Salt, water and nephron: Mechanisms of action and link to hypertension and chronic kidney disease

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SUMMARY AT A GLANCE

- Na⁺ and water regulations are tightly connected, involving neurohumeral multi-organ regulations of AVP, glucocorticoids, RAAS activation, muscle catabolism, urea generation and salt-retention.
- High-salt and low-water intake, common in the general population, is potentially pathogenic and linked to the genesis of hypertension and CKD.
- The tissue (mainly skin and muscle) storage pool of Na⁺ likely participates in Na⁺ and water homeostasis; heavier tissue Na⁺ storage may be pathological and has been associated with resistant hypertension and left ventricular hypertrophy.
- When appropriate, a modest increase (~0.5 -1.5 L/day) in water intake and reduction in dietary salt to a recommended range can blunt AVP increase and will likely be beneficial.

Although classically considered separately, sodium (Na⁺) is associated more with volume status in the body, and water more with body-fluid tonicity, mounting evidence indicates a complex and tight link between Na⁺ and water regulation, involving multiple organ systems. An update on this topic with a focus on kidney-related regulations is provided. The
role of Na\textsuperscript{+}-overload and water insufficiency in the development of hypertension (HTN) and chronic kidney disease (CKD), as well as the potential benefits of dietary modifications, is discussed.

**SALT AND WATER REGULATION, AN UPDATE**

Regulation of salt and water is highly influenced by the circulating levels of vasopressin (arginine vasopressin [AVP], also known as antidiuretic hormone, ADH). AVP is produced by hypthalamus magnocellular neurons (MCN) of the paraventricular and supraoptic nuclei and parvocellular neurons. AVP is not only stored in the posterior pituitary and released upon stimulation (detailed below), but also travels through portal vessels to the anterior pituitary, where AVP stimulates adrenocorticotropic hormone (ACTH) release, thus participating in the regulation of glucocorticoids. Additionally, AVP stimulates cortisol release from the adrenal medulla.\textsuperscript{1,2}

Arginine vasopressin is secreted to the circulation in an equimolar ratio with copeptin, a 39-amino acid glycopeptide, cleaved from the C-terminal part of the AVP precursor. Compared to AVP, copeptin is more stable and easily measurable (median level in healthy adults, <5 pmol/L\textsuperscript{3,4}). Copeptin levels track nicely with the circulating levels of AVP, both at the baseline and in response to stimulations. It is considered a surrogate marker for AVP.

Arginine vasopressin exerts its functions through interaction with its receptors, which are a group of G-protein-coupled (V\textsubscript{1} and V\textsubscript{2}) receptors. V\textsubscript{1} receptors (V\textsubscript{1R}) signal through 1,2-DAG signalling pathways and are further divided into V\textsubscript{1aR} and V\textsubscript{1bR}. V\textsubscript{1aRs} are predominantly expressed in the vascular smooth muscle cells, where they mediate vasoconstriction. V\textsubscript{1aRs} are also expressed in cardiac myocytes, kidney vasa recta, medullary interstitial cells and collecting duct principle cells. Precise functions of V\textsubscript{1aRs} in these systems are being actively investigated. Additionally, V\textsubscript{1aR} participates in habit-forming circadian signals.\textsuperscript{5} V\textsubscript{1bRs}

![Fig. 1](image)

**Fig. 1** (A) and (B) Arginine vasopressin (AVP)-mediated intraglomerular hypertension (HTN), a consequence of combined afferent vasodilation (modulation of tubuloglomerular feedback, TGF) and efferent vasoconstriction (activation of renin angiotensin-aldosterone system, RAAS). AVP positively influences renal sodium absorption at the levels of thick-ascending limb and distal tubules. It also increases urea cycling and intramedullary urea retention by upregulating urea transporters A1 (UT-A1) and A3 (UT-A3) expression. (C) AVP V\textsubscript{2} receptor-mediated regulation of aquaporin 2 (AQP2) and ENaC. Additionally, hypertonicity of extracellular fluids can efficiently activate tonicity response element binding protein (TonEBP) which upregulates AQP2 expression (independent of AVP). TonEBP also upregulates the expression of osmoprotective genes in the renal epithelial cells, critical in preserving medullary structural integrity.
are expressed in the anterior pituitary cells and adrenal medulla, mediating AVP-stimulated ACTH and cortisol release.\(^6\) V2Rs signal through adenylyl cyclase (primarily adenylyl cyclase type 6) signalling pathways. V2Rs are expressed mainly in the kidney collecting ducts, regulating water absorption through controlling aquaporin 2 (AQP2) gene expression and protein distribution (Figure 1A,C). V2R-mediated water absorption is critical to body-fluid toxicity and is functionally non-redundant; loss-of-function and gain-of-function mutations in V2Rs induce clinically significant nephrogenic diabetes insipidus and syndrome of inappropriate ADH, respectively.\(^7\)–\(^9\)

