Epithelial Na$^+$ sodium channels in magnocellular cells of the rat supraoptic and paraventricular nuclei

Ryoichi Teruyama,1 Mayumi Sakuraba,2 Lori L. Wilson,1 Narine E. J. Wandrey,1 and William E. Armstrong2

1Department of Biological Sciences, Louisiana State University, Baton Rouge, Louisiana; and 2Department of Anatomy and Neurobiology, University of Tennessee Health Science Center, Memphis, Tennessee

Submitted 5 August 2011; accepted in final form 31 October 2011

Teruyama R, Sakuraba M, Wilson LL, Wandrey NE, Armstrong WE. Epithelial Na$^+$ sodium channels in magnocellular cells of the rat supraoptic and paraventricular nuclei. Am J Physiol Endocrinol Metab 302: E273–E285, 2012. First published November 1, 2011; doi:10.1152/ajpendo.00407.2011.—The epithelial Na$^+$ channels (ENaCs) are present in kidney and contribute to Na$^+$ and water homeostasis. All three ENaC subunits (α, β, and γ) were demonstrated in the cardiovascular regulatory centers of the rat brain, including the magnocellular neurons (MNCs) in the supraoptic nucleus (SON) and the paraventricular nucleus (PVN). However, the functional significance of ENaCs in vasopressin (VP) and oxytocin (OT) synthesizing MNCs is completely unknown. In this study, we show with immunocytochemical double-labeling that the α-ENaC is colocalized with either VP or OT in MNCs in the SON and PVN. In addition, parvocellular neurons in the dorsal, ventrolateral, and posterior subregions of the PVN (not immunoreactive to VP or OT) are also immunoreactive for α-ENaC. In contrast, immunoreactivity to β- and γ-ENaC is colocalized with VP alone within the MNCs. Furthermore, immunoreactivity for a known target for ENaC expression, the mineralocorticoid receptor (MR), is colocalized with both VP and OT in MNCs. Using single-cell RT-PCR, we detected mRNA for all three ENaC subunits and MR in cDNA libraries derived from single MNCs. In whole cell voltage clamp recordings, application of the ENaC blocker benzamil reversibly reduced a steady-state inward current and decreased cell membrane conductance approximately twofold. Finally, benzamil caused membrane hyperpolarization in a majority of VP and about one-half of OT neurons in both spontaneously firing and quiet cells. These results strongly suggest the presence of functional ENaCs that may affect the firing patterns of MNCs, which ultimately control the secretion of VP and OT.

aldosterone; vasopressin; oxytocin

THE NEUROHYPOPHYSIAL HORMONES vasopressin (VP) and oxytocin (OT) are synthesized in the magnocellular neurons (MNCs) located within the paraventricular nucleus (PVN) and the supraoptic nucleus (SON) of the hypothalamus and released from the neurohypophysis into the general circulation in response to physiological demands. The secretion of VP increases in response to hyperosmolality, hypovolemia, and hypertension and produces antidiuretic and pressor effects (59). In addition to the well-known effects of OT during parturition and lactation, plasma OT also increases with hyperosmolality and hypernatremia (32) and induces natriuresis (13, 31).

The non-voltage-dependent, amiloride-sensitive epithelial Na$^+$ channels (ENaCs) are present in the apical membrane of epithelial cells in a variety of tissues, such as urinary bladder, renal collecting duct, distal colon, sweat and salivary glands, lung, and taste buds, and are known to mediate the transport of Na$^+$ across epithelia (7, 21). Thus, together with the Na$^+$/K$^+$ - ATPase present in the basal membrane of epithelial cells, ENaCs regulate transepithelial Na$^+$ transport; however, entry across the apical membrane through ENaCs is the rate-limiting step under most circumstances (21). The ENaCs located in the distal nephron in kidney are known to finely regulate blood pressure and extracellular fluid volume by modulating Na$^+$ excretion and reabsorption (7, 21). Activity of ENaCs is regulated largely by the adrenal mineralocorticoid aldosterone through its mineralocorticoid receptor (MR) (42).

