Excess Dietary Salt Alters Angiotensinergic Regulation of Neurons in the Rostral Ventrolateral Medulla

Julye M. Adams, John J. McCarthy, Sean D. Stocker

Abstract—Excess dietary salt intake contributes to or exacerbates some forms of hypertension by increasing sympathetic nerve activity (SNA) and arterial blood pressure (ABP) through angiotensin II (Ang II) type 1 receptor activation in the rostral ventrolateral medulla (RVLM). Despite this interaction among dietary salt, Ang II, and the RVLM, no studies have directly examined whether dietary salt by itself alters Ang II–dependent responses and regulation of RVLM neurons, SNA, and ABP. Therefore, the present study directly tested this hypothesis. Male Sprague-Dawley rats were fed normal chow and given access to water or 0.9% NaCl solution for 14 days. Unilateral injection of Ang II (0.6, 6, and 60 pmol) into the RVLM produced a significantly greater increase in renal SNA and mean ABP of rats drinking 0.9% NaCl versus water. However, dietary salt did not alter mRNA levels of RVLM Ang II type 1a receptors or the SNA and ABP responses to stimulation of the dorsolateral funiculus. Additional experiments demonstrate that blockade of RVLM Ang II type 1 receptors significantly reduced renal SNA, splanchnic SNA, and mean ABP of rats drinking 0.9% NaCl but not water. Blockade of ionotropic glutamate receptors had no effect. Altogether, these findings suggest that elevated dietary salt enhances the sympathoexcitatory actions of Ang II in the RVLM via changes in the intrinsic properties of RVLM neurons. Moreover, elevated dietary salt intake differentially affects the tonic activity of the peripheral versus brain RVLM Ang II type 1 receptors to regulate baseline SNA and ABP. (Hypertension. 2008; 52:932-937.)

Key Words: sympathetic nervous system ■ blood pressure ■ hypertension ■ brain ■ sodium

The brain renin-angiotensin system plays an important role in cardiovascular regulation through its ability to modulate sympathetic nerve activity (SNA) and arterial blood pressure (ABP). One of the major centers postulated to mediate the sympathoexcitatory actions of brain angiotensin II (Ang II) is the rostral ventrolateral medulla (RVLM).1,2 The RVLM contains tonically active, bulbospinal neurons that mediate a number of sympathetically mediated reflexes and contribute to sympathoexcitatory disease states.3 The RVLM contains a high density of Ang II type 1 (AT1) receptors,1 and microinjection of Ang II into the RVLM produces an AT1 receptor–mediated increase in SNA and ABP.2 Overexpression of constitutively active AT1 receptors in the RVLM increases ABP,4 and blockade of RVLM AT1 receptors has been reported to reduce ABP in several forms of experimental hypertension5,6 and other sympathoexcitatory conditions.7,8

A major factor that contributes to the pathogenesis of hypertension and modulates the excitability of RVLM sympathetic-regulatory neurons is dietary salt intake. Evidence from both clinical and experimental models indicate that excess dietary salt intake exacerbates the level of hypertension, including Ang II–dependent models, via increases in sympathetic vasomotor tone.9,10 In several instances, findings from experimental models suggest that the elevated sympathetic tone depends on altered neurotransmission and AT1 receptor activation in the RVLM.5,6 However, few studies have directly examined whether excess dietary salt intake by itself affects the regulation of RVLM sympathetic-regulatory neurons. To date, the limited data indicate that elevated dietary salt intake enhances both sympathoexcitatory and sympathoinhibitory responses evoked by a number of neurotransmitters exogenously applied to the RVLM.11–13 However, these enhanced responses could not be attributed to changes in downstream sympathetic pathways13,14 or vascular reactivity.12,13,15

Despite the ability of dietary salt intake to modulate the excitability of RVLM neurons and the role of RVLM AT1 receptors in salt-dependent hypertension, no studies have determined whether dietary salt intake by itself alters Ang II–dependent responses or regulation of RVLM neurons, SNA, and ABP. The present findings indicate that excess dietary salt intake enhances the sympathoexcitatory responses evoked by exogenously applied Ang II in the RVLM without changes in mRNA levels for RVLM AT1 receptors or downstream sympathoexcitatory pathways. Additional experiments indicate that elevated dietary salt intake tonically activates RVLM AT1 receptors to maintain baseline SNA and ABP.
Materials and Methods

