Disorders of plasma sodium are the most common electrolyte disturbances in clinical medicine, yet they remain poorly understood. Severe hyponatraemia and hypernatraemia are associated with considerable morbidity and mortality, however, and even mild hyponatraemia is associated with worse outcomes when it complicates conditions such as heart failure, although which is cause and which effect is often uncertain. Distinguishing the cause(s) of hyponatraemia may be challenging in clinical practice, and controversies surrounding its management remain. Here, we describe the common causes of disorders of plasma sodium, offer guides to their investigation and management, and highlight areas of recent advance and of uncertainty.

Sources and selection criteria

We incorporated the latest consensus from systematic reviews and publications identified by a literature search through Medline and Web of Science with the search strategy terms “hyponatraemia,” “hypernatraemia,” and “sodium.” We found fewer than a dozen randomised controlled trials of treatment of any description. Despite their frequency, plasma sodium disorders have not been reviewed by the Cochrane Library, Clinical Evidence, or Best Evidence.

Control of sodium balance

Under normal conditions, plasma sodium concentrations are finely maintained within the narrow range of 135-145 mmol/l despite great variations in water and salt intake. Sodium and its accompanying anions, principally chloride and bicarbonate, account for 80% of the extracellular fluid osmolality, which is normally 285-295 mosm/kg and calculated as (2× [Na]mmol/l + [urea]mmol/l + [glucose]mmol/l. The main determinant of the plasma sodium concentration is the plasma water content, itself determined by water intake (thirst or habit), “insensible” losses (such as metabolic water, sweat), and urinary dilution. The last of these is under most circumstances the most important and is predominantly determined by arginine vasopressin, which is synthesised in the hypothalamus and then stored in and released from the posterior pituitary. In response to arginine vasopressin, concentrated urine is produced by water reabsorption across the renal collecting ducts. This is mediated by specialised cellular membrane transport proteins called aquaporins.

Hyponatraemia

Determining the cause of hyponatraemia may be straightforward if an obvious precipitating cause is present—for example, in the setting of vomiting or diarrhoea, when both sodium and total body water are low, and especially if the patient (typically elderly) is taking diuretics. In hospital practice, diagnosing the cause is often less clear cut. Here, hyponatraemia almost always reflects an excess of water relative to sodium, commonly by dilution of total body sodium secondary to increases in total body water (water overload) and sometimes as a result of depletion of total body sodium in excess of concurrent body water losses. The clinical classification of hyponatraemia according to the patient’s extracellular fluid volume status, as hypovolaemic, euvoalaemic, or hypervolaemic (box 1), is useful to help with the diagnosis. In practice, however, distinguishing euvoalaemic and hypervolaemic hyponatraemia may not be straightforward.

The symptoms of hyponatraemia are related to both the severity and the rapidity of the fall in the plasma sodium.

Summary points

- Sodium disorders are common, particularly in hospital patients and elderly people
- Mild sodium disorders may be asymptomatic and self-limiting, but severe sodium disorders are associated with considerable morbidity and mortality
- The causes of sodium imbalance are often iatrogenic and therefore avoidable
- Assessing hydration status and measuring sodium in plasma and urine are key to diagnosing the cause of hyponatraemia
- The cause of hypernatraemia will usually be evident from the history
- Little evidence from randomised controlled trials exists for the treatment of sodium disorders
- Slow correction of sodium is usually safe, with careful monitoring of clinical status and plasma sodium
sodium concentration. A decrease in plasma sodium concentration creates an osmotic gradient between extracellular and intracellular fluid in brain cells, causing movement of water into cells, increasing intracellular volume, and resulting in tissue oedema, raised intracranial pressure, and neurological symptoms. Patients with mild hyponatraemia (plasma sodium 130–135 mmol/l) are usually asymptomatic. Nausea and malaise are typically seen when plasma sodium concentration falls below 125–130 mmol/l. Headache, lethargy, restlessness, and disorientation follow, as the sodium concentration falls below 115–120 mmol/l. With severe and rapidly evolving hyponatraemia, seizures, coma, permanent brain damage, respiratory arrest, brain stem herniation, and death may occur. In more gradually evolving hyponatraemia, the brain self regulates to prevent swelling over hours to days by transport of, firstly, sodium, chloride, and potassium and, later, organic solutes including glutamate, taurine, myo-inositol, and glutamine from intracellular to extracellular compartments. This induces water loss and ameliorates brain swelling, and hence leads to few symptoms in patients with chronic hyponatraemia.

**History, examination, and investigation**

An accurate history may reveal a clue to the cause of the hyponatraemia and establish the rapidity of the symptoms. The key diagnostic factors (box 2) are the hydration status of the patient and the urine “spot sodium” concentration, which is available quickly and allows the crucial distinction in hypovolaemic hyponatraemia between renal (high; > 30 mmol/l) and extrarenal (low; < 30 mmol/l) salt loss. Urinary sodium is similarly helpful in patients in whom volume status is difficult to assess, as patients with dilutional hyponatraemia will have a urinary sodium < 30 mmol/l, but may be >30 if dietary access to salt restricted. Plasma osmolality is almost always low in hyponatraemia, and urine is less than maximally dilute (inappropriately concentrated); so, although usually measured, plasma and urine osmolalities are rarely discriminant.