Interestingly, both mRNA and protein for all three ENaC subunits (α, β, and γ) and MR were demonstrated in regions implicated in cardiovascular control, such as the MNCs in the SON and PVN, and also in the hippocampus, choroid plexus, ependyma, and brain blood vessels in rats (4). The physiological function of the ENaC in neurons is not well understood; however, the locations of ENaC in the brain suggest a role in cardiovascular regulation. Intracerebroventricular (icv) infusion of aldosterone increases blood pressure, presumably via upregulation of ENaC in the brain in Wistar rats (68, 69) and in an animal model of salt-sensitive hypertension, Dahl salt-sensitive (Dahl-S) rats (25). Importantly, icv injections of the ENaC blocker, the amiloride analog benzamil, significantly attenuated hypertension in these animals (27, 47). These findings suggest that aldosterone-mediated activation of ENaCs in brain could contribute to the development of hypertension and that central ENaC inhibition may be a potential new target in the treatment of cardiovascular disease (63). However, the significance of ENaCs in regulating MNC electrical activity is unknown. In the present study, we confirmed the presence of ENaCs and MR in MNCs and determined the extent of their selective expression in VP and OT neurons, and for the first time we show that ENaCs contribute to the membrane potential in these neurons.

MATERIALS AND METHODS

Animals

Male Sprague-Dawley rats were used (180–210 g body wt; Harlan Laboratories, Indianapolis, IN). The rats were housed in a room on a 12:12-h light-dark cycle, with access to food and water available ad libitum. All protocols were approved by the Institutional Animal Care and Use Committees at the University of Tennessee and Louisiana State University.

Electrophysiology

Slice preparation. The rats were deeply anesthetized with pentobarbital sodium (50 mg/kg ip) and perfused through the heart with
cold artificial cerebrospinal fluid (ACSF; see below), in which NaCl was replaced by an equiosmolar amount of sucrose. Brains were removed, and coronal slices (250 μm) containing SON were obtained by a vibrating blade microtome (Leica VT1000S; Leica, Mannheim, Germany).

**Recording.** Whole cell patch clamp recordings were acquired digitally at 20 kHz and filtered at 5 kHz with a Digidata 1440A and an Axopatch 700A or 700B (Molecular Devices, Foster City, CA) amplifier in conjunction with PClamp 10 software (Molecular Devices) on a Windows platform PC. The MNCs in the SON were identified visually using an Olympus BX51WI microscope and a ×40 water immersion lens (0.8 na) under IR illumination (780 ± 30 nm) using a charge-coupled device camera. Recordings were taken using borosilicate electrodes (4–8 MΩ resistance) produced with a horizontal electrode puller (Model P-87 Flaming/Brown Micropipette puller; Sutter Instruments, Novato, CA). The patch solution for voltage clamp experiment contained (in mM) 100 n-gluconic acid, 100 CsOH, 20 CaCl2, 10 HEPES, 1 CaCl2, 2 MgCl2, 10 NaCl, 2 adenosine 5′-triphosphate (ATP), and 0.4 guanosine 5′-triphosphate (GTP) and for current clamp experiment contained (in mM) 140 K-gluconate, 1 MgCl2, 2 ATP (Mg2+), 0.4 GTP (Na+), and 1 EGTA. The patch solutions also contained 0.2% biocytin (Sigma) to fill the patched cell (64, 65). The ACSF consisted of (in mM) 125 NaCl, 2.5 KCl, 1 MgSO4, 1.25 Na2HPO4, 26 NaHCO3, 20 d-glucose, 2 CaCl2, and 0.4 ascorbic acid. The medium was saturated with 95% O2-5% CO2, with a pH of 7.3–7.4 and an osmolality of 290–300 mOsm/kg H2O. Picrotoxin and DNQX (100 and 10 μM, respectively) were also added to ACSF to suppress the synaptic activity. Solutions were warmed to 33–34°C during the recordings.

**Post Hoc Immunocytochemical Identification of Cell Types**

The slices were fixed (4% paraformaldehyde-0.1% picric acid in 0.15 M sodium phosphate buffer, pH 7.2–7.4) and processed for double-immunofluorescence labeling to identify the cell type (64, 65). To identify VP neurons, a specific VP-neurophysin (NP) polyclonal antibody (53) raised in rabbits was used at a 1:20,000 dilution (provided by Dr. Mark A. Knepper (National Institutes of Health, Bethesda, MD). The production and characterization of these ENaC subunits antibodies were described previously in great detail (42). The anti-VP-NP (PS41) and the anti-OT-NP (PS38) were raised in mouse against VP-NP or OT-NP, respectively, and used at a 1:500 dilution (see above). The slices were first incubated with one of the anti-ENaC subunits for 48–72 h at 4°C, followed by the incubation with either anti-VP-NP or anti-OT-NP for 48–72 h at 4°C.

The monoclonal anti-mineralocorticoid receptor antibody (MRN3 3F10) developed by C. E. Gomez-Sanchez was obtained from the Developmental Studies Hybridoma Bank developed under the auspices of the National Institute of Child Health and Human Development and maintained by The University of Iowa, Department of Biology, Iowa City, IA. The production and characterization of the MR antibody were described previously in great detail (24). The anti-VP-NP (Rob-VP) and the anti-OT-NP (Rob-OT) antiserum used for double labeling with MR antibody were provided by Alan Robinson (UCLA). Rob-VP and -OT antiserum (53) were raised in rabbit against VP-NP or OT-NP, respectively, and used at 1:20,000 and 1:10,000 dilutions, respectively.