Animals
All of the experimental procedures conform to the National Institutes of Health Guide for the Care and Use of Laboratory Animals and were approved by the University of Kentucky Institutional Animal Care and Use Committee. Male Sprague-Dawley rats (200 to 250 g; Charles River Laboratories) were housed in a temperature-controlled room (22 ± 1°C) with a 14:10-hour light/dark cycle. Rats were fed standard rat chow containing 0.23% NaCl (Harlan Teklad Global Diet #2018) and given access to deionized water for ≥ 7 days before experiments began. Then rats were fed standard chow and given access to deionized water or 0.9% NaCl for 14 days. This protocol produces reliable and physiological increases in daily salt intake of rodents.11

General Procedures and RVLM Microinjections
Rats were anesthetized with a mixture of urethane (750 mg/kg IV) and α-chloralose (75 mg/kg IV) and prepared for recordings of renal or splanchnic SNA and ABP as described previously.11 Animals were artificially ventilated with oxygen-enriched room air and paralyzed with gallamine triethiodide (25 mg/kg IV). End-tidal CO2 and body temperature were maintained at 4% to 4.5% and 37 ± 1°C, respectively. An adequate depth of anesthesia was assessed by either the absence of a withdrawal reflex (before neuromuscular blockade) or a pressor response to foot pinch. Supplemental doses of anesthetic (10% initial dose) were given as necessary. RVLM microinjections were performed as described previously in our laboratory.11 Initially, L-glutamate (1 nmol) was injected into the RVLM at 3 different sites separated by 300 μm in the rostral–caudal plane to identify the site that produced the largest increase in ABP; subsequent injections were performed at those coordinates. For all experiments, injections (60 nl) were performed over 5 seconds. Injection sites were marked at the end of experiments with 0.2% rhodamine beads.

In the first set of experiments, Ang II (0.6, 6, and 60 pmol) was injected unilaterally into the RVLM. The Ang II dose was selected randomly, and 2 doses were tested per animal (1 per side). Injections were separated by a minimum of 30 minutes. In a second set of experiments, the AT1 receptor antagonist losartan (1 nmol/side), kynurenic acid (3 nmol/side), or isotonic saline was injected bilaterally into the RVLM. Each drug was tested in a separate group of rats. In kynurenic acid experiments, a blood sample (0.2 μl) was also collected from the arterial line for analysis of plasma electrolytes with an I-STAT1 analyzer and 6 cartridge (Abbott; East Windsor, NJ). In preliminary experiments (n = 4), the dose of losartan completely eliminated the sympathoexcitatory response to 60 pmol Ang II in control rats: renal SNA (control, 50 ± 6%; losartan, 0; P < 0.001) and ABP (control, 20 ± 2 vs losartan, 0 ± 0 mm Hg; P < 0.001). The dose of kynurenic acid was based on a previous study.12 To control for any effect of AT1 receptor blockade in the circulation, a third experiment administered the identical dose of losartan intravenously (2 nmol/200 μl).

Spinal Cord Stimulation
Rats were prepared for renal SNA and ABP recording as described above. The spinal cord was stabilized with a spinal clamp, and a pneumothorax was performed to limit respiratory movements. Current trains (5 seconds, 1-ms pulse; 500 μA) of various frequencies (10, 20, and 50 Hz) were delivered through a monopolar stimulating electrode lowered into the T1 dorsolateral funiculus. To quantify changes in renal SNA, spike-triggered averages of renal SNA were constructed from 60 sweeps (0.5 Hz; 1-ms pulse) at different stimulus intensities (50, 100, and 300 μA).

RT-PCR Analysis of Ang II Receptors
Rats were deeply anesthetized with 5% isoflurane and perfused transcardially with cold oxygenated artificial cerebrospinal fluid (124 mmol/L NaCl, 26 mmol/L NaHCO3, 0.6 mmol/L NaH2PO4, 3 mmol/L KCl, 1.6 mmol/L MgCl2, 1.5 mmol/L CaCl2, 11 mmol/L glucose, pH 7.4). The brain stem was rapidly removed and sectioned at 300 μm in oxygenated artificial cerebrospinal fluid (4°C) using a vibratome. The RVLM was isolated under a microscope, immediately frozen in liquid nitrogen, and stored at −80°C. Additional samples were collected from the renal cortex.

