Specific Na\(^+\) Sensors Are Functionally Expressed in a Neuronal Population of the Median Preoptic Nucleus of the Rat

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Whole-cell patch-clamp recordings were performed on acute brain slices of male rats to investigate the ability of the neurons of the median preoptic nucleus (MnPO) to detect fluctuation in extracellular osmolarity and sodium concentration ([Na\(^+\)]\(_{\text{out}}\)). Local application of hypotonic and hypertonic artificial CSF hyperpolarized and depolarized the neurons, respectively. Similar responses obtained under synaptic isolation (0.5 mM TTX) highlighted the intrinsic ability of the MnPO neurons to detect changes in extracellular osmolarity and [Na\(^+\)]\(_{\text{out}}\). Manipulating extracellular osmolarity, [Na\(^+\)]\(_{\text{out}}\) and [Cl\(^-\)]\(_{\text{out}}\) showed in an independent manner that the MnPO neurons responded to a change in [Na\(^+\)]\(_{\text{out}}\) exclusively. The specific Na\(^+\) response was voltage insensitive and depended on the driving force for Na\(^+\) ions, indicating that a sustained background Na\(^+\) permeability controlled the membrane potential of the MnPO neurons. This specific response was not reduced by Gd\(^3+\), amiloride, or benzamil, ruling out the participation of mechanosensitive cationic channels, specific epithelial Na\(^+\) channels, and Phe-Met-Arg-Phe-gated Na\(^+\) channels, respectively. Combination of in situ hybridization, using a riboprobe directed against the atypical Na\(^+\) channel (Na\(_X\)), and immunohistochemistry, using an antibody against neuron-specific nuclei protein, revealed that a substantial population of MnPO neurons expressed the Na\(_X\) channel, which was characterized recently as a concentration-sensitive Na\(^+\) channel. This study shows that a neuronal population of the MnPO acts as functional Na\(^+\) sensors and that the Na\(_X\) channel might represent the molecular basis for the extracellular sodium level sensing in these neurons.

Key words: lamina terminalis; sodium homeostasis; salt appetite; sodium sensors; hydromineral homeostasis; Na\(_X\); sodium channel

Introduction

Constant physiological osmolarity of plasma and CSF requires detectors that continuously monitor sodium ion concentration ([Na\(^+\)]) and hydration within these extracellular fluids. One major challenge in neurobiology of the hydromineral homeostasis is to reveal the existence, at the level of the brain, of specific sensors involved in the control of the water and mineral balance. Fluctuations in plasma and CSF [Na\(^+\)] not only trigger short-term adjustments, such as the release of antidiuretic and natriuretic hormones from the neurohypophysis (Bourque et al., 1994; Hussy et al., 2000; Voisin and Bourque, 2002), but also long-term regulation that controls thirst and specific appetite for salt (Weisgerger et al., 1979; Denton et al., 1996). It has been hypothesized that specific brain Na\(^+\) sensors initiate sodium intake (Weisgering et al., 1979; Denton et al., 1996) as well as natriuresis (Cox et al., 1987; Denton et al., 1996), and the recent discovery of coincident detectors of extracellular fluid osmolarity and [Na\(^+\)] in the supraoptic nucleus (SON) established the cellular basis for Na\(^+\) detection in this nucleus (Voisin et al., 1999; Voisin and Bourque, 2002). However, physiological experiments indicated the presence of Na\(^+\) sensors in the periventricular region of the brain (Cox et al., 1987; Park et al., 1989; Denton et al., 1996), and recent molecular biology data combined with behavioral studies revealed that Na\(^+\) sensors located in the preoptic region were associated with salt intake (Watanabe et al., 2000; Hiyama et al., 2002). The location of the median preoptic nucleus (MnPO) along the third ventricle, as well as the absence of tight junctions between ciliated cells that form the ventricular ependyma separating the CSF from the MnPO, make this nucleus a strategic locus for detection of CSF [Na\(^+\)] (Landas and Philipp, 1987; McKinley et al., 1987; Oldfield and McKinley, 1995). Functionally, neuronal subpopulations of the MnPO have been shown to display an osmosensitive profile (Honda et al., 1990; Travis and Johnson, 1993; Aradachi et al., 1996). However, the influence of osmotic challenges has been controversial (Honda et al., 1990; Travis and Johnson, 1993), and the intrinsic ability of these neurons to detect changes in environmental osmolarity and [Na\(^+\)] has yet to be determined. Moreover, the adjustment of the extracellular osmolarity with NaCl did not distinguish between the presence of osmoreceptors or Na\(^+\) sensors in this nucleus. Using a hypothalamic slice preparation and manipulating the extracellular osmolarity and [Na\(^+\)] separately, we investigated the possibility that neurons of the MnPO might act as intrinsic...
Table 1. Composition of the different aCSFs used in the study

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<th>Hypotonic acSF</th>
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acSF of different composition and osmolarity were transiently and locally applied over the region of the ventral MnPO to investigate how extracellular osmolarity and extracellular concentration of sodium and chloride ions might alter the excitability of the ventral MnPO neurons. After pH adjustment with NaOH (10 mM), aCSF osmolarity was measured with a vapor pressure osmometer, and mannitol was added to aCSF, when necessary. Note that final aCSF [Na⁺] was corrected accordingly.