After incubations with primary antibodies, the slices were incubated in a cocktail of appropriate secondary antibodies for 2 h at room temperature. The secondary antibodies used were DyLight 488-conjugated goat anti-rabbit and DyLight 594-conjugated goat anti-mouse IgG (Jackson ImmunoResearch, West Grove, PA). The brain slices were examined, and confocal images (1,024 × 1,024) were acquired with a confocal microscope (Leica TCS SP2 spectral confocal microscope). The optical section thickness was 1 μm. These were viewed in stacks of three to five sections using ImageJ software (NIH).

**Table 1. Primer sequences used to detect gene expressions of interest**

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer Sequence</th>
<th>Amplicon Size, bp</th>
</tr>
</thead>
<tbody>
<tr>
<td>α-ENaC</td>
<td>Forward 5'-GTTCCTGTCAGCTACGAAAAAGGAGAG-3'</td>
<td>429</td>
</tr>
<tr>
<td></td>
<td>Reverse 5'-GTTAGGAGCAGGATGAG-3'</td>
<td>429</td>
</tr>
<tr>
<td>β-ENaC</td>
<td>Forward 5'-ACCCCTGACGAAGGAGGAT-3'</td>
<td>260</td>
</tr>
<tr>
<td></td>
<td>Reverse 5'-AAAGGAGCAGGATGAG-3'</td>
<td>260</td>
</tr>
<tr>
<td>γ-ENaC</td>
<td>Forward 5'-CTCTAAGTGCTCAGTACA-3'</td>
<td>301</td>
</tr>
<tr>
<td></td>
<td>Reverse 5'-GAGAGGCTCAGGAAAACCTGAT-3'</td>
<td>301</td>
</tr>
<tr>
<td>MR</td>
<td>Forward 5'-GCTCAACATTGGCTCACAGTACA-3'</td>
<td>463</td>
</tr>
<tr>
<td></td>
<td>Reverse 5'-GCCAGAGGCTCAGGAAAACCTGAT-3'</td>
<td>463</td>
</tr>
<tr>
<td>OT</td>
<td>Forward 5'-AGGGGGGAGAGCAGGATGAG-3'</td>
<td>440</td>
</tr>
<tr>
<td>VP</td>
<td>Reverse 5'-ACCCCTAAGGAGGAGGAT-3'</td>
<td>440</td>
</tr>
</tbody>
</table>

ENaC: epithelial Na+ channel; MR, mineralocorticoid receptor; OT, oxytocin; VP, vasopressin.
Single-Cell RT-PCR

Single-cell harvest for single-cell RT-PCR. The brains were sliced as described in Slice preparation. Small pieces of brain (~2 × 2 mm) containing the SON were dissected from the slices under a stereomicroscope. These pieces were incubated in oxygenated ACSF (35°C) containing protease type XIV (1.2 mg/ml; Sigma Chemical, St. Louis, MO) for 20–30 min and then washed with sodium isethionate solution (in mM: 140 sodium isethionate, 2 KCl, 4 MgCl₂, 23 glucose, 15 HEPES, pH 7.3). The enzyme-treated tissues were triturated in sodium isethionate solution using three successively smaller fire-polished pipettes to release individual MNC cell bodies. The supernatant containing dissociated neurons from each trituration step was transferred to a plastic Petri dish (Nunc, Rochester, NY) on an inverted microscope stage, and cells were allowed to settle for ~5 min. A background flow of ~1 ml/min of HEPES-buffered saline solution (HBSS) was then established. HBSS consisted of (in mM) 138 NaCl, 3 KCl, 1 MgCl₂, 2 CaCl₂, 10 HEPES, and 20 dextrose, pH 7.3 (adjusted with 1 N NaOH), and osmolarity = 300–305 mOsmol/l.

Electrode glass (Corning 7052 capillary glass; Garner Glass, Claremont, CA) was autoclaved to prevent RNAse contamination. Electrodes were pulled on a Sutter Instrument (Novato, CA) Model P-87 Flaming/Brown Micropipette puller, fire-polished, and filled with HBSS made with RNAse-free water. Positive pressure was applied to the pipette while navigating to the cell to minimize contamination. The electrode with an attached cell was lifted into a stream of ACSF and washed for 5 min before the cell was sucked into the pipette.