Total RNA was isolated using TRizol (Invitrogen; Carlsbad, Calif). All RNA samples were treated with TURBO DNase (Ambion; Austin, Tex) to remove genomic DNA. Non–RT-PCR using 50 ng of total RNA and Taqman Gene Expression Assay for β-actin confirmed DNase treatment successfully removed genomic DNA because no signal was detected after 40 cycles. First-strand cDNA synthesis from total RNA was performed using SuperScript III First-Strand Synthesis SuperMixRT (Invitrogen). Taqman Gene Expression Assays (Applied Biosystems; Foster City, Calif) were used to determine the receptor expression of AT1α (Agtr1a, Rn01435427_s1), AT1β (Agtr1b, Rn02132799_m1), and AT2 (Agtr2, Rn00560677_s1). Gene expression was normalized to 3 endogenous control genes (β-actin, Rn00667869_m1; β-2 microglobulin (B2m, Rn00560865_m1), and hydroxymethylbilane synthase (Hmb2, Rn01421880_g1) and then quantified using the comparative Ct method (ΔΔCt).

Data Analysis
All data are expressed as mean ± SE. Changes in integrated SNA were calculated by subtracting background noise after hexamethonium (30 mg/kg IV). Successful recordings of renal or splanchnic SNA were not obtained in every experiment for technical reasons. For Ang II injections, the 1-second peak SNA and ABP response was compared with a 30-second baseline segment immediately before the injection. SNA responses to Ang II are only reported for those injections performed ipsilateral to the nerve isolation. Therefore, n values for ABP are much greater than SNA and are reported in the figure legends. Data from losartan and kynurenic acid experiments were averaged in 1-minute bins and compared with baseline values. For spinal cord stimulation studies, peak mean ABP (1 second) or renal SNA (10 ms) was compared with baseline values (30 seconds and 250 ms, respectively). All data were analyzed by a 1- or 2-way ANOVA with repeated measures when appropriate (time factor). All post hoc tests were performed with independent or paired t tests with a layered Bonferroni correction. A P < 0.05 was statistically significant.

Results
The RVLM was initially identified by the site that produced the largest increase in ABP in response to L-glutamate. As reported previously,11,13 injection of glutamate into the RVLM of rats drinking 0.9% NaCl versus water evoked a significantly larger increase in renal SNA (253 ± 4 versus 201 ± 6%, respectively; P < 0.01) and ABP (48 ± 2 versus 31 ± 3 mm Hg, respectively; P < 0.01). There were no differences in baseline mean ABP, heart rate, and renal or splanchnic SNA between groups despite a significant elevation in 24-hour fluid ingestion and daily sodium intake (Table). The elevation in sodium intake did not alter hematocrit or plasma Na+, K+, and Cl− concentrations.

Elevated Dietary Salt Enhances SNA and ABP Response to Microinjection of Ang II Into the RVLM
A major goal of the present study was to determine whether elevated dietary salt intake enhanced the SNA and ABP responses evoked by Ang II in the RVLM. Injection of Ang II into the RVLM produced dose-dependent increases in renal SNA and ABP of both groups (Figure 1). However, rats drinking 0.9% NaCl versus water displayed a significantly greater increase in renal SNA and ABP after injection of 0.6

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and 6 pmol Ang II (Figure 1). In fact, injection of 0.6 pmol Ang II in rats drinking 0.9% NaCl produced a similar increase in renal SNA and ABP of rats drinking water and receiving 6 pmol Ang II. There were no significant differences in renal SNA or mean ABP between groups at 6 pmol Ang II.

To determine whether the enhanced SNA and ABP responses to RVLIN injection of Ang II could be attributed to differences downstream of RVLIN cell bodies, descending sympathoexcitatory spinal pathways were activated by electric stimulation of the dorsolateral funiculus. Rats drinking water or 0.9% NaCl displayed frequency-dependent increases in mean ABP (Figure 2); however, there were no significant differences in the magnitude of these responses between groups (P>0.7 from overall ANOVA). Spike-triggered averaging of renal SNA indicated that both groups had stimulus-dependent increases in renal SNA (Figure 2), but the magnitude of these increases were not different between rats drinking water versus 0.9% NaCl (P>0.6 from overall ANOVA).

Elevated Dietary Salt Does not Change AT₁ mRNA Expression in the RVLIN

In a separate set of experiments, we examined whether elevated dietary salt altered expression of AT₁a, AT₁b, or AT₂ receptor levels in the RVLIN. AT₁a and AT₂ receptor mRNA were expressed in the RVLM but not AT1b mRNA (Ct values >35). However, there were no significant differences in mRNA levels of RVLM AT₁a and AT₂ receptors between rats drinking water versus 0.9% NaCl (Figure 3), despite a significant reduction of AT₁a mRNA in the renal cortex of rats drinking 0.9% NaCl. These observations were consistent regardless of the housekeeping gene used for normalization and calculation of the 2⁻¹⁰ΔCt values (data not shown for β2 microglobulin or hydroxymethylbilane synthase). Dietary salt did not alter the expression of any housekeeping gene (data not shown).