Major physiological signals for the posterior pituitary AVP release are serum osmolality elevation, hypovolemia and circadian rhythmicity. Serum osmolality is an exquisitely sensitive but less potent AVP stimulant. A small rise in osmolality, even within physiological range (280–295 mOsm/kg), can trigger a brisk rise of circulating AVP. Hypovolemia, on the other hand, is a potent but much less sensitive stimulant. Greater than ~7% of volume depletion is needed to trigger AVP release. Once triggered, the rise can be sustained and amounts to a large magnitude. Circadian rhythmicity of AVP, peaking at around midnight, is habit-forming, independent of serum osmolality and volume status. Insufficient AVP release can be associated with enuresis. In 2017, the Food and Drug Administration USA (FDA) approved the clinical use of desmopressin acetate (a selective V2 receptor agonist) for patients with nocturnal polyuria. An habitual V1aR-mediated signal, triggering water-intake prior to sleep in anticipation of a no-water-intake period (sleeping) has also been described.\(^5\)

Arginine vasopressin-mediated water absorption and Na\(^+\) retention are accomplished via V2-mediated effects in the kidneys (Figure 1A,C). In the distal nephrons, through engaging V2 receptors, AVP activates adenylyl cyclase-mediated cyclic AMP/PKA pathways, positively influencing AQP2 gene transcription, mRNA translation and protein translocation to the apical membrane. AVP also participates in the building of intramedullary hypertonicity to facilitate osmotic-gradient-driven water absorption via AQP2. AVP enhances Na\(^+\) absorption at multiple tubular segments including the thick-ascending limb of Henle (NKCC2), distal convoluted tubules (NCC) and collecting ducts (ENaC). It also increases urea cycling and medullary urea retention through upregulating the expression of urea transporters A1 (UT-A1) and A3 (UT-A3).\(^10\) AVP thus not only stimulates water conservation, but also retains Na\(^+\) and urea. The Na\(^+\) retentive effect, demonstrated in both animals and humans,\(^11\) can result in a positive Na\(^+\) balance. Although V1aR-mediated counter V2R force can mitigate the Na\(^+\) retention, the risk for AVP-provoked deleterious effects remains.\(^12\)

In glomeruli, AVP exerts multiple effects (Figure 1A,B). As a result of heightened tubular Na\(^+\) and urea retention induced by AVP, tubular fluids passing through the distal nephrons contain more water/less solutes (Na\(^+\) and urea). Such a fluid composition modulates tubuloglomerular feedback (TGF) at the macula densa, causing afferent arteriole vasodilation. Simultaneously, AVP stimulates renin production in AT-1 receptor-dependent and -independent (via PKA/CREB\(^13\)) manners in the macula densa and in the collecting ducts, respectively. The activation of the renin and its downstream angiotensin-aldosterone system (RAAS) constricts efferent arterioles, resulting in glomerular hyperfiltration.\(^14\) Moreover, AVP has also been shown to stimulate mesangial contraction and proliferation.\(^15\) These collective AVP-mediated effects are consistent with epidemiological data showing concentrated urine (high AVP state) and serum copeptin levels proportionally correlated with the occurrence and severity of proteinuria,\(^16\) a marker of long-term kidney damage.