Following aspiration, the contents of the electrode were ejected into a chilled, 0.5-µl presiliconized RT tube containing a cellular mixture [1.9 µl of diethylpyrocarbonate (DEPC)-treated water, 1.0 µl of dNTP (10 mM), 0.7 µl of BSA (143 µg/µl), 0.7 µl of oligo(dT) (0.5 µg/µl), and 0.7 µl of SUPERase-in (40 U/µl)]. The mixture was stored at −80°C or used immediately for RT.

RT. The mixture was heated to 65°C and then placed on ice for ≥1 min. Single-stranded cDNA was synthesized from the cellular mRNA after 16 µl of RT Master Mix (6.0 µl of DEPC-treated water, 2.0 µl of 10× RT buffer, 4.0 µl of MgCl₂ (25 mM), 2.0 µl of DTT (0.1 M), 1.0 µl of RNAse Out, and 1.0 µl of Superscript III) was added. This mixture was incubated at 42°C for 50 min and then terminated by heating to 70°C for 15 min. The reactions were collected by a brief centrifugation and then incubated in 0.5 µl of RNAse H (2 U/µl) for 20 min to remove any remaining RNA from the reaction. The cDNA can be stored at −80°C or used immediately for PCR.

PCR. The single-cell cDNA generated from the RT step was subjected to conventional PCR using a programmable thermal cycler from MJ Research (Waltham, MA) and primers designed specifically to amplify the cDNA of interest (Table 1). Identification of each cDNA was based on the predicted size of each PCR product. These primers listed in Table 1 have been used successfully (4, 23). Negative controls for contamination from extraneous and genomic DNA from other sources were run for every batch of neurons. To ensure that there was no contamination from genomic DNA, reverse transcriptase was omitted. Replacing the cellular...
template with water controlled for contamination from extraneous sources.

RESULTS

Immunocytochemical Localization of ENaC Subunits in the SON and the PVN

The ENaC subunit antibodies gave consistent results in 22 animals at dilutions of 1:1,000 for α-subunits, 1:250 for β-subunits, and 1:2,000 for γ-subunits, as shown in Figs. 2–7. The most prominent immunoreactivity was observed in the SON and the PVN within coronal sections containing these nuclei. In addition, immunoreactivity was found in cuboidal epithelium of the choroid plexus and in the ventricular ependyma. Overall, the localization of α-, β-, and γ-ENaC subunits within the rat hypothalamus was similar to previous reports (3, 4). Here, we report a more detailed localization of α-, β-, and γ-ENaC within the SON and PVN performed by confocal double immunocytochemistry, using specific antibodies against ENaC subunits and antibodies against VP- or OT-NP.

An intense immunoreactivity to α-ENaC subunit was confined largely to somata of both OT and VP neurons within the SON (Fig. 2). Prominent α-ENaC immunoreactivity was also observed in somata and proximal parts of dendrites of magnocellular cells within the PVN (Fig. 3). Whereas most of these α-ENaC-immunoreactive MNCs were located in a cluster of cells in the posterior magnocellular region, α-ENaC-immunoreactive magnocellular cells were found scattered among parvocellular cells in other regions of the PVN (dorsal, medial, ventrolateral, and posterior parvocellular regions). The lateral cluster of the cells was composed mostly of VP neurons (Fig. 3, A and D); however, some OT neurons scattered around these VP neurons were immunoreactive to α-ENaC. Therefore, the α-ENaC immunoreactivity appears to be colocalized with either VP or OT MNCs. In addition to the MNCs, some parvocellular neurons that were not immunoreactive to VP- or OT-NP within dorsal, ventrolateral, and posterior parvocellular regions were also immunoreactive to α-ENaC subunit. Only sporadic α-ENaC-immunoreactive parvocellular and MNCs were observed within the medial parvocellular region of the PVN.