**Box 1: Classification of hyponatraemia**

<table>
<thead>
<tr>
<th>Hypovolaemia</th>
<th>Euvolaemic</th>
<th>Hypervolaemic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extrarenal loss, urine sodium &lt;30 mmol/l</td>
<td>Clinical state may not help diagnostically</td>
<td>Usually easy to diagnose clinically</td>
</tr>
<tr>
<td>• Dermal losses, such as burns, sweating</td>
<td>Plasma urea tends to be low rather than high</td>
<td>Heart failure</td>
</tr>
<tr>
<td>• Gastrointestinal losses, such as vomiting, diarrhoea</td>
<td>Urine sodium &gt;30 mmol/l, but may be &lt;30</td>
<td>Cirrhosis with ascites</td>
</tr>
<tr>
<td>• Pancreatitis</td>
<td>If dietary access to salt restricted</td>
<td>Nephrotic syndrome</td>
</tr>
<tr>
<td>Renal loss, urine sodium &gt;30 mmol/l</td>
<td>Fluid restriction (&lt;1 litre/day)</td>
<td>Nephrotic syndrome</td>
</tr>
<tr>
<td>• Diuretics</td>
<td>* Demeclocycline (600-1200 mg/day)</td>
<td>Diuretics</td>
</tr>
<tr>
<td>• Salt wasting nephropathy</td>
<td>Intravenous 3% (hypertonic) saline if severe symptoms and of acute onset (&lt;48 h)</td>
<td>Salt wasting nephropathy</td>
</tr>
<tr>
<td>• Cerebral salt wasting</td>
<td>Fluid deprivation, demeclocycline, or both may help</td>
<td>Congestive cardiac failure</td>
</tr>
<tr>
<td>• Mineralocorticoid deficiency (Addison's disease)</td>
<td>TREAT underlying condition</td>
<td>Hypertension</td>
</tr>
</tbody>
</table>

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**Management of hyponatraemia**

As the duration of hyponatraemia may be difficult to judge, the presence of symptoms and their severity should guide the treatment strategy (figure). Acute hyponatraemia developing within 48 hours carries a risk of cerebral oedema, so prompt treatment is indicated with apparently small risk of central pontine myelinoly-
**Box 2: Examination and investigations in patient with hyponatraemia**

**Evaluation of volume status**
- Skin turgor
- Pulse rate
- Postural blood pressure
- Jugular venous pressure
- Consider central venous pressure monitoring
- Examination of fluid balance charts

**General examination for underlying illness**
- Congestive cardiac failure
- Cirrhosis
- Nephrotic syndrome
- Addison's disease
- Hypopituitarism
- Hypothyroidism

**Investigations**
- Urinary sodium
- Plasma glucose and lipids
- Renal function
- Thyroid function
- Peak cortisol during short synacthen test
- Plasma and urine osmolality
- If indicated: chest x-ray, and computed tomography and magnetic resonance imaging of head and thorax

*Pseudohyponatraemia due to artefactual reduction in plasma sodium in the presence of marked elevation of plasma lipids or proteins should no longer be seen with the measurement of sodium by ion specific electrodes; hyperglycaemia causes true hyponatraemia, irrespective of laboratory method.
†May be unhelpful in pituitary apoplexy, in which patients may still "pass" the test.
‡For SIADH: plasma osmolality < 270 mosm/kg with inappropriate urinary concentration (> 100 mosm/kg), in a euvoalaemic patient after exclusion of hypothyroidism and glucocorticoid deficiency.

Hyponatraemia minimises central pontine myelinolysis. Unfortunately, no consensus exists about the optimal rate of correction of hyponatraemia. Although many people advocate a target rate not exceeding 8 mmol/l on any day of treatment, others suggest 12 mmol/l/day or even more if the patient has symptoms—for example, raising the sodium concentration by 1-2 mmol/l per hour until symptoms have resolved, with close monitoring of plasma sodium. The evidence base for using hypertonic saline (3% sodium chloride) in acute symptomatic hyponatraemia is slight, and we recommend that this should be used only after specialist advice has been sought and with frequent (one to two hourly) measurement of plasma sodium. Some authors recommend that a loop diuretic such as furosemide should be given with the hypertonic saline infusion to enhance free water clearance, but caution is needed as this may cause too rapid a rise in sodium.¹³

**New developments for management of hyponatraemia**

Fluid restriction (≤ 1 litre/day) is the initial approach of treating chronic asymptomatic hyponatraemia < 130 mmol/l. Depressingly, no long term trials of the efficacy in practice of this apparently simple approach have been done, but in short term trials it seems to have little effect.¹⁴ ¹⁵ Demeclocycline, which inhibits arginine vasopressin action in the kidney collecting duct, is the current “drug of choice” for treating chronic asymptomatic hyponatraemia due to SIADH if fluid restriction alone does not restore sodium concentrations. Lithium exerts similar renal effects but is less desirable because of inconsistent effects and more side effects (renal impairment, central nervous system effects, thyroid disorders). Urea has been proposed as an alternative option but is poorly tolerated.

