Human cells dwell in salt water. Their well-being depends on the ability of the body to regulate the salinity of extracellular fluids. By controlling water intake and excretion, the osmoregulatory system normally prevents the plasma sodium concentration from straying outside its normal range (135 to 142 mmol per liter). Failure of the system to regulate within this range exposes cells to hypotonic or hypertonic stress. This review considers the causes and consequences of an abnormal plasma sodium concentration and offers a framework for correcting it.

Plasma Sodium Concentration and Extracellular Tonicity

The plasma sodium concentration affects cell volume. The term “tonicity” describes the effect of plasma on cells — hypotonicity makes cells swell and hypertonicity makes them shrink. Hypernatremia always indicates hypertonicity. Hyponatremia usually indicates hypotonicity, but there are exceptions (e.g., hyperglycemic hyponatremia and pseudohyponatremia) that are not covered in this review.

Plasma Sodium Concentration and the Electrolyte and Water Content of the Body

Solute concentrations (osmolalities) must be equal inside and outside of cells because water channels (aquaporins) make cell membranes permeable to water. The “sodium pump” (Na⁺/K⁺–ATPase) functionally excludes sodium from cells, exchanging it for potassium by means of active transport. Although sodium is largely extracellular and potassium is intracellular, body fluids can be considered as being in a single “tub” containing sodium, potassium, and water, because osmotic gradients are quickly abolished by water movement across cell membranes. As such, the concentration of sodium in plasma water should equal the concentration of sodium plus potassium in total body water. This theoretical relationship was validated empirically by Edelman et al., who used isotopes to measure exchangeable body cations and water.

Edelman and colleagues described the relation between these variables with the following equation:

\[
[\text{Na}^+]_{\text{plasma H}_2\text{O}} = 1.11 \times \frac{(\text{Na}^+_{e} + \text{K}^+_{e})}{\text{total body H}_2\text{O}} - 25.6,
\]

where \(\text{Na}^+_{e}\) is exchangeable sodium, \(\text{K}^+_{e}\) exchangeable potassium, and \(\text{H}_2\text{O}\) water. This equation has an intercept (–25.6); the regression line relating plasma sodium
The ratio of exchangeable (Na\(^+\) + K\(^-\)) to total body water does not pass through zero because not all exchangeable sodium is free in solution.\(^4\) A substantial amount of sodium is bound to large polyanionic macromolecules called proteoglycans, which make up the ground substance of bone, connective tissue, and cartilage (Fig. 1).\(^1\) The sodium concentration of cartilage is nearly twice that of plasma. The osmotic force created by the high sodium concentration (about 40 mm Hg for every difference in concentration of 1 mmol per liter) maintains the high water content in the tissue, allowing it to withstand pressures that can exceed 20,000 mm Hg during exercise.\(^5\)

When it became known that much of the sodium in the body is bound to bone, cartilage, and connective tissue, it was hypothesized that these tissues could serve as sodium reservoirs, taking up or releasing sodium in response to the needs of the body.\(^6\) Despite early evidence supporting the concept of a sodium reservoir,\(^7\) this theory lost favor\(^8\) and was not pursued for half a century. However, the past decade has seen renewed interest in stored sodium.\(^9\) In patients who consume high-salt diets, sodium can accumulate in the body, seemingly disappearing without a change in the plasma sodium concentration, body weight, or extracellular fluid volume.\(^10\) Sodium, potassium, and water balance do not always account for changes in the plasma sodium concentration during recovery from hyponatremia.\(^11\) Proteoglycans in skin serve as a sodium reservoir, and the number of negative charges available to bind sodium varies in response to the sodium concentration of interstitial tissue.\(^12\) In experiments in rats, chronic hyponatremia has been shown to be a more potent cause of osteopenia than vitamin D deficiency, and loss of sodium from bone exceeded the loss of calcium from bone. The activity of osteoclasts is increased in chronic hyponatremia owing to a direct effect of sodium and possibly vasopressin on these cells.\(^13\)

In humans, chronic hyponatremia is associated with osteoporosis and fractures. During extreme-endurance athletic events lasting several hours, bone density decreases measurably, and the decrease in bone density correlates remarkably closely with changes in the plasma sodium concentration.\(^14\)

**Figure 1. Internal and External Solute and Water Balance and the Plasma Sodium Concentration.**

The plasma sodium concentration is determined according to the ratio of the content of sodium and potassium in the body (the numerator of the ratio) to total body water (the denominator of the ratio). This concentration is altered by net external balances (intake minus output) of sodium, potassium, and water and by internal exchange between sodium that is free in solution and sodium that is bound to polyanionic proteoglycans in bone, cartilage, and skin.
A simplified version\textsuperscript{19} of the equation reported by Edelman et al. is
\[
\text{plasma } [\text{Na}^+] = \frac{\text{total body } (\text{Na}^+ + \text{K}^+)}{\text{total body } H_2O}.
\]