Bilateral Injection of Losartan, but not Kynurenic Acid, Lowers SNA and ABP in Rats Drinking 0.9% NaCl

A final set of experiments examined whether elevated dietary salt intake altered the contribution of AT₁ receptor activation

**Table. Characteristics of Rats Drinking Water or 0.9% NaCl for 14 Days**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Water</th>
<th>0.9% NaCl</th>
</tr>
</thead>
<tbody>
<tr>
<td>Body weight, g</td>
<td>375±6 (50)</td>
<td>383±4 (50)</td>
</tr>
<tr>
<td>Daily food intake, g</td>
<td>28±1 (50)</td>
<td>29±1 (50)</td>
</tr>
<tr>
<td>Daily ingested fluid, mL</td>
<td>36±1 (50)</td>
<td>57±2* (50)</td>
</tr>
<tr>
<td>Daily Na⁺ intake, mEq/day</td>
<td>2.6±0.1 (50)</td>
<td>11.6±0.3* (50)</td>
</tr>
<tr>
<td>Baseline mean ABP, mm Hg</td>
<td>118±2 (45)</td>
<td>116±2 (45)</td>
</tr>
<tr>
<td>Baseline heart rate, bpm</td>
<td>397±5 (45)</td>
<td>391±5 (45)</td>
</tr>
<tr>
<td>Baseline renal SNA, mV</td>
<td>0.115±0.009 (42)</td>
<td>0.123±0.009 (43)</td>
</tr>
<tr>
<td>Baseline splanchnic SNA, mV</td>
<td>0.058±0.009 (15)</td>
<td>0.046±0.004 (16)</td>
</tr>
<tr>
<td>Plasma Na⁺, mEq/L</td>
<td>140±1 (5)</td>
<td>139±1 (5)</td>
</tr>
<tr>
<td>Plasma K⁺, mEq/L</td>
<td>4.7±0.2 (5)</td>
<td>5.0±1.0 (5)</td>
</tr>
<tr>
<td>Plasma Cl⁻, mEq/L</td>
<td>109±2 (5)</td>
<td>107±1 (5)</td>
</tr>
<tr>
<td>Hematocrit, %</td>
<td>43±1 (5)</td>
<td>44±1 (5)</td>
</tr>
</tbody>
</table>

Values (mean±SE) were pooled from all experiments including the online supplement. Value in parentheses indicates number of animals. Note that n values for renal and splanchnic SNA are smaller than those for mean ABP and heart rate because SNA recordings were not obtained in all experiments for technical reasons. Plasma electrolytes were analyzed in animals receiving injection of kynurenic acid.

*Indicates significant difference from rats drinking water (P<0.001).
Discussion
Excess dietary salt intake contributes to or exacerbates many forms of hypertension, including Ang II–dependent models, by increasing SNA and vasomotor tone.9,10 Evidence in anesthetized animals indicates that the elevated SNA and ABP depend, in part, on AT1 receptor activation in the RVLM.5,6 Despite the interaction among dietary salt, Ang II, and the RVLM in the pathogenesis of hypertension, no studies have directly examined whether dietary salt by itself alters Ang II–dependent regulation of RVLM neurons, SNA, and ABP. The present study provides several novel observations: (1) elevated dietary salt intake enhances the SNA and ABP responses to exogenously applied Ang II in the RVLM; (2) these exaggerated sympathoexcitatory responses were not associated with changes in RVLM AT1a mRNA levels or the excitability of downstream sympathoexcitatory pathways; and (3) blockade of RVLM AT1, but not ionotropic glutamate, receptors significantly lowered SNA and ABP in rats drinking 0.9% NaCl. Altogether, these findings suggest that elevated dietary salt enhances the sympathoexcitatory actions of Ang II in the RVLM via changes in the intrinsic properties of RVLM neurons. Furthermore, excess dietary salt intake activates an angiotensinergic pathway to the RVLM to regulate baseline SNA and ABP.