To study the effect of osmotic and natriuretic challenges on the excitability of MnPO neurons, modified aCSF containing different concentrations of Na⁺ and Cl⁻ ions, as well as different osmolarity, was delivered in the vicinity of the MnPO by using a rapid solution changer and manifold (model RSC-160; Bio-Logic). aCSF osmolarity was assessed by using a vapor pressure osmometer (Wescor), and adjustment to the appropriate osmolarity values was performed by adding mannitol to aCSF after pH adjustment, when necessary. Composition and osmolarity of the various aCSF used for that study are detailed in Table 1.

Drugs expected to affect neurons of the vMnPO in a steady-state manner were directly added to the extracellular solution and bath applied. Gadolinium and TTX, as well as amiloride and benzamil, were purchased from Sigma (St. Louis, MO).

Statistical analysis. All results are expressed as means ± SEM. Statistical comparisons were performed using paired Student’s t test for absolute values (cell input resistance); p < 0.05 was considered significant. Results expressed as percentages (pharmacological data) were considered to be nonparametric data. They were analyzed using the Mann–Whitney U test. Statistical significance was determined at p < 0.05.

Immunohistochemistry and in situ hybridization protocols. A combination of immunohistochemistry (IHC) for neuron-specific nuclei protein (NeuN) and in situ hybridization (ISHH) immunohistochemistry for NaX mRNA was performed to allow simultaneous visualization of NeuN immunoreactivity and NaX mRNA in the same tissue sections that contained the MnPO. To match the anatomical observations with the electrophysiological recordings, IHC and ISHH experiments were performed on rats of the same age as those used for the electrophysiological study (100–150 gm). Brain sections (30 µm) were processed using the avidin–biotin bridge method with hydrogen peroxide as a substrate. The
HIC reaction was performed in RNase-free condition using a mouse monoclonal antibody (MAB377; Chemicon, Temecula, CA) that recognized NeuN at a concentration of 1:5000. After the HIC procedure, the brain sections were processed for ISHH using a 35S-labeled atypical Na\(^+\) channel Na\(_{v}\) cRNA probe for mRNA. The Na\(_{v}\) cRNA probe was generated from a rat cDNA containing in a pBS(SK\(^-\)) plasmid from Stratagene (La Jolla, CA) (a kind gift from Dr. Waxman, Yale University, West Haven, CT) (Black et al., 1994). The insert corresponds to the cDNA sequence encoding a portion of domain 4: nucleotides 880–1308 of the sequence M96578 (Gautron et al., 1992) or nucleotides 4671–5119 of the sequence Y09164 (Akopian et al., 1997). To produce the antisense probe, DNA was digested with EcoRI and T7 polymerase. Radiolabeled sense RNA copies (control) were also prepared to verify the specificity of Na\(_{v}\) probe. To produce the sense probe, DNA was digested in a mixture containing sterile KPBS, the chromagen 3,3′-diaminobenzidine tetrahydrochloride (0.5 mg/ml), and hydrogen peroxide (0.003%). Thereafter, hybridization histochemical localization of Na\(_{v}\) mRNA was performed using a 35S-labeled Na\(_{v}\) cRNA probe. Protocols for riboprobe synthesis, hybridization, and autoradiographic localization of Na\(_{v}\) mRNA signal were adapted from Simmons et al. (1989) and Dumont et al. (1999). Immunoreactive sections for NeuN were mounted onto poly-l-lysine-coated slides and were dessicated under vacuum overnight, fixed in 4% paraformaldehyde for 30 min, and digested by proteinase K (10 \(\mu\)g/ml in 100 mM Tris HCl, pH 8.0) and 50 \(\mu\)M EDTA, at 37°C for 25 min. The brain sections were then rinsed in sterile DEPC water, followed by a solution of 100 mM triethanolamine (TEA), pH 8.0, acetylated in 0.25% acetic anhydride in 100 mM TEA, and cleared in xylene, and coverslipped with a mixture of distyrene, tricresyl phosphate, and xylene (DPX).

Results
In the first set of experiments, vMnPO neurons were maintained around their resting potential (range, –63 to –60 mV) or were slightly depolarized with current injection to generate a regular spiking activity (range, –55 to –45 mV). Baseline for stable membrane potential was established for 2 to 4 min under isotonic aCSF. Hypotonic aCSF (Table 1, column 1) was then locally applied for 1 min over the region of the vMnPO. Transient application of the hypotonic solution triggered membrane hyperpolarization, with a concomitant abolition of the firing activity in cells that were sufficiently depolarized to fire spikes during the control period. Membrane hyperpolarization amounted to 6.0 ± 0.4 mV (\(n = 20\)) and was fully reversed after 4–5 min. Local application of hypertonic aCSF (Table 1, column 2) produced a transient depolarization of the cell (3.9 ± 0.2 mV; \(n = 22\)) that was often accompanied by a sustained discharge of action potentials. Neurons returned to their initial membrane potential after reexposure to isotonic condition, and the action potentials stopped. As illustrated in Figure 1A, individual neurons of the vMnPO were able to respond to opposite changes in extracellular tonicity. In 6 of the 20 neurons that displayed a hypotonicity-induced hyperpolarization (top trace), a subsequent application of a hypertonic stimulus was tested after the recovery period. In all of the neurons tested, the hypertonic aCSF induced a transient depolarization (bottom trace). The ability of vMnPO neurons to respond to consecutive hypertonic stimuli was then tested. In 11 neurons of the vMnPO held at −60 mV, two consecutive local applications of hypotonic aCSF hyperpolarized the neurons by 7 ± 0.9 and 7 ± 0.6 mV, respectively. Furthermore, in four other neurons, three consecutive hypertonic challenges induced membrane hyperpolarization amounting to 6.9 ± 1.1, 6.9 ± 1.1, and 7.8 ± 0.9 mV, respectively.

