijdra A, Bakker WH, van Gijn J. Volume depletion and natriuresis in patients with a ruptured intracranial aneurysm. *Ann Neurol* 1985; 18: 211–216 [PMID: 4037761 DOI: 10.1002/ana.410180208]

**56** Berl T, Quitinna-Pelletier F, Verbalsis JG, Schrier RW, Bichet DG, Ouyang J, Czerwiec FS. Oral tolvaptan is safe and effective in chronic hyponatremia. *J Am Soc Nephrol* 2010; 21: 705–712 [PMID: 20185637 DOI: 10.1681/ASN.2009080857]

**57** Peters JP, Welt LG, Sims EA, Orloff J, Needham J. A salt-wasting syndrome associated with cerebral disease. *Trans Assoc Am Physicians* 1950; 63: 57–64 [PMID: 14855556]

**58** Palmer BF. Hyponatremia in patients with central nervous system disease: SIADH versus CSW. *Trends Endocrinol Metab* 2003; 14: 182–187 [PMID: 12714279 DOI: 10.1016/S1043-2760(03)00048-1]

**59** Sivakumar V, Rajeshkher V, Chandy MJ. Management of neurosurgical patients with hyponatremia and natriuresis. *Neurosurgery* 1994; 34: 269–274; discussion 274 [PMID: 8177388 DOI: 10.1227/00006123-199402000-00010]

**60** Hayslett JP, Kashgarian M. A micropuncture study of the renal handling of lithium. *Pflügers Arch* 1979; 380: 159–163 [PMID: 573443 DOI: 10.1007/BF00582152]

**61** Maesaka JK, Venkatesan J, Piccione JM, Dockr R, Dreisbach AW, Wetherington J. Plasma natriuretic factor(s) in patients with intracranial disease, renal salt wasting and hyperuricosuria. *Life Sci* 1993; 52: 1875–1882 [PMID: 8502124 DOI: 10.1016/0024-3205(93)90008-G]

**62** Maesaka JK, Wolf-Klein G, Piccione JM, Ma CM. Hypouricemia, abnormal renal tubular urine transport, and plasma natriuretic factor(s) in patients with Alzheimer’s disease. *J Am Geriatr Soc* 1993; 41: 501–506 [PMID: 8468882 DOI: 10.1111/j.1532-5415.1993.tb01885.x]

**63** Youmans SJ, Wein MR, Wirkowski E, Maesaka JK. Demonstration of natriuretic activity in urine of neurosurgical patients with renal salt wasting. *F1000Res* 2013; 2: 126 [PMID: 24358843 DOI: 10.12688/f1000research.2-126.v2]

**64** Beltowski J, Wójcicka G. Regulation of renal tubular sodium transport by cardiac natriuretic peptides: two decades of research. *Med Sci Monit* 2002; 8: RA39–RA52 [PMID: 11859295]

**65** Decaux G, Namias B, Gubitus B, Soufrant A. Evidence in hyponatremia related to inappropriate secretion of ADH that V1 receptor stimulation contributes to the increase in renal uric acid clearance. *J Am Soc Nephrol* 1996; 7: 805–810 [PMID: 8738818]

**66** Wijdicks EF, Vermeulen M, Hijdra A, van Gijn J. Hyponatremia and cerebral infarction in patients with ruptured intracranial aneurysms: is fluid restriction harmful? *Ann Neurol* 1985; 17: 137–140 [PMID: 3977297 DOI: 10.1002/ana.410170206]

**67** Ellinson DH, Berl T. Clinical practice. The syndrome of inappropriate antidiuresis. *N Engl J Med* 2007; 356: 2064–2072 [PMID: 17507705 DOI: 10.1056/NEJMcp066837]

**68** Sherlock M, O’Sullivan E, Agha A, Behan LA, Rawluk D, Brennan P, Tormey W, Thompson CJ. The incidence and pathophysiology of hyponatraemia after subarachnoid haemorrhage. *Clin Endocrinol (Oxf)* 2006; 64: 250–254 [PMID: 16487432 DOI: 10.1111/j.1365-2265.2006.02432.x]

**69** Sherlock M, O’Sullivan E, Agha A, Behan LA, Owens D, Finucane F, Rawluk D, Tormey W, Thompson CJ. Incidence and pathophysiology of severe hyponatraemia in neurosurgical patients. *Postgrad Med J* 2009; 85: 171–175 [PMID: 19417163 DOI: 10.1136/pgmj.2008.072819]

**70** Hannon MJ, Behan LA, O’Brien MM, Tormey W, Ball SG, Javadpour M, Sherlock M, Thompson CJ. Hyponatremia following mild/moderate subarachnoid hemorrhage is due to SIAD and glucocorticoid deficiency and not cerebral salt wasting. *J Clin Endocrinol Metab* 2014; 99: 291–298 [PMID: 24248182 DOI: 10.1210/jc.2013-3032]

**71** Verbalis JG. Hyponatremia with intracranial disease: not often cerebral salt wasting. *J Clin Endocrinol Metab* 2014; 99: 59–62 [PMID: 24384014 DOI: 10.2107/jcem.2013.8289]

**72** Steele A, Gowrishankar M, Abrahamson S, Mazur CD, Feldman RD, Halperin ML. Postoperative hyponatremia despite near-isotonic saline infusion: a phenomenon of desalinization. *Ann Intern Med* 1997; 126: 20–25 [PMID: 8992919 DOI: 10.7326/0003-4819-126-1-199701010-00003]

**73** Maesaka JK, Gallagher D, Mattana J, Imbriano L. Unexpectedly High Frequency of Reset Osmostat and Cerebral-renal Salt Wasting in Non-edematous Hyponatremia: Value of Determining total Extracellular and Intracellular Water Volumes and Fractional Excretion of Urate. *Proceedings of the American Society of Nephrology, Annual Kidney Week*; 2013 Nov 5–10; Atlanta, Georgia. PUB 258, p 932

**74** Maesaka JK, Levit MF, Abramson RG. Effect of saline infusion on phosphate transport in intact and thyroparathyroidectomized rats. *Am J Physiol* 1973; 215: 1421–1429 [PMID: 4760455]

**75** Robertson GL. Regulation of arginine vasopressin in the syndrome of inappropriate antidiuresis. *Am J Med* 2006; 119: S36-S42 [PMID: 16843083 DOI: 10.1016/j.amjmed.2006.05.006]

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