Fig. 3. Colocalization of α-ENaC subunit and VP-NP immunoreactivity in coronal section of the rat paraventricular nucleus (PVN). A and D: α-ENaC subunit immunoreactivity labeled with DyLight 488-conjugated secondary antibody. Prominent α-ENaC immunoreactivity is observed in somata and proximal parts of dendrites of MNCs within the PVN. Most of these α-ENaC-immunoreactive magnocellular cells are located in a cluster of cells in the posterior magnocellular (pm) region. The parvocellular cells in dorsal (dp), ventrolateral (vlp), and posterior parvocellular regions are also immunoreactive to α-ENaC subunit. B: VP-NP immunoreactivity labeled with DyLight 594-conjugated secondary antibody in same section and image plane as in A. Note that the VP-NP immunoreactive cells form a cluster in the lateral portion of the PVN. E: OT-NP immunoreactivity labeled with DyLight 594-conjugated secondary antibody in same section and image plane as in D, C and F: the merged images revealed that α-ENaC immunoreactivity is colocalized with both VP and OT immunoreactivities within MNCs in the PVN. Note that the lateral cluster of the cells is composed mostly of VP, MNCs are immunoreactive to α-ENaC as well as those sparsely located OT MNCs around these VP neurons. However, the parvocellular cells immunoreactive to α-ENaC within dp, vlp, and posterior parvocellular regions are not immunoreactive to VP or OT.
Immunoreactivity of β-ENaC in the SON was also confined within the somata of MNCs (Fig. 4). Although the entire area of the SON has diffuse staining that stands out from the rest of the brain area, indicating that almost all MNCs have some degree of immunoreactivity, not all of the MNCs exhibited intense somatic immunoreactivity. Double immunocytochemistry either with VP- or OT-NP revealed that these β-ENaC-immunoreactive cells were all VP and not OT neurons (Fig. 4, C and F). Not many MNCs possessed intense β-ENaC immunoreactivity in the PVN (Fig. 5); however, unlike in the SON, the immunoreactivity in these cells was not confined within the soma. Prominent immunoreactivity is seen in the dendritic processes of these neurons, located mostly in the lateral portion of the PVN. Again, double immunocytochemistry exposed that β-ENaC-immunoreactive neurons were immunoreactive to VP-NP and not OT-NP (Fig. 5, arrows). Moreover, there was no apparent β-ENaC immunoreactivity in the parvocellular cells in the PVN.

Immunoreactivity to γ-ENaC was present in many MNCs in the SON, with intense staining observed in a minority of MNCs and their sparsely distributed thick dendritic processes. (Fig. 6, arrows). Intense γ-ENaC immunoreactivity was also found in thick processes (probable dendrites) and somata of a cluster of MNCs within the posterior magnocellular portion of the PVN (Fig. 7). A more striking and consistent pattern of colocalization was present in the somata and dendrites of the MNCs within the SON (Fig. 6, C and F) and PVN (Fig. 7, C and F) after reaction with VP-NP antibody. Some γ-ENaC-immunoreactive MNCs were also immunoreactive to OT in the SON and the PVN, but only barely detectable levels of OT immunoreactivity were observed in all cases. Only weak γ-ENaC immunoreactivity was indicated in the parvocellular cells in the PVN.

An intense immunoreactivity to MR was confined largely to somata of a majority of MNCs in the SON (Fig. 8, A and D). Moreover, the MR immunoreactivity appears to be colocalized with either VP or OT MNCs in the SON (Fig. 8, C and F). In the PVN, most of prominent MR immunoreactivity was located in a cluster of MNCs in the posterior magnocellular region (Fig. 9, A and D). MR immunoreactivity was also colocalized with either VP or OT in MNCs in the PVN. In addition to the MNCs, some parvocellular neurons that were not immunoreactive to VP- or OT-NP within dorsal, ventrolateral, and posterior parvocellular regions were also immunoreactive to MR.

**Single-Cell RT-PCR Detection of ENaC Subunits in the MNCs**

Libraries of cDNA were derived from 12 cells dissociated from SON tissue. *Cells 1–11* had mRNA for OT-NP and/or VP-NP (Fig. 10). Unlike immunocytochemical identification...
that demonstrates that most MNCs are phenotypically distinct, it has been well documented with single-cell RT-PCR that there is a variable amount of OT and VP mRNA coexpression in virtually all of the MNCs in the SON (23, 70, 71). Therefore, without quantitative RT-PCR it is only appropriate to state here that these dissociated cells are confirmed as MNCs producing OT or VP. Of these MNCs, mRNA for \( \alpha \)-ENaC was found in cells 1, 2, 4, and 5, mRNA for \( \beta \)-ENaC was found in cells 1 and 2, mRNA for \( \gamma \)-ENaC was found in cells 5, 9, 10, and 11, and mRNA for MR was found in cells 1, 3, 5, 6, 10, and 11. In addition, cell 12, although it did not contain VP or OT mRNA, contained \( \alpha \)-ENaC, \( \gamma \)-ENaC, and MR. Although single-cell RT-PCR was not strong enough to obtain robust expression of each subunit in every cell, all of these mRNAs were found in a cDNA library derived from punched SON tissues. Taste receptor cells in tongue epithelia are known to express ENaC; therefore, the cDNA acquired from tongue epithelial tissue served as a positive control.