The application of hypotonic or hypertonic aCSF was done locally, i.e., over the MnPO area. However, spillover of the modified aCSF over the adjacent region, which could modify the excitability of upstream neuronal cells projecting to the vMnPO neurons, could not be excluded. Therefore, the intrinsic ability of the MnPO neurons to detect changes in aCSF tonicity was verified in a series of experiments in which synaptic transmission was fully abolished by perfusing a potent blocker of the voltage-gated Na\(^+\) channel within the recording chamber (0.5 \(\mu\)M TTX). Under this steady-state condition, local application of hypotonic and hypertonic stimuli hyperpolarized and depolarized the MnPO neurons by 5.5 ± 0.5 mV (\(n = 7\)) and 3.9 ± 0.4 mV (\(n = 5\)), respectively (Fig. 1B). Consecutive applications of hypertonic and hypotonic aCSF were tested in three neurons, and all of them displayed hyperpolarization and depolarization in response to the appropriate stimulus.

Of a total of 75 neurons tested under standard condition and in the presence of TTX, 72% of the neurons (\(n = 54\)) showed a significant change in membrane potential in response to the local change in extracellular tonicity; these were considered to be osmoresponsive. The remaining neurons did not respond to osmotic stimuli, as illustrated in Figure 1C, for a hypertonic challenge; these were considered to be non-osmoresponsive. The specificity of the MnPO to contain osmoresponsive neurons was then verified. Ten neurons located in the median septum, a region rostral to the MnPO in the sagittal brain slice, were tested for similar changes in extracellular tonicity. These neurons were either maintained around their resting membrane potential or slightly depolarized with current injection. None of these neurons showed a membrane hyperpolarization or depolarization in response to local application of a hypotonic or a hypertonic stimulus, respectively. A typical example of a non-osmoresponsive septal neuron is illustrated in Figure 1D.

vMnPO neurons detect [Na\(^+\)]\(_{out}\) rather than extracellular osmolarity
Changing the extracellular Na\(^+\) salt concentration ([Na\(^+\)]\(_{out}\)) to alter the osmotic pressure suggests that the osmoresponsive vMnPO neurons were able to detect either variation of the extracellular osmolarity or the extracellular concentration of Na\(^+\) and/or Cl\(^-\) ions ([Na\(^+\)]\(_{out}\) and [Cl\(^-\)]\(_{out}\), respectively). In the
next series of experiments, we investigated the possibility that the vMnPO neurons acted as osmoreceptors by detecting alteration of the extracellular osmolarity without changing the [NaCl]_{out}. To test this hypothesis, we applied a hyperosmotic stimulus while maintaining [Na\(^+\)]_{out} at 150 mM, the standard concentration found in the extracellular environment of central neurons. Local application of hyperosmotic–isonatriuric solution (Table 1, column 3) on vMnPO neurons maintained at their resting potential did not change the membrane potential of the neuron (n = 5) (Fig. 2A, top trace). It should be noted that the effect of a hyperosmotic–isonatriuric aCSF was not tested because aCSF containing 150 mM NaCl could not be hypo-osmotic. In the next series of experiments, we varied [Na\(^+\)]_{out} while keeping constant the osmolarity of the aCSF that was directly applied over the vMnPO region. Neurons of the vMnPO were maintained at their resting potential or slightly depolarized to generate action potential discharges. Decreasing [Na\(^+\)]_{out} from 150 to 100 mM while maintaining osmolarity at 300 mOsm/l (by adding D-mannitol) (Table 1, column 4) induced a reversible membrane hyperpolarization of 6.2 ± 0.3 mV (n = 36) (Fig. 2A, middle trace). The amplitude of the hyperpolarization was similar to that which was obtained with a hypo-osmotic–isonatriuric (hypotonic) aCSF (6.0 ± 0.4 mV; n = 20) (Fig. 2A, bottom trace). This observation suggests that osmolarity had no additional effect on the neuronal response induced by hyponatremia. It should be noted that the effect of increased [Na\(^+\)]_{out} in iso-osmotic solution was not investigated, because [Na\(^+\)]_{out} >150 mM was always associated with hyperosmolarity.

This series of results obtained by varying osmolarity and [Na–Cl]_{out} independently indicated that vMnPO neurons did detect a change in [NaCl]_{out} rather than a change in extracellular osmolarity. In our experimental protocol, varying [Na\(^+\)]_{out} induced a concomitant change in [Cl\(^-\)]_{out}. We tested the possibility that the vMnPO neurons responded to a change in [Cl\(^-\)]_{out} rather than to a change in [Na\(^+\)]_{out}. First, hypotonic–isochloride aCSF (Table 1, column 5) was obtained by adding 50 mM choline chloride to the hypotonic aCSF. As illustrated in Figure 2B (top trace), local application of the hypotonic–isochloride aCSF induced a hyperpolarization that amounted to 6.6 ± 0.4 mV (n = 4), a response similar to the one obtained previously with a low Na\(^+\) and low Cl\(^-\) hypotonic aCSF (6.3 ± 0.3 mV). Second, local application of a hypotonic–isochloride aCSF (Table 1, column 6) depolarized the vMnPO neurons by 0.7 ± 0.3 mV (n = 5) (Fig. 2B, bottom trace). In contrast, raising [Cl\(^-\)]_{out} without changing [Na\(^+\)]_{out} (Table 1, column 7) had no effect on the membrane potential of the vMnPO neurons (n = 5). These results show that alteration of the [Na\(^+\)]_{out} did influence the membrane potential of the vMnPO neurons, indicating that these neurons were able to detect a change in [Na\(^+\)]_{out}.