The plasma sodium concentration is altered by changes in overall sodium and potassium balance (the numerator of the simplified equation) and water balance (the denominator) (Fig. 1). To understand or roughly predict changes in the plasma sodium concentration, the overall tonicity of the diet and intravenous fluids and the overall tonicity of gastrointestinal fluids, sweat, and urine must be considered. Like the plasma sodium concentration, which is determined by the concentrations of sodium and potassium in body water, the tonicity of these fluids is defined by their concentrations of sodium plus potassium.

It is not possible to predict the effect of administering intravenous fluids on the plasma sodium concentration without considering concurrent urinary losses. The electrolyte concentration (sodium plus potassium) of urine, and not its osmolality (which includes electrolytes, urea, and glucose), determines the effect of urine on the plasma sodium concentration. Urine is hypotonic if its electrolyte concentration is lower than that of plasma; because it is partly composed of electrolyte-free water, excretion of hypotonic urine will increase the plasma sodium concentration. Conversely, urine is hypertonic if its electrolyte concentration is higher than that of plasma; excretion of hypertonic urine will lower plasma sodium concentrations.\textsuperscript{20}

Isosmolar or hyperosmolar urine containing mostly urea (an end product of protein metabolism) may be nearly electrolyte-free.\textsuperscript{20,21} Excretion of urea owing to recovery from azotemia, catabolism, or a high-protein diet will cause hypernatremia unless there is replacement of the free water that has been lost.\textsuperscript{22} Because it increases excretion of electrolyte-free water, urea has been used to treat hyponatremia.\textsuperscript{23,24}

Consequently, in most tissues, the sodium concentrations of plasma and interstitial fluid are nearly identical, with a small difference created by intravascular albumin.\textsuperscript{1,25} In contrast, brain capillaries have tight endothelial junctions and are lined by astrocytic foot processes, creating a blood–brain barrier that sodium cannot cross (Fig. 2).\textsuperscript{26} Consequently, an abnormal plasma sodium concentration causes water to enter or leave brain tissue. Because of the confines of the skull, only a small degree of brain swelling or shrinkage is compatible with life.

Since the plasma sodium concentration affects brain volume, it is not surprising that the cell-volume receptors that are responsible for adjusting thirst and vasopressin secretion are located in the brain. Osmoreceptors, which are more accurately called tonicity receptors, are hypothalamic neurons that express transient receptor potential cation channel subfamily vanilloid member 1 (TRPV1) and member 4 (TRPV4) channels on their cell membranes.\textsuperscript{27,28}

Transient receptor potential cation channels belong to a large family of molecules. They were first identified as photoreceptors in fruit flies and were later discovered to serve as receptors for a variety of sensations throughout nature.\textsuperscript{29} For example, transient receptor potential V (or vanilloid) (TRPV) channels respond to capsaicin, a vanilloid that causes the burning sensation associated with the ingestion of chili peppers. TRPV1, which is a member of the TRPV family of receptors, was identified in mutant \textit{Caenorhabditis elegans} roundworms that did not avoid hyperosmotic environments. Insertion of a mammalian TRPV4 gene into the genome of mutant \textit{C. elegans} roundworms restored normal worm behavior. The TRPV1 gene is required for normal functioning of isolated osmoreceptor neurons, and genetically engineered mice that lack genes for TRPV1 and TRPV4 have abnormal osmoregulation. Polymorphisms in genes encoding the TRPV4 channel have been identified in humans. Healthy aging men who are positive for the TRPV4 P19S polymorphism are more likely to have mild hyponatremia than are men without this polymorphism.\textsuperscript{30}

In normal osmoregulation, both thirst and vasopressin secretion are inhibited when the plasma sodium concentration is decreased below 135 mmol
per liter. In the absence of vasopressin, urine osmolality decreases to as low as 50 mOsm per kilogram. In persons who consume a typical Western diet, with an output of urinary solute of about 900 mOsm daily, a urinary solute concentration of 50 mOsm per liter yields 18 liters of urine (750 ml per hour).20