From an evolutionary view, AVP, mediating water and Na\(^+\) conservation, fits well with limited dietary sources of water and salt. With modernization and industrialization; however, salt has become a readily-available and inexpensive commodity. Our daily salt intake has risen from of 1–3 g by our ancestors to an average of 10–15 g. Coupled with insufficient water intake,\(^17\) such a dietary habit creates a challenge that demands the kidneys to conserve water but dispose of excess salt. The kidneys, in order to conserve water, have to retain extra Na\(^+\) (and urea) to strengthen the intramedullary hypertonicity. During this process, extra Na\(^+\) is being ‘saved’. The surplus of body-fluid Na\(^+\) rises osmolality,\(^18\) stimulating AVP release. It is conceivable that the combined effects of high-salt and low-water intake exert a major role in the epidemics of non-communicable chronic diseases including HTN and CKD (Figure 2).

Na\(^+\) homeostasis has been viewed classically as almost exclusively regulated by the kidneys. This notion has been challenged; however, in light of new evidence showing multiple organ system involvement in Na\(^+\) regulation and the
discovery of a non-osmotic storage pool of Na\textsuperscript{+}, potentially contributing to a unique circaseptan rhythmicity of urine Na\textsuperscript{+} excretion. Valuable information was obtained through examining a simulated space-flight to Mars in which intake and output of healthy adults \((n = 10)\) were precisely recorded. Two studies of 105 and 520 days yielded a tight correlation between salt intake and blood pressure elevation, as well as changes in serum and urine aldosterone (reduction) and free cortisol products (elevation).\textsuperscript{19,20} High-salt diet (12 g/day) raised urine Na\textsuperscript{+} without triggering extra water intake (lacking thirst stimulation). The kidneys achieve the salt-unloading and water-conservation through activating an array of metabolic processes, including an elevated hepatic urea genesis. Moreover, through activating glucocorticoid receptors in the skeletal muscle, high-salt intake accelerates muscle catabolism, presumably providing building blocks and energy equivalents to facilitate hepatic urea genesis.

The discovery of a circaseptan rhythmicity of urine Na\textsuperscript{+} excretion is clinically important. It negates the value of a single 24 h urine collection as the gold-standard for judging Na\textsuperscript{+} intake. In fact, a single 24 h urine study has a mere 50–50 predictive value. It is clear that, depending on the timing in the circaseptan, a single 24 h urine Na\textsuperscript{+} excretion could substantially deviate from one’s Na\textsuperscript{+} intake, despite being in a stable condition.\textsuperscript{21} Urine Na\textsuperscript{+} excretion derived from an average of seven consecutive days yields a more accurate estimation of daily Na\textsuperscript{+} consumption.\textsuperscript{22,23}

With the advent of Na\textsuperscript{+} imaging technique, an osmolality- and volume-independent pool of Na\textsuperscript{+} in the skin (epidermal and dermal regions) and muscle has been revealed,\textsuperscript{24,25} potentially contributing to Na\textsuperscript{+} homeostasis/circaseptan rhythm. Heavy Na\textsuperscript{+} signal in the skin seems deleterious and has been shown to be an independent risk factor for refractory HTN and left ventricular hypertrophy.\textsuperscript{21,26,27} Along the same line, recent studies have linked high-salt intake to vascular inflammation, impairment of endothelial surface glycocalyx layer\textsuperscript{28} and endothelial nitric oxide response, contributing to artery stiffness,\textsuperscript{29} reduced brain oxygen delivery and cognitive dysfunction.\textsuperscript{30}

COMBINED SALT OVERCONSUMPTION AND UNDER-HYDRATION (SUBOPTIMAL WATER INTAKE), CULPRITS FOR HTN AND CKD?

Except for several methodologically-challenged studies,\textsuperscript{31} both animal and human studies have demonstrated a positive correlation between salt intake and blood pressure.\textsuperscript{32,33} Knowledge derived in the last few years from studies of salt-triggered metabolic reactions and maladaptive signalling mechanisms has afforded a better understanding of the underlying pathogenesis. With short-term (<7 days) high-salt intake, the rise in serum osmolality and, hence, AVP, is short-lived. This is because serum osmolality/AVP-mediated vasoconstriction is promptly sensed by baroreceptors, activating inhibitory GABAergic neurons in the hypothalamus and turning off AVP production. Chronic high-salt intake, on the contrary, activates an aberrant pathway involving BDNF/Trk activation, which leads to KCC2 channel inhibition; the latter collapses the transmembrane Cl\textsuperscript{−} gradient, causing the AVP-producing MCN to erroneously respond to the inhibitory GABAergic signal as an AVP stimulatory signal.\textsuperscript{34,35} In essence, sustained high-salt intake turns the negative feedback loop into a positive one, resulting in an exaggerated AVP production and vasoconstriction.\textsuperscript{36,37} A superimposed under-hydration (signified by a high-normal serum Na\textsuperscript{+} concentration) would further augment AVP secretion. Moreover, high-salt intake has been shown to elevate endothelin-1 (ET-1) which participates in the regulation of the skin storage pool of Na\textsuperscript{+}, influencing blood pressure and diurnal pressure rhythm.\textsuperscript{30,38} These results are consistent with the observations of an elevated HTN risk in adults with high plasma copeptin\textsuperscript{37} and a remarkable blood pressure reductive response to long-term dietary salt reduction.\textsuperscript{39}