**Cellular mechanisms underlying detection of CSF [Na\(^+\)]**

If Na\(^+\) ions are the major current carrier associated with the observed changes in membrane potential of vMnPO neurons, iso-osmotic variation in [Na\(^+\)]_{out} might change the membrane potential with graded magnitude, revealing the Na\(^+\) specificity of

hypotonic aCSF did not alter the membrane potential of this vMnPO neuron, which was considered as a non-osmoreceptive neuron. D. Local application of hypotonic aCSF did not change the membrane potential of the neurons located in a region adjacent to the MnPO, the median septum. Note that spike amplitude has been truncated in the traces presented here. These results indicate that detection of the extracellular tonicity involved intrinsic properties of a specific neuronal population of the vMnPO. HP, Holding potential.

Figure 1. Intrinsic osmosensitivity is a unique property of the MnPO neurons. A, Depolarization of a ventral MnPO neuron with current injection triggered a regular spiking activity. Local and transient (1 min) application of hypotonic aCSF (270 mOsm/l; 100 mM NaCl) over the vMnPO region hyperpolarized the neuron with a concomitant abolition of the spike discharges (top trace). The same neuron was held below the spike threshold, and local application of hypotonic aCSF (330 mOsm/l; 170 mM NaCl) depolarized the neuron that discharged a prolonged burst of spikes (bottom trace). Details of the spiking activity was illustrated before (*) and after (**) application of the modified aCSF. B, TTX (0.5 μM) was bath applied to block synaptic inputs onto the recorded vMnPO neuron. Under this steady-state synaptic blockage, transient and local application of a hypotonic aCSF hyperpolarized the neuron (top trace). In contrast, local application of a hypertonic aCSF induced a depolarization (bottom trace). C, Local application of
the response. As illustrated in Figure 3A, local application of different iso-osmotic–hypoxynatriuric stimuli (300 mOsm/l; 142, 130, and 100 mM NaCl) hyperpolarized the vMnPO neurons by 3.2 ± 0.3 mV (n = 5), 4.1 ± 0.3 mV (n = 4), and 6 ± 0.4 mV (n = 20), respectively. In addition, the hyperpolarization induced by local application of iso-osmotic–hypoxynatriuric aCSF was not accompanied with a change in input resistance (1.35 ± 0.15 GΩ under control and 1.37 ± 0.15 GΩ under the iso-osmotic–hypoxynatriuric aCSF; n = 12; p = 0.18) (Fig. 3B). Similarly, no change in input resistance was observed during the response to hypernatremia (1.33 ± 0.17 GΩ in control vs 1.33 ± 0.18 GΩ under hypoxynatriuric aCSF; n = 8; p = 0.83) (Fig. 3C).

In the next series of experiments, we used the voltage-clamp mode to further characterize the cellular mechanism(s) underlying detection of CSF [Na⁺]. vMnPO neurons were clamped at their resting potential (−60 mV). In the presence of TTX (0.5 μM) and TEA-Cl (10 mM), local application of iso-osmotic–hypoxynatriuric aCSF triggered a reversible outward current that amounted to +5.9 ± 0.4 pA (n = 26) (Fig. 4A). Intensity-to-voltage relationships obtained with a ramp protocol ranging from −100 to −10 mV were tested before, during, and after the application of the iso-osmotic–hypoxynatriuric aCSF (rate of change fixed at 16 mV/sec; n = 13) (Fig. 4B). To test the action of an iso-osmotic–hypoxynatriuric aCSF, vMnPO neurons were bathed with the iso-osmotic–hypoxynatriuric aCSF (300 mOsm/l; 100 mM NaCl) for 5–10 min (control). Local application of iso-osmotic–hypoxynatriuric aCSF (300 mOsm/l; 150 mM NaCl) compared with control induced a reversible inward current of −5.7 ± 0.4 pA (n = 9) (Fig. 5A). Similar intensity-to-voltage relationships were triggered before, during, and after the application of the iso-osmotic–hypoxynatriuric aCSF (Fig. 5B). These two experimental protocols, performed on the same vMnPO neurons when possible (n = 6), allowed the isolation of the Na⁺-evoked current, which showed no voltage dependency between −100 and −10 mV (Figs. 4C, 5C). The inward direction of the current observed during application of iso-osmotic–hypoxynatriuric aCSF materialized influx of Na⁺ ions within the neuron through a background permeability, whereas the outward current observed during application of iso-osmotic–hypoxynatriuric aCSF reflected a reduction of an Na⁺ influx.

The presence of such a constitutive Na⁺ permeability was investigated in the dorsal part of the MnPO. Ten neurons located above the anterior commissure were bathed in an iso-osmotic aCSF, and transient application of iso-osmotic–hypoxynatriuric aCSF triggered an outward current in 7 of the 10 cells tested (70%), the amplitude of which amounted to +4.7 ± 1.2 pA (data not shown).