Although individual responses vary, vasopressin is usually detectable at a plasma sodium concentration above 135 mmol per liter, and levels of vasopressin increase linearly with increasing sodium levels.32 The hormone may also be secreted in response to circulatory inadequacy,33 or it may be secreted “inappropriately,” and sometimes ectopically, with no osmotic or hemodynamic stimulus.34 (Vasopressin secretion
without an osmotic or hemodynamic abnormality to account for it is termed “inappropriate.” Once secreted, vasopressin binds to its V2 receptor on basolateral membranes of principal cells lining the renal collecting duct. In the presence of vasopressin, aquaporins are inserted into the luminal membrane, allowing water to flow out, attracted by the high solute concentration of the surrounding medullary interstitium. When the plasma sodium level increases to approximately 145 mmol per liter, vasopressin levels are normally high enough to result in maximally concentrated urine (about 1200 mOsm per kilogram). The presence of a dilute urine when the plasma sodium concentration is above 145 mmol per liter implies either deficient vasopressin secretion (as in neurogenic diabetes insipidus) or failure of the kidneys to respond to vasopressin (as in nephrogenic diabetes insipidus) (see Case 1 in the Supplementary Appendix, available with the full text of this article at NEJM.org).

However, even complete diabetes insipidus (with total absence of vasopressin or no tubular response to vasopressin) generally does not cause hyponatremia, because thirst prompts replacement of urinary losses of water. Hypernatremia develops if water is unavailable, if the urge to drink is impaired (hypodipsia), or if patients are too young, old, or sick to seek water themselves.

Maximal dilution of urine prevents hyponatremia unless water intake is extraordinarily large (>1 liter per hour) (e.g., in patients with schizophrenia who drink water compulsively) or the rate of urinary solute excretion is extremely low (e.g., in beer drinkers who eat very little). Except in these scenarios, hypotonic urine is associated with an impaired ability of the body to dilute urine because of diminished sodium transport in renal-diluting sites (most commonly because of the use of diuretics), the presence of vasopressin, or, rarely, an inherited activating mutation of the vasopressin receptor. Because vasopressin, along with renin, angiotensin, aldosterone, and the sympathetic nervous system, participates in the neurohumoral response to inadequate circulation, vasopressin-mediated hyponatremia may complicate hypovolemia or states that lead to edema (e.g., heart failure and cirrhosis).

Many causes of hyponatremia (e.g., hypovolemia, medications, cortisol deficiency, nausea, pain, or stress) are reversible — either by treatment or the passage of time. Once the cause of hyponatremia resolves, the normal osmoreceptor response to a low plasma sodium concentration inhibits vasopressin secretion, resulting in excretion of maximally dilute urine and resolution of hyponatremia.

Administration of a vasopressin antagonist will also result in the excretion of dilute urine (also known as a “water diuresis” or “aquaresis”) despite the continued presence of vasopressin. Therefore, vasopressin antagonists increase the plasma sodium concentration and are approved by the Food and Drug Administration for the treatment of hyponatremia caused by the syndrome of inappropriate antidiuretic hormone secretion (SIADH) or by heart failure.

**Urinary Sodium and the Plasma Sodium Concentration**

Urinary excretion of sodium is relatively independent of plasma sodium levels; measurement of urinary sodium levels can help to distinguish between SIADH and hypovolemic hyponatremia. Excretion of sodium responds to intravascular volume, increasing with volume expansion and decreasing with volume depletion. Water retention due to SIADH expands extracellular fluid volume, resulting in increased urinary excretion of sodium despite hyponatremia.

Excretion of sodium in SIADH restores normal extracellular volume, but hypertonic urinary losses also exacerbate hyponatremia. For this reason, SIADH should not be treated with isotonic solutions such as 0.9% saline or Ringer’s lactate, because infused sodium will be excreted in smaller volumes of urine, leading to net retention of electrolyte-free water. Such a sequence is common in patients with subarachnoid hemorrhage; saline is prescribed to maintain cerebral perfusion, but vasopressin released by neurologic injury concentrates the urine (see Case 2 in the Supplementary Appendix).

**Causes of Rapid Changes in the Plasma Sodium Concentration**

The plasma sodium concentration will decrease rapidly if the amount of water ingested or infused exceeds the capacity of the kidneys to excrete free water. The plasma sodium concentration increas-
es rapidly if large amounts of concentrated salt are ingested or infused or if there are large, unreplaced losses of electrolyte-free water because of aquareesis or osmotic diuresis (most commonly due to glycosuria). Loss or gain of approximately 3 ml of water per kilogram of body weight will change the plasma sodium concentration by approximately 1 mmol per liter.51 Maximally dilute urine, whether resulting from untreated diabetes insipidus, spontaneous recovery from hyponatremia, or administration of a vasopressin antagonist, will increase the plasma sodium concentration by about 2.5 mmol per liter per hour. In the absence of urinary loss of water, 1 ml of 3% saline per kilogram of body weight will increase the plasma sodium concentration by about 1 mmol per liter.51 Therefore, in a woman with a body weight of 50 kg, the increase in the plasma sodium level caused by a maximum water diuresis is similar to the increase caused by infusion of approximately 125 ml of 3% saline per hour.