The development of orally active antagonists selective for the antidiuretic (renal V2 receptor) action of arginine vasopressin therefore has exciting therapeutic prospects in the management of hyponatraemia. Such "aquaretics" (for example, tolvaptan, lixivaptan) induce a water diuresis without affecting urinary electrolyte or solute excretion. Emerging, if short term, clinical trials have shown the expected effects of aquaresis and correction of hyponatraemia in cirrhosis, heart failure, and SIADH,¹⁶—¹⁸ and the drugs seem to be well tolerated; thirst is the only major side effect reported. Moreover, restriction of fluid intake may not be necessary with these agents.¹⁷ Although V1a (vasoconstrictor) receptor antagonism would not directly affect hyponatraemia, combined V1a/V2 receptor antagonists (such as conivaptan) are in phase III trials after showing promising effects in patients with heart failure in association with hyponatraemia, in which their additional antivasoconstrictor effects seem to helpfully reduce total peripheral resistance and increase cardiac output.¹⁷ Before their possible place in the pantheon of therapeutics can be determined, however, arginine vasopressin receptor antagonists need to show efficacy and lack of toxicity in long term trials with medically important morbidity and mortality end points.

**Hypernatraemia**

Hypernatraemia is much less common than hyponatraemia.²⁴ It reflects a net water loss or a hypertonic...
sodium gain, with inevitable hyperosmolality. Severe symptoms are usually evident only with acute and large increases in plasma sodium concentrations to above 158-160 mmol/L. Importantly, the sensation of intense thirst that protects against severe hypernatraemia in health may be absent or reduced in patients with altered mental status or with hypothalamic lesions affecting their sense of thirst (adipsia) and in infants and elderly people. Non-specific symptoms such as anorexia, muscle weakness, restlessness, nausea, and vomiting tend to occur early. More serious signs follow, with altered mental status, lethargy, irritability, stupor, or coma. Acute brain shrinkage can induce vascular rupture, with cerebral bleeding and subarachnoid haemorrhage.

History, examination, and investigation
Often the cause is evident from the history (box 3). Measurement of urine osmolality in relation to the plasma osmolality and the urine sodium concentration help if the cause is unclear. Patients with diabetes insipidus present with polyuria and polydipsia (and not hypernatraemia unless thirst sensation is impaired). Central diabetes insipidus and nephrogenic diabetes insipidus may be differentiated by the response to water deprivation (failure to concentrate urine) followed by the V2 agonist desmopressin, causing concentration of urine in patients with central diabetes insipidus.

Management
In patients with hypernatraemia that has developed over a period of hours, rapid correction of plasma sodium (falling by 1 mmol/L per hour) improves the prognosis without the risk of convulsions and cerebral oedema.10 Management of a shocked patient needs specialist input and close monitoring, preferably in a high dependency unit. Intravenous normal saline should be used to correct the extracellular fluid depletion, with calculation of the free water deficit to determine how much 5% dextrose to give.11 In patients with hypernatraemia of longer or unknown duration, reducing the sodium concentration more slowly is prudent. Patients should be given intravenous 5% dextrose for acute hypernatraemia or half-normal saline for hypernatraemia of longer or unknown duration, should be used to correct the extracellular fluid depletion, with calculation of the free water deficit to determine how much 5% dextrose to give.11 In patients with hypernatraemia of longer or unknown duration, reducing the sodium concentration more slowly is prudent. Patients should be given intravenous 5% dextrose for acute hypernatraemia or half-normal saline for chronic hypernatraemia if unable to tolerate oral water. Central diabetes insipidus is treated with desmopressin, either as intranasal spray or tablets, with careful monitoring to avoid the complications of water intoxication (delaying one dose each week to allow polyuria and thirst to “breakthrough” in patients susceptible to hyponatraemia with desmopressin may be prudent). Treatment of nephrogenic diabetes insipidus includes removal of precipitating factors, with caution. Acute brain shrinkage can induce vascular rupture, with cerebral bleeding and subarachnoid haemorrhage.

Final thoughts
Despite the frequent occurrence and the poor outcomes of serious disorders of sodium balance, few hard data are available to guide the clinician. This area needs clinical trials, notably of existing approaches to correction of plasma sodium concentration in patients with hypernatraemia: a multicentre, randomised, placebo-controlled trial. Hypertension 2003;42:132-9.

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References
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