Finally, we investigated the pharmacology of the ionic channels underlying the intrinsic background Na⁺ permeability of MnPO neurons. Na⁺ ion flow through stretch-inactivated channels (SICs) has been shown to constitute the cellular basis for depolarizing current to make the neuron fire (holding potential, −55 mV). The bottom trace was obtained from a neuron maintained around its resting potential (holding potential, −60 mV). In these neurons, transient and local application of hyperpolaric–isochloride aCSF (270 mOsm/l; 100 mM NaCl and 50 mM choline chloride) drove the membrane potential to hyperpolarization. C, Two neurons maintained at different membrane potential (holding potential, −55 mV) (top trace); holding potential, −63 mV (bottom trace). Local application of hyperpolaric–isochloride aCSF (330 mOsm/l; 150 mM NaCl and 20 mM Na⁺-gluconate) onto these neurons triggered depolarization of the cells. Spike amplitude has been truncated in all of the panels presented. These results show that neurons of the vMnPO did respond to a change in extracellular (Na⁺) and not to an alteration of the extracellular osmolarity or [Cl⁻]. HP, Holding potential.

Figure 2. Ventral MnPO neurons detect changes in extracellular Na⁺ level but not osmolarity. A, Local application of a hyperosmotic–isochloride stimulus (330 mOsm/l; 150 mM NaCl) had no apparent effect on the excitability of the vMnPO neuron (top trace). In contrast, local application of an iso-osmotic–hypoxynatriuric aCSF (300 mOsm/l; 100 mM NaCl) hyperpolarized the vMnPO neuron (middle trace). The amplitude of the hyperpolarization was similar to the one obtained with a hypotonic aCSF (270 mOsm/l; 100 mM NaCl; bottom trace). B, Traces obtained from two different vMnPO neurons. The top trace was obtained with slight injection of hypotonic aCSF; the amplitude of the hyperpolarization was similar to the one obtained with a hypotonic aCSF (270 mOsm/l; 100 mM NaCl; bottom trace).
Na⁺/H⁺ sensing in the supraoptic neuroendocrine cells (Voisin et al., 1999). Because the open probability of these channels was greatly reduced by the lanthanide trivalent gadolinium (Gd³⁺), we investigated the involvement of mechanosensitive channels in the

Figure 3. The specific response to a natriuric challenge depended on the driving force for Na⁺ ions and was not associated with a change in input resistance. A. In the same vMnPO neuron, local application of iso-osmotic–hyponatriuric aCSF with a reduction of 8, 20, and 50 mOsm/l in the [Na⁺]o compared with regular aCSF (150 mM) induced a cellular response of graded amplitude. B. Local application of iso-osmotic–hyponatriuric aCSF (300 mOsm/l; 100 mM NaCl) triggered an outward current. C. The specific current (Δ current) evoked by hypotremia was isolated by digital subtraction of the I–V relationships recorded in A and B. Note the absence of voltage dependency over the range of potentials tested. TTX (0.5 μM) and TEA-Cl (20 mM) were present in isotonic aCSF, as well as in iso-osmotic–hyponatriuric aCSF, to block voltage-gated Na⁺ and K⁺ currents. For these experiments, extracellular osmolarity (300 mOsm/l) was achieved with or without adding mannitol to aCSF (iso-osmotic–hyponatriuric aCSF and control, respectively).

C, Mean ramp current obtained from the protocol described in B (n = 6 MnPO neurons), showing a parallel shift of the current in the outward direction during transient application of iso-osmotic–hyponatriuric aCSF (top). Mean control current is represented by open circles, and mean current recorded during application of modified aCSF is represented by filled circles. Digital subtraction of the traces shown in the top panel resolved the I–V relationship of the mean current evoked by application of iso-osmotic–hyponatriuric aCSF (bottom). HP, Holding potential.

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Figure 4. Transient application of iso-osmotic–hyponatriuric aCSF on vMnPO neurons triggered a voltage-independent outward current. A. A vMnPO neuron was recorded under the voltage-clamp mode and maintained at a potential of −60 mV. Local application of iso-osmotic–hyponatriuric aCSF (300 mOsm/l; 100 mM NaCl) triggered an outward current. B. Three I–V relationships from −100 to −10 mV (16 mV/sec) were elicited under control (label A), iso-osmotic–hyponatriuric aCSF (label B), and back to iso-osmotic–isonatriuric aCSF (label C). The specific current (Δ current) evoked by hypotremia was isolated by digital subtraction of the I–V relationships recorded in A and B. Note the absence of voltage dependency over the range of potentials tested. TTX (0.5 μM) and TEA-Cl (20 mM) were present in isotonic aCSF, as well as in iso-osmotic–hyponatriuric aCSF, to block voltage-gated Na⁺ and K⁺ currents. For these experiments, extracellular osmolarity (300 mOsm/l) was achieved with or without adding mannitol to aCSF (iso-osmotic–hyponatriuric aCSF and control, respectively).

C, Mean ramp current obtained from the protocol described in B (n = 6 MnPO neurons), showing a parallel shift of the current in the outward direction during transient application of iso-osmotic–hyponatriuric aCSF (top). Mean control current is represented by open circles, and mean current recorded during application of modified aCSF is represented by filled circles. Digital subtraction of the traces shown in the top panel resolved the I–V relationship of the mean current evoked by application of iso-osmotic–hyponatriuric aCSF (bottom). HP, Holding potential.
Figure 5. Transient application of iso-osmotic–hypernatremic aCSF on vMnPO neurons triggered a voltage-independent inward current. A, A vMnPO neuron was maintained at a membrane potential of −60 mV. Local application of iso-osmotic–hypernatremic aCSF (300 mOsm/l; 150 mM NaCl) compared with control solution (300 mOsm/l; 100 mM NaCl) triggered an inward current. B, I–V relationships from −100 to −10 mV (16 mV/sec) were elicited before (label A), during (label B), and after (label C) the application of iso-osmotic–hypernatremic aCSF. The specific current (Δ current) evoked by hypernatremia was isolated by digital subtraction of the I–V relationships recorded in A and B. TTX (0.5 μM) and TEA-Cl (20 mM) were present in control aCSF, as well as in iso-osmotic–hypernatremic aCSF to block voltage-gated Na⁺ and K⁺ currents. For these experiments, extracellular osmolarity (300 mOsm/l) was achieved by adding mannitol to the control aCSF and by removing glucose from the iso-osmotic–hypernatremic aCSF. C, Mean ramp current obtained from the protocol described in B and obtained from the six MnPO neurons illustrated in Figure 4C. Transient application of iso-osmotic–hypernatremic aCSF evoked an inward shift of the current (top). Mean current recorded under control is represented by open circles, and mean current recorded during application of modified aCSF is represented by filled circles. Digital subtraction of the traces shown in the top panel resolved the I–V relationship of the mean current evoked by application of iso-osmotic–hypernatremic aCSF (bottom). HP, Holding potential.