Turning to the contribution of salt excess and water inadequacy to the risk of kidney dysfunction, the detrimental effects have also been tied to AVP. Studies in 5/6 nephrectomized Brattleboro rats (genetically lacking AVP genes) show that, compared to their wild-type counterparts, these rats exhibit a milder degree of kidney dysfunction.\textsuperscript{40} After given back dDAVP (a V\textsubscript{2} receptor agonist), they developed more severe kidney damage with worsening serum creatinine, proteinuria and anaemia. Similarly, water-drinking lowers AVP and reduces proteinuria and interstitial inflammation.\textsuperscript{41,42} The AVP effects on albuminuria and renal function have also been demonstrated in humans.\textsuperscript{43,44} Due to structural defects in the kidney medulla, ADPKD patients typically exhibit a degree of obligatory polyuria. The water loss, consequently, is associated with higher levels of serum osmolality and AVP (copeptin), especially after fasting. Even with preserved kidney function, an abnormal urine protein excretion is evident.\textsuperscript{44} The pathogenic role of AVP has also been demonstrated in several multicentre prospective interventional trials, showing renal protection with V\textsubscript{2}R blockade in ADPKD.\textsuperscript{45,46}

Arginine vasopressin-mediated kidney dysfunction is not unique to ADPKD. Rather, it exerts negative effects in the non-ADPKD population.\textsuperscript{47} Elevated copeptin in kidney transplant recipients is associated with an accelerated decline of kidney function.\textsuperscript{48} In a prospective study from 2002 to 2008 \((n = 2148)\), concentrated urine (a high AVP state) was significantly associated with a more rapid and progressive loss of kidney function.\textsuperscript{49} Similarly, Ponte et al. examined data from December 2009 to March 2013 \((n = 1128)\), showing that circulating copeptin is positively correlated with the occurrence of proteinuria, declining kidney function and kidney shrinkage.\textsuperscript{50} Recently, in a 5 year prospective study of non-CKD and non-diabetes adults \((n = 12 041)\), Kuwabara et al. show that serum Na\textsuperscript{+} concentration (in the range of 137–147 mmol/L) and calculated...
serum osmolality, despite falling within reference range, are positively associated with a cumulative incidence of CKD (eGFR of <60 mL/min per BSA). A recent multicenter interventional study of CKD stage 3 patients (n = 631) showed a smaller decline in serum creatinine clearance (no significant change in the eGFR) resulting from a one-year increase in self-reported fluid intake by 0.6 to 0.8 L/day; this was associated with a reduction in serum copeptin (-2.2 pmol/L). 

### SUMMARY

Na+ and water regulation is complex, involving neuro-humeral multi-organ regulations of AVP, glucocorticoids, RAAS activation, muscle catabolism, urea generation and salt-retention. Several lines of evidence lend strong support to a pathogenic role of high-salt and low-water intake, common in the general population, in the genesis of HTN and CKD. Moreover, although detailed regulatory mechanisms are yet to be elucidated, the non-osmotic storage pool of Na+ in the skin/muscle likely participates in the Na+ and water homeostasis; heavier tissue Na+ storage may be pathological and has recently been suggested to be a risk for resistant HTN and left ventricular hypertrophy. Given even a modest increase (~0.5–1.5 L/day) in water intake and reduction in dietary salt can blunt increments of AVP and blood pressure, public education to raise awareness and promote these preventive measures would be cost effective and beneficial.

### DISCLOSURE

The author has no conflict of interest to report.

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