**CONSEQUENCES OF AN ABNORMAL PLASMA SODIUM CONCENTRATION**

Extreme hypotonicity ruptures cell membranes; extreme hypertonicity damages the cytoskeleton and causes breaks in DNA, ultimately leading to apoptosis.52 Given time, cells protect their volume and survival by adjusting intracellular solute contents.53 Organic osmolytes are small intracellular molecules (e.g., glutamate, taurine, and myo-inositol) that are found throughout nature; their concentrations can vary without perturbing cell functions.54 Hypotonicity promotes the release of osmolytes from cells through volume-sensitive leak pathways, while, concurrently, osmolyte-accumulating transporters (e.g., the taurine transporter TauT and the myo-inositol transporter SMIT) are down-regulated. With hypertonicity, TauT and SMIT are up-regulated.53,54 These adaptations allow cells to maintain intracellular solute concentrations that are equal to the osmolality of hypotonic or hypertonic plasma, with little change in cell volume.53,54

Although osmotic disturbances affect all cells, clinical manifestations of hyponatremia and hypernatremia are primarily neurologic, and rapid changes in plasma sodium concentrations in either direction can cause severe, permanent, and sometimes lethal brain injury (Tables 1 and 2 and Fig. 3).59,60 If severe hypernatremia develops over a period of minutes (e.g., after massive ingestion of salt that may occur in a suicide attempt), vascular injury created by a suddenly shrinking brain causes intracranial hemorrhage. Brain swelling from an abrupt onset of hyponatremia results in increased intracranial pressure, impairing cerebral blood flow and sometimes causing herniation (Fig. 3). Adaptive changes in brain osmolytes permit survival, but they may also contribute to symptoms.55 For example, in acute hyponatremia, adaptive release of glutamate, an excitatory neurotransmitter, may increase the susceptibility to seizures; depletion of the transmitter from nerve terminals may account for some of the neurologic symptoms of chronic hyponatremia.56

The foot processes of astrocytes, which encircle both brain capillaries and neurons, express aquaporins (such as aquaporin-4) that allow water to cross the blood–brain barrier. Astrocytes protect neurons from osmotic stress; in response to hypotonicity, a cell-to-cell transfer of taurine to adjacent astrocytes allows neurons to maintain their volume while astrocytes swell.56 Within 24 to 48 hours after this transfer, astrocytes restore their volume through loss of organic osmolytes, but this makes them vulnerable to injury from rapid normalization of the plasma sodium concentration. Because of the down-regulation of transporters, recovery of lost brain osmolytes may take a week or longer.55,56 Therefore, rapid correction of hyponatremia is a hypertonic stress to astrocytes that are depleted of osmolytes, triggering apoptosis, disruption of the blood–brain barrier, and, eventually, brain demyelination (see the Supplementary Appendix). In experiments in animals, brain demyelination has been prevented by repletion of myo-inositol,58 by lowering the plasma sodium concentration again promptly (within 12 to 24 hours after rapid correction of hyponatremia),59 or by administration of minocycline (which prevents proliferation of glial cells).60

Brain injury after rapid correction of chronic hyponatremia manifests as a biphasic illness called the osmotic demyelination syndrome: an initial reduction in symptoms is followed by a gradual onset of new neurologic findings (see Case 3 in the Supplementary Appendix).61 The clinical spectrum of the osmotic demyelination syndrome is broad and can include seizures, behavioral abnormalities, and movement disorders.62
require ventilator support — have a full functional disability or death, many patients — even those who are normally protective against oxidative injury. Acute hypernatremia may also cause brain demyelination, without the biphasic clinical course of the osmotic demyelination syndrome (Fig. 3). Acute hypernatremia, like chronic hyponatremia, causes a reversible encephalopathy. Particularly in infants, organic osmolytes gained in the adaptation to chronic hypernatremia are lost slowly. Therefore, rehydration resulting in rapid correction of chronic hypernatremia causes seizures and a bulging fontanelle indicating cerebral edema (Fig. 3).