response induced by hypotonic (Table 1, column 1) and hyper- tonic (Table 1, column 2) stimuli. In the presence of TTX (0.5 μM), local application of hypotonic and hyper tonic aCSF hyperpolarized and depolarized the vMnPO neurons, respectively, as demonstrated previously. After recovery, Gd³⁺ (100 μM) was bath applied for 5–6 min before a second local application of hypotonic and hypertonic aCSF that also contained 100 μM Gd³⁺. The amplitude of both the hyperpolarization and depolarization was not reduced in the presence of Gd³⁺ (Fig. 6A), and the percentage of the responses induced by hypotonic and hyper tonic aCSF remained unchanged (99.9 ± 6.7% in control vs 109.1 ± 7.1% in the presence of Gd³⁺; n = 12; p > 0.99). Then, we tried to antagonize the responses by using blockers for specific Na⁺ channels that are constitutively active. Bath application of amiloride (3 μM) remained without effect on the hyperpolarization induced by hypotonic aCSF (Fig. 6B), thus unaffecting the percentage of the response (99.9 ± 9.6% in control vs 104 ± 8.7% in the presence of amiloride; n = 8; p > 0.99) (Fig. 6C). In addition, bath application of benzamil (30 μM), an analog of amiloride, did not reduce the amplitude of the hypotonicity-induced response, for which the percentage remained constant (99.9 ± 10.9% in control vs 99.9 ± 9.1% in the presence of benzamil; n = 3; p > 0.99) (Fig. 6C). Similar inefficacy of these molecules was also observed for the depolarization induced by hypertonic aCSF.

Together, our results indicate that neurons of the MnPO dis-
play a sustained permeability through which Na\(^+\) flux depends on the electrochemical gradient of Na\(^+\) ions. The pharmacological profile of the sustained Na\(^+\) permeability remains, however, to be determined.

**Neurons of the MnPO express the atypical Na\(^+\) channel Na\(_X\)**

Recently, the atypical Na\(^+\) channel Na\(_X\) has been demonstrated to act as a concentration-sensitive Na\(^+\) channel in dissociated subfornical neurons (Hiyama et al., 2002). Despite that the pharmacology of the Na\(_X\) current has not been determined yet, this current shares numerous properties with the sustained Na\(^+\) current that we report here. We, therefore, performed *in situ* hybridization experiments using a riboprobe directed against the Na\(_X\) mRNA to possibly identify the Na\(^+\) background permeability to the Na\(_X\) channel. Results obtained with *in situ* hybridization indicated that Na\(_X\) mRNA are expressed in discrete regions of the hypothalamus. As illustrated in Figure 7A, Na\(_X\) mRNA are strongly expressed in the subfornical organ (SFO), the neurons of which have been demonstrated to express Na\(_X\) channels (Watanabe et al., 2000; Hiyama et al., 2002). Note that *in situ* hybridization performed with the sense riboprobe did not reveal any staining in the SFO region. These results, therefore, did validate the specificity of the riboprobe used in the present study. More importantly, our data revealed that Na\(_X\) mRNA are constitutively expressed in the MnPO (0.18 ± 0.03 μCi/gm; *n* = 7) (Fig. 7B), in which we identified neuronal populations that act as genuine Na\(^+\) sensors. Interestingly, the SON that contains coincident detectors of CSF [Na\(^+\)] and osmotic pressure (Voisin et al., 1999) did not show any staining that is indicative of Na\(_X\) mRNA-containing cells (data not shown). This latter result indicates that neuroendocrine cells of the SON did not express detectable levels of the atypical Na\(^+\) channel mRNA.

The atypical Na\(^+\) channel Na\(_X\) has been first cloned from rat cortical astrocytes and then in neurons (Gautron et al., 1992). To determine whether the Na\(_X\) channel gene is encoded by neuronal populations in the MnPO, we performed immunohistochemistry for NeuN combined with *in situ* hybridization immunohistochemistry for Na\(_X\) mRNA to allow simultaneous visualization of NeuN immuno-reactivity and Na\(_X\) mRNA in the same tissue sections that contained the MnPO (Fig. 8). We found prominent double-
labeled cells localized in both the dorsal and ventral portions of the MnPO, clearly indicating that neurons of the MnPO express the \( \text{Na}_\text{C} \) channel (Fig. 8A, D, E). However, adjacent brain regions, such as the septum (Fig. 8C) or the bed nucleus of the stria terminalis (Fig. 8G, H), did not show any apparent \( \text{Na}_\text{mRNA} \)-labeled neurons.