**Benzamil-Sensitive Current in MNCs**

To demonstrate the functional expression of ENaC, we looked for evidence of a benzamil-sensitive current in MNCs using whole cell voltage clamp. However, in the initial part of the study, a clear response to application of either amiloride or benzamil was observed only in a minority of MNCs (~7% of the recorded cell population). The majority of MNCs displayed small changes or no change at all. This percentage was inconsistent with the number of MNCs that were immunoreactive to the antibodies against ENaC subunits. One of the possibilities for the lack of response may be an intracellular Na\(^{+}\)-dependent rundown observed in the cells expressing ENaC (21, 33, 60). A rise in intracellular Na\(^{+}\) concentration results in a slow decrease in the current mediated by ENaC in these cells. We tried to minimize the rise in intracellular Na\(^{+}\) concentration by addition of 1–2 \( \mu \)M amiloride in the perfusion buffer during brain slice preparation and in the incubation medium. Prior to recordings, the brain slices were transported to the chamber of the patch clamp rig that was perfused continuously with ACSF containing 1 \( \mu \)M benzamil. The benzamil was then washed out after establishment of stable patch clamp. In this way, robust responses to benzamil/amiloride were observed in 12 of 19 VP and 5 of 13 OT neurons.
The steady-state current was measured while the cell was held at −70 mV in voltage clamp. Brief hyperpolarizing pulses (15 mV, 200 ms) were injected every 5 s to monitor the input resistance of the cell. Washout of amiloride from the bath resulted in an increase in a resting inward current and a decreased resistance. Subsequent bath application of the ENaC blocker benzamil (1 μM) reduced the resting inward current and decreased conductance ~1.5-fold (Fig. 9). The effect was reversed by wash of benzamil by ACSF. Figure 11 shows an example of the effect of benzamil on a VP neuron, and we observed this in six of nine VP neurons. The response was also observed in 1 of 5 OT neurons tested.

Next, we looked for the effect of benzamil on the membrane potential and the firing pattern using whole cell current clamp recording with no injected current. The removal of amiloride from the bath resulted in an increase in a resting inward current and a decreased resistance. Subsequent bath application of the ENaC blocker benzamil (1 μM) reduced the resting inward current and decreased conductance ~1.5-fold (Fig. 9). The effect was reversed by wash of benzamil by ACSF. Figure 11 shows an example of the effect of benzamil on a VP neuron, and we observed this in six of nine VP neurons. The response was also observed in 1 of 5 OT neurons tested.

The effect was reversed by washout of benzamil from the bath and was repeatable.

**DISCUSSION**

Amiloride and its analogs are known inhibitors of the degenerin/epithelial sodium channel (Deg/ENaC) superfamily of ion channels. The Deg/ENaC superfamily includes the ENaC and the acid-sensitive ion channels (ASIC) as members. ASIC1a and ASIC2a subunits were detected, and an acid- and amiloride-sensitive current was detected in SON MNCs (48). However, the concentration of amiloride required to inactivate the ASICs in the MNCs in that study was high (10−100 μM), and 1 μM amiloride had little effect on the ASIC activity. Amiloride is also a known inhibitor of several other ion transporters, including the Na⁺/H⁺ exchanger (NHE) and the Na⁺/Ca²⁺ exchanger (NCX) (36). However, amiloride at low doses is reasonably specific for ENaCs compared with the NHE and NCX (36). Nevertheless, the amiloride analog benzamil can be used to further increase selectivity for ENaC compared with the NHE or NCX. Benzamil, compared with amiloride, is ninefold more potent toward ENaCs, with markedly lower relative potency (0.08-fold) toward the NHE (14). The concentration of benzamil (1–2 μM) used in our study is
well below the IC50 of benzamil for these ion exchangers. Therefore, when combined with the immunocytochemical and single-cell RT-PCR data, the electrophysiological results from this study strongly suggest the presence of functional ENaCs in the MNCs.

The release of OT and VP from the neurohypophysis depends largely on the pattern of electrical activity of their synthesizing neurons (12, 49). During the release of VP in response to hypovolemia (29), hypotension (34), and hyperosmolality (8), VP neurons increase their firing rate and adapt a phasic firing pattern comprising alternating periods of activity (7–15 Hz) and silence, each lasting tens of seconds. Plasma OT also increases in response to hypernatremia (32), and OT neurons respond to hyperosmolarity with increases in firing rate (52, 67). In the present study, the benzamil-sensitive current was identified as an inward leak current, suggesting its potential for modulating membrane potential. The activation of the benzamil-sensitive current, presumably mediated by ENaCs, would depolarize the membrane potential and allow MNCs to initiate bursting activity. Computational studies in the supraoptic MNCs showed that a Na+ leak current is critical to the depolarizations underlying phasic firing (54). Therefore, by changes in membrane potential, the modulation of ENaCs may significantly contribute to the regulation of firing activities of these neurons and ultimately affect the release of VP and OT.