A plasma sodium concentration that is even slightly outside the normal range increases the risk of death, but few deaths associated with abnormalities in the plasma sodium concentration are related to neurologic complications. The underlying disorders that produce an abnormal plasma sodium concentration may be responsible for excess mortality, but there may also be non-neurologic adverse consequences of prolonged osmotic disturbances, and observational studies have shown decreased mortality among hospitalized patients in whom the plasma sodium concentration was corrected. Taurine and myo-inositol, organic osmolytes that are lost from many cells in the adaptation to hyponatremia, are normally protective against oxidative injury. An experimental model of chronic SIADH in rats showed that prolonged hyponatremia resulted in hypogonadism, loss of body fat, skeletal-muscle sarcopenia, and cardiomyopathy.

**Correction of an Abnormal Plasma Sodium Concentration**

Clinicians who treat patients with hyponatremia and hypernatremia should respond promptly to the immediate dangers posed by an acute disturbance, while being mindful of adaptations that make excessive correction potentially harmful. Aggressive interventions are indicated when the plasma sodium concentration has decreased or increased rapidly or when an abnormal plasma sodium concentration is causing severe symptoms. Therapy should be guided by frequent monitoring of the plasma sodium concentration and not by formulas alone.
Fatal brain swelling — a rare complication of hyponatremia that clinicians are most concerned about — has only been reported in hypotonic patients with intracranial disease and in a few specific conditions that cause the plasma sodium concentration to decrease rapidly, such as postoperative hyponatremia and self-induced water intoxication that develops over a few hours (Table 1). Because the brain cannot swell by much more than 5%, correction of hyponatremia by this amount would be expected to prevent the most serious complications of acute water intoxication; empirical observations support this prediction. An increase in the plasma sodium concentration of 4 to 6 mmol per liter is enough to reverse impending brain herniation or stop active seizures in patients with severe acute hyponatremia. Such an increase can be reliably achieved with 100-ml bolus infusions of 3% saline (2 ml per kilogram in small patients), administered at 10-minute intervals to a total of three doses, if necessary, to control symptoms. Milder symptoms of acute hyponatremia should be treated with enough 3% saline to avoid a worsening of hyponatremia because of delayed absorption of ingested water or excretion of hypertonic urine.

Hyponatremia is usually a chronic condition and it should be presumed to be chronic when the actual duration is unclear; to reduce symptoms and improve potential outcomes, chronic hyponatremia should be corrected gradually with the use of fluid restriction, salt tablets, slow infusions of 3% saline, furosemide, urea, or vasopressin antagonists, or by treatment of the underlying cause. Severe symptoms of hyponatremia may require
more aggressive initial interventions, but there is no need to increase the plasma sodium concentration by more than 4 to 6 mmol per liter per day. Regardless of how chronic hyponatremia is treated, inadvertent overcorrection, most commonly caused by excretion of dilute urine, is common and can be very dangerous (see Case 3 in the Supplementary Appendix). If the plasma sodium concentration is less than 120 mmol per liter, or if there are risk factors for osmotic demyelination, correction of the plasma sodium concentration by more than 8 mmol per liter per day should be meticulously avoided through replacement of lost water or prevention of water loss with desmopressin.

Repeat therapeutic lowering of the plasma sodium concentration is justified if the correction of hyponatremia exceeds 8 mmol per liter per day and there are risk factors for osmotic demyelination or if the correction is 10 to 12 mmol per liter per day without these risk factors (Table 1) — although the benefit of this strategy has not been confirmed in humans. To prevent inadvertent overcorrection (see Case 4 in the Supplementary Appendix), desmopressin can be administered preemptively, in anticipation of, rather than in response to, unwelcome urinary losses of water; hyponatremia is corrected with a slow infusion of 3% saline while the urine is kept concentrated with repeated doses of desmopressin.

Limiting correction of chronic hyponatremia so that the plasma sodium concentration is decreased by less than 0.5 mmol per liter per hour reduces the risk of cerebral edema and seizures associated with rehydration. However, the fear of these complications, which have been reported only in young children, should not deter the aggressive rehydration of adults with acute hypernatremia to avoid brain hemorrhage or osmotic demyelination (Table 2). In contrast to the risk of inadvertent overcorrection in patients with hyponatremia, there is little risk of inadvertent overcorrection in patients with hypernatremia, and adults with hypernatremia are often undertreated.

**CONCLUSIONS**

Disorders of plasma sodium concentration expose cells to hypotonic or hypertonic stress. Although all cells are affected, clinical manifestations of hyponatremia and hypernatremia are primarily neurologic, and rapid changes in plasma sodium concentrations in either direction can...
cause severe, permanent, and sometimes lethal brain injury. Because the brain adapts to an abnormal plasma sodium level, excessive correction of a chronic disturbance can be injurious and should be avoided.

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REFERENCES

47. Leaf A, Bartert 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;32:686-78.

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