**Discussion**

The present study indicates that fluctuations in extracellular \([\text{Na}^+]_{\text{out}}\) directly affect the excitability of a neuronal population of the MnPO by changing its membrane potential. The cellular mechanism involved in \([\text{Na}^+]_{\text{out}}\) detection was based on a background \( \text{Na}^- \) permeability active at the resting membrane potential. This permeability is likely identified to the \( \text{Na}_\text{C} \) channel, as strongly suggested by the expression of this atypical \( \text{Na}^- \) channel by MnPO neuronal population.

**Osmoreceptors or specific Na\(^+\) sensors in the MnPO?**

*In vivo*, extracellular electrophysiological recordings showed that a subset of MnPO neurons projecting to the supraoptic neuroendocrine cells increased their firing rate during systemic injection of hypertonic saline (McKinley et al., 1992; Aradachi et al., 1996) and during application of hypertonic saline directly into the MnPO (Honda et al., 1990). These studies certainly demonstrated the osmoreponsiveness of a population of MnPO neurons projecting to the SON and suggested the osmosensitivity of these neurons. Here we showed that, under complete synaptic isolation of the ventral region of the MnPO, extracellular hyperosmotic and hypotonicity depolarized and hyperpolarized a majority of neurons, highlighting the innate ability of a single vMnPO neuron to respond to opposite changes in the extracellular osmolarity (osmosensitivity). Although our results were in agreement with previous *in vivo* studies (Honda et al., 1990; McKinley et al., 1992; Aradachi et al., 1996), they were in contradiction, at least partially, with *in vitro* results reporting that hyperosmolality was associated with a decreased spike activity in few vMnPO neurons (Travis and Johnson, 1993). However, these authors also reported that hypo-osmolality decreased, or increased the electrical activity, depending on the vMnPO neuron tested. These contrasting data might arise from the presence of two separate neuronal populations in the vMnPO that respond to a similar osmotic challenge with an opposite change in their membrane potential. The effects of consecutive applications of hyponatriuric and hypernatriuric aCSF in a significant number of neurons have only revealed a stereotyped response: hyponatriuric aCSF always triggered membrane hyperpolarization, or outward current, whereas hypernatriuric aCSF triggered depolarization, or inward current. These results almost rule out the presence of two different populations with opposite capabilities. Alternatively, different vMnPO neurons might receive opposite synaptic inputs from upstream osmosensitive neurons. Application of hyperosmotic or hypotonic aCSF was done locally, i.e., over the vMnPO region, thus limiting the stimulation of upstream osmosensitive neurons. Moreover, neuronal responses were reproduced under synaptic isolation, excluding opposite synaptic inputs to the vMnPO.

Determining whether vMnPO neurons, as well as neurons located in the dorsal part of the nucleus, were osmoreceptors or \( \text{Na}^- \) sensors was of primary interest in the characterization of the role of the MnPO in hydromineral homeostasis. In the previous studies, the osmotic challenge was always achieved by manipulating \([\text{Na}^+]_{\text{out}}\) in either the extracellular solution (Travis and Johnson, 1993) or the extracellular fluid compartment (Honda et al., 1990). Here, we independently manipulated \([\text{Na}^+]_{\text{out}}\) and extracellular osmolality to evaluate the ability of the vMnPO neurons to discriminate between aCSF osmolality and aCSF \([\text{Na}^+]\). Our results showed that local application of hypertonic aCSF (330 mOsm/l; 170 mM NaCl) depolarized vMnPO neurons. Surprisingly, vMnPO cells did not respond to extracellular hyperosmolality (330 mOsm/l) when \([\text{Na}^+]_{\text{out}}\) was kept constant (150 mM NaCl). Because it is generally accepted that deviation of 20 mOsm/l from the set point corresponds to an approximate change of \([\text{Na}^+]_{\text{out}}\) by 10 mM, we did expect that setting the extracellular osmolality to 330 mOsm/l with mannitol would mimic a response obtained with a \([\text{Na}^+]_{\text{out}}\) amounting to 165–170 mM. This was not the case. Furthermore, we showed that vMnPO neurons did respond to hyponatremia without a change in extracellular osmolality.

Our results clearly indicate that a subpopulation of MnPO neurons located in both parts of the nucleus are specifically endowed with the detection of \([\text{Na}^+]_{\text{out}}\) and, thus, favor the role of MnPO neurons as cerebral \( \text{Na}^- \) sensors, whose existence have long been hypothesized in periventricular tissue (Cox et al., 1987; Park et al., 1989; Denton et al., 1996). We now provide evidence for functional \( \text{Na}^- \) sensors in the MnPO that likely belong to a complex neuronal network of interconnected structures within the lamina terminalis (Honda et al., 1990; Oldfield et al., 1991, 1992; McKinley et al., 1999; McKinley, 2003). This network might function as a central osmoregulatory unit that might control vasopressin and may be oxytocin secretion, together with the innate \( \text{Na}^- \) and osmolality sensitivity of magnocellular neurons (MCNs) (Voisin et al., 1999; Voisin and Bourque, 2002).

**The cellular mechanism underlying Na\(^+\) detection in the MnPO neurons**

The association of the background \( \text{Na}^+ \) permeability with the high input resistance of the MnPO neurons (input resistance, >1 G\(\Omega\)) might represent the cellular mechanism of sodium sensing in this region of the lamina terminalis. Indeed, a small increase in the leak \( \text{Na}^+ \) current (\([\text{Na}^-]_{\text{out}}\) above 150 mM) appears sufficient to depolarize the MnPO neurons, leading to spiking activity. Conversely, a slight reduction of the leak \( \text{Na}^+ \) current (\([\text{Na}^-]_{\text{out}}\) below 150 mM) hyperpolarizes the neurons, thus reducing their excitability.