An intriguing finding in this study is that β- and γ-ENaC subunits were preferentially located in VP neurons, whereas the α-ENaC subunit was located in both VP and OT MNCs. It is generally agreed that each of the three ENaC subunits contributes to the formation of the functional channel complex (11, 19), although the actual subunit composition of the channel remains uncertain and controversial. Interestingly, α-subunits alone or with either the β- or γ-subunit can induce low but measurable amiloride-sensitive currents (11, 20). Moreover, alteration of subunit composition is suggested as a cause of the variability in single-channel properties of amiloride-sensitive ENaCs in native tissues (20). Thus, this differential subunit expression suggests that subunit composition differs between VP and OT neurons and may be physiologically important. This may be the molecular mechanism accounting for our finding that more VP (≈70%) than OT neurons (≈40%) were responsive to benzamil.

It is well documented that the α-subunit is regulated independently of β- and γ-subunits. Binding of aldosterone to the

---

**Fig. 7. Colocalization of γ-ENaC subunit with VP-NP or OT-NP immunoreactivity in coronal section of the rat PVN.** A and D: γ-ENaC subunit immunoreactivity labeled with DyLight 488-conjugated secondary antibody. Note that intense immunoreactivity to γ-ENaC is observed in the MNCs and their thick processes (probable dendrites). These immunoreactive cells form a cluster in the lateral portion of the PVN. B: VP-NP immunoreactivity labeled with DyLight 594-conjugated secondary antibody in same section and image plane as in A. Note that the VP-NP immunoreactive cells form a cluster in the lateral portion of the PVN. E: OT-NP immunoreactivity labeled with DyLight 594-conjugated secondary antibody in same section and image plane as in D. C: the merged images revealed that γ-ENaC immunoreactivities appear to be colocalized only with VP within the MNCs located mostly in the lateral portion of the PVN. F: the merged image of D and E. Note that γ-ENaC immunoreactivities do not appear to be colocalized with OT.
MR was shown to predominantly increase the expression of α-subunits in rat kidney (18, 41, 42, 51, 62). Interestingly, the aldosterone regulation of ENaC subunits is somewhat tissue specific. For example, β- and γ-ENaC mRNA are selectively induced by mineralocorticoids in colon (5, 18, 51), whereas mRNA for all three subunits is primarily regulated by glucocorticoids in lung tissue (51). The presence of MR in the brain, including the SON and the PVN, was reported previously (2, 4, 28, 55). Nevertheless, the finding of MR in VP and OT MNCs in present study suggests that the expression of ENaC in the MNC is also promoted by aldosterone. Intracerebroventricular infusion of aldosterone causes hypertension Dahl-S rats, but icv infusion of a mineralocorticoid receptor antagonist prevents the salt-induced hypertension (25). Importantly, icv infusion of amiloride or benzamil prevents salt-induced hypertension in Dahl-S rats (26, 69). Obviously, these findings do not provide information about the upregulation of specific ENaC subunits by aldosterone in the MNC but suggest an aldosterone-mediated activation of ENaC in these neurons.

In addition to aldosterone, VP is also known mainly to induce the expression of β- and γ-ENaC subunits in the kidney (16, 17, 46, 57). Of the three VP receptor subtypes (V1a, V1b, and V2), characterized, VP appears to act through the V2 vasopressin receptor to induce β- and γ-subunit gene expression (46). VP also stimulates translocation of preexisting intracellular pools of ENaC subunits to the apical membrane of the principal cells in the collecting duct, promoting Na+ reabsorption in kidney (58). VP neurons not only secrete VP at the nerve terminals in the neurohypophysis but also release VP in the extracellular space of the SON and PVN from their soma and dendrites (37, 39, 45). A recent study demonstrated that the V2-like receptor mediates the ability of somatodendritically released VP to facilitate cell volume regulation in VP neurons (56). In addition, somato-dendritic release of VP is known to modulate electrical activity of VP neurons primarily via the V1a receptor (15, 40, 43, 44). Nevertheless, the presence of V2-like receptors on VP neurons implies the possibility that the β- and γ-subunits in VP neurons could be regulated by somato-dendritic release of VP by mechanisms similar to their regulation in the nephron.

Another finding of interest in the current study is the presence of α-ENaC immunoreactivity in a subpopulation of the parvocellular neurons in the dorsal and ventrolateral parvocellular regions of the PVN. These PVN parvocellular neurons were not immunoreactive for OT-NP or VP-NP and were found in regions that send projections to either the rostral ventrolateral medulla that contain sympathetelic premotor neurons or the spinal sympathetic preganglionic neurons to affect the autonomic nervous system (30, 50, 61). In addition, there was no clear immunoreactivity for β- and γ-subunits in these parvocellular populations. This solitary expression of α-ENaC in the parvocellular cell population suggests a different role for ENaCs in these neurons.