One possibility was that the sustained \( \text{Na}^+ \) permeability, and thus the \( \text{Na}^- \) sensing ability, might be attributable to the expression of SICs, highly permeable to \( \text{Na}^- \), as shown in the hypothalamic MCNs (Voisin et al., 1999). However, the involvement of the SICs in the vMnPO neurons was unlikely, as demonstrated by the inability of Gd\(^{3+}\) to abolish the responses to hypotonic and hypertonic aCSF. Although sensitivity to Gd\(^{3+}\) was often considered to be an intrinsic property of mechanoreceptors, a different class of mechanosensitive channels expressed in corneal epithelial cells, or in colon DRG cells, were insensitive to Gd\(^{3+}\) (Watanabe et al., 1997; Su et al., 2000). In the study by Su et al. (2000), the open probability of the mechanosensitive channels was reduced by amiloride, a pharmacological agent that was also shown to block constitutively active specific \( \text{Na}^- \) channels (Horisberger, 1998), such as the epithelial \( \text{Na}^- \) channels involved in \( \text{Na}^- \) homeostasis (Garty and Palmer, 1997). However, hyperpolarization induced by the hypotonic stimulus was not antagonized by either amiloride or benzamil, a potent amiloride analog. This ruled out the involvement of such amiloride-sensitive \( \text{Na}^- \) channels in the cellular mechanism of \( \text{Na}^- \) detection used by the MnPO neurons.

The unsuccessful pharmacological characterization of the na-
triuretic responses suggests that detection of CSF [Na\(^+\)] in MnPO neurons was mediated by specific channels highly permeable to Na\(^+\) but with unknown pharmacology. An attractive possibility might be the presence of the Na\(_{\text{ax}}\) channel on the MnPO neurons, a specific Na\(^+\) channel that has been shown to mediate sensing of Na\(^+\) levels in the CNS (Hiyama et al., 2002). Several electrophysiological properties, such as TTX resistance, voltage independence, and insensitivity to [Cl\(^-\)]\(_{\text{out}}\) and to a change in extracellular osmolarity, are shared with the Na\(^+\) current that we report here. These similarities might favor the existence of such a channel in the MnPO neurons.

**MnPO neurons encode the atypical sodium channel Na\(_{\text{ax}}\) gene**

To verify the latter possibility, we used a riboprobe directed against the Na\(_{\text{ax}}\) mRNA in combination with the immunohistochemical detection of NeuN. This anatomical study revealed that a substantial neuronal population of the MnPO expressed Na\(_{\text{ax}}\) mRNA, supporting the possibility that the background Na\(^+\) current flows through the atypical Na\(^+\) channel Na\(_{\text{ax}}\). In addition, the anatomical data shed new light on the cellular mechanisms that are likely involved in Na\(^+\) detection by specialized brain areas. Previous studies have shown that the Na\(_{\text{ax}}\) gene, recently renamed Na\(_{\text{ax}}\) gene (Goldin et al., 2000), was highly expressed in the circumventricular organs (CVOs) that lack a regular blood–brain barrier (Watanabe et al., 2000; Hiyama et al., 2002). The use of high cellular resolution techniques such as in situ hybridization and immunohistochemistry showed that additional structures located along the third ventricle, such as the MnPO and the periventricular nucleus (data not shown), also express Na\(_{\text{ax}}\) channel. These results emphasize the role of these structures in the detection of Na\(^+\) concentration fluctuation within the CSF. In addition, the presence of such channels in the CVOs and periventricular regions, but not in the SON, indicate the diversity of molecular mechanisms dedicated to the detection of osmotic and mineral changes in the structures that constitute the interface between the brain and the fluid compartments (CSF and blood).

**Physiological significance**

A majority of results reported in our study were obtained with a drastic change in Na\(^+\) concentration, and one might reasonably question the role of such Na\(^+\) sensors in the context of a highly regulated body Na\(^+\) concentration \textit{in vivo}. However, we showed that a physiological reduction of \~5% in aCSF [Na\(^+\)] (150–142 mM NaCl) was able to initiate a neuronal response, i.e., a membrane hyperpolarization. This important result is in the range of CSF [Na\(^+\)] that is known to induce salt appetite. Indeed, \textit{in vivo} manipulation of CSF osmotic pressure that reduced CSF by 10–15 mM NaCl triggered a robust sodium intake (Weisinger et al., 1979). Under normal conditions, the human body can maintain [Na\(^+\)]\(_{\text{out}}\) between 145 and 135 mM as a physiological range. More drastic alteration of Na\(^+\) levels as those used in this study have been, however, reported (Moritz and Ayus, 2002). Hyponatremia is defined as a serum Na\(^+\) level inferior to 135 mEq, mainly attributable to electrolyte disorders, and might lead to encephalopathy. Severe hyponatremia (107 mM NaCl) has been reported after pituitary tumor (Lin et al., 2002), and cerebral salt wasting syndrome induces acute hyponatremia (120 mM NaCl), usually consecutive to viral infection (Ti et al., 1998; Singh et al., 2002).

**References**


Aradachi H, Honda K, Negoro H, Kubota T (1996) Median preoptic neurons projecting to the supraoptic nucleus are sensitive to haemodynamic changes as well as to rise in plasma osmolality in rats. J Neuroendocrinol 8:35–43.