Fig. 8. Colocalization of MR with both VP-NP and OT-NP immunoreactivity in coronal section of the rat SON. A and D: MR immunoreactivity labeled with DyLight 488-conjugated secondary antibody. Note that the intense immunoreactivity to MR appeared to be confined in somata of the MNCs within the SON. B: VP-NP immunoreactivity labeled with DyLight 594-conjugated secondary antibody in same section and image plane as in A. E: OT-NP immunoreactivity labeled with DyLight 594-conjugated secondary antibody in same section and image plane as in D. C and F: the merged images revealed that MR is colocalized with both VP-NP and OT-NP immunoreactivities within MNCs in the SON (yellow).
These findings, along with the report that α-subunits alone can carry low but measurable amiloride-sensitive currents (11, 20), suggest the ENaCs in these nonneuroendocrine parvocellular cells may have different biophysical characteristics from those in the MNCs.

In taste receptor cells, ENaCs act as Na$^+$ sensors and play an important role in salt taste transduction (22, 35, 38). The existence of neuronal elements that are sensitive to Na$^+$ was suggested, since the effects induced by icv administration of hypertonic NaCl were shown to originate from the changes in CSF Na$^+$ concentration but not in CSF osmolarity (9, 10). Therefore, it is possible that ENaCs in MNCs also act as Na$^+$ sensors, where they could underlie the specific Na$^+$ sensitivity attributed previously to MNCs in the SON (66).

ACKNOWLEDGMENTS

We thank Drs. J. T. Caprio and E. L. Gleason for reading earlier versions of this manuscript, Dr. M. A. Knepper for providing ENaC antibodies, and Grant Bertolet for technical assistance.

GRANTS

This work was supported by National Institutes of Health Grants R21-HL-0932728 (R. Teruyama), R21-HL-09327285 (R. Teruyama), R01-NS-2394-19 (W. E. Armstrong), R01-NS23941-19S (W. E. Armstrong), and R56-NS-23941-20 (W. E. Armstrong) and in part from a grant to the Louisiana State University College of Science from the Howard Hughes Medical Institute Biomedical Education Program.

DISCLOSURES

The authors have nothing to disclose.

AUTHOR CONTRIBUTIONS

R.T. did the conception and design of the research; R.T., M.S., L.L.W., and N.E.W. performed the experiments; R.T. and M.S. analyzed the data; R.T. interpreted the results of the experiments; R.T., L.L.W., and N.E.W. prepared the figures; R.T. drafted the manuscript; R.T. and W.E.A. edited and revised the manuscript; R.T. approved the final version of the manuscript.

REFERENCES

libraries of cDNA were derived from 12 cells dissociated from SON tissue. Libraries of cDNA were derived from 12 cells dissociated from SON tissue. Libraries of cDNA were derived from 12 cells dissociated from SON tissue.

Fig. 10. Single-cell RT-PCR to detect transcripts in individual dissociated cells. OT and VP mRNA were found in cells 1, 2, 3, and 5, α-ENaC mRNA was found in cells 2, 4, and 5, β-ENaC mRNA was found in cells 1, 3, and 4, and γ-ENaC mRNA was found in cells 2, 3, 4, and 5. All of these mRNAs were found in cDNA library derived from punched SON tissues. Taste receptor cells in tongue epithelia are known to express ENaC, and therefore, the cDNA acquired from tongue epithelial tissue (TET) served as a positive control. Negative controls (NC) for contamination from extraneous and genomic DNA from other sources were run for every batch of neurons. Amplifications with no cDNA yielded negative results.


Fig. 11. A representative recording from a VP neuron showing the effect of benzamil on the current measured at −70 mV. The steady-state current was measured when the cell was held at −70 mV in voltage clamp. Brief hyperpolarizing pulses (200 ms, −15 mV) were injected every 5 s to monitor the input resistance of the cell. Washing amiloride from the bath resulted in an increase in a resting inward current and a decreased resistance. Subsequent bath application of ENaC blocker benzamil (1 μM) reduced the resting inward current and decreased conductance approximately 1.5-fold. The effect was reversed by wash of benzamil by artificial cerebrospinal fluid (ACSF).
Fig. 12. An example of the effect of benzamil on the membrane potential and the firing pattern of VP neuron. The removal of benzamil from the bath caused depolarization and firing of the VP cell. Subsequent bath application of 1 μM benzamil caused membrane hyperpolarization and cessation of the firing. No current was injected during the recording.