Injection of Ang II into the RVLM produces an AT1 receptor–mediated increase in SNA and ABP.1,2 These sympathoexcitatory effects are likely attributable to direct postsynaptic actions because Ang II–induced depolarization of bulbospinal RVLM neurons persists after synaptic blockade.17,18 Although Ang II has been reported to presynaptically modulate glutamate release in other brain regions,2 the available data for the RVLM5,19 including this study, indicate that AT1 and glutamatergic receptors independently regulate the activity of RVLM neurons. Interestingly, the present

Histology
All injection sites were centered in the RVLM defined as the triangular region located 0 to 600 μm caudal to the caudal pole of the facial nucleus and bordered dorsally by nucleus ambiguus, medially by the inferior olive or pyramidal tracts, and laterally by the spinal trigeminal nucleus (Figure S3).

A

Figure 3. Quantitative RT-PCR of AT1a and AT2 receptor expression in the RVLM and AT1a receptor expression in the renal cortex of rats drinking water (open; n=5) or 0.9% NaCl (filled; n=5) for 14 days. Values were calculated using the $2^{-\Delta\Delta Ct}$ method and normalized to β-actin. * indicates significant difference from rats drinking water (P<0.01).

B

Figure 4. Mean ABP, renal SNA, and splanchnic SNA response to bilateral injection of losartan (n=11 per group; A) or kynurenic acid (n=5 per group; B) into the RVLM of rats drinking water (●) or 0.9% NaCl (●) for 14 days. Injections were performed at 0 minutes and separated by 2 to 3 minutes. SNA values are based on a smaller number of animals (losartan, n=7 per group; kynurenic acid, n=4). * indicates significant difference between groups (P<0.05).
findings provide strong evidence that excess dietary salt intake enhances the sympathoexcitatory actions of Ang II (and L-glutamate). In fact, the plot of the SNA and ABP responses to Ang II (Figure 1) suggest the sensitivity, not the maximal response, has changed. These exaggerated responses cannot be attributed to an alteration in downstream spinal sympathetic pathways or neuroeffector junctions because electric stimulation of the dorsolateral funiculus produced similar increases in both renal SNA and ABP of rats drinking water versus 0.9% NaCl. Moreover, several investigators have demonstrated that dietary salt does not alter vascular reactivity.12,13,15

The mechanism(s) underlying the greater responsiveness of RVLM neurons during elevated dietary salt intake is not known. Although changes in dietary salt intake have been reported previously to alter brain AT1 mRNA levels,20–22 analysis of RVLM AT1a (and AT1b) mRNA levels in the present study revealed no difference between rats drinking water versus 0.9% NaCl. Thus far, we have been unsuccessful extending this observation to the protein level because Western blot analyses reveal dense bands at the predicted molecular weight of ~45 kDa in the RVLM and renal cortex, but these bands were still present in the renal cortex of AT1a−/− mice and the brain region dorsolateral to the RVLM in rats, where no Ang II binding has been reported23 (Figure S4). A second noteworthy observation is that excess dietary salt intake enhances the sympathoexcitatory and sympathoinhibitory responses to a number of neurotransmitters (>5) in the RVLM.11–13 Therefore, elevated dietary salt either alters the expression of every receptor or, more likely, alters the intrinsic excitability of RVLM sympathetic-regulatory neurons. Such possibilities may include a general change in membrane conductance (ie, potassium) because this would permit enhanced responses to both excitatory and inhibitory synaptic inputs.

Blockade of AT1 receptors in the RVLM decreases ABP in models of experimental hypertension including Dahl salt-sensitive hypertensive rats.5,6 The present findings indicate that excess dietary salt by itself increases the contribution of RVLM AT1 receptor activation to the maintenance of baseline SNA and ABP. This seems at odds with a previous report that systemic administration of losartan decreases ABP in animals maintained on a low-salt but not a high-salt diet.24 However, the dose of systemic losartan needed to block RVLM AT1 receptors has not been defined, and the hypertensive actions of systemic losartan in animals on a low-salt diet may be attributed to tissues other than the RVLM. Despite a greater activation of AT1 receptors in the RVLM of rats drinking 0.9% NaCl, baseline SNA and ABP were not different between groups. The greater AT1 receptor activation may be offset by a withdrawal of other excitatory inputs or an increase in inhibitory input. The transient increase in SNA and ABP observed in rats drinking water or 0.9% NaCl after losartan injection is likely attributed to the potassium salt. However, SNA tended to remain above baseline levels in control rats. The explanation for this observation is not clear, but previous studies have indicated that bilateral injection of losartan into the RVLM did not change8,19 or increased25 SNA or ABP in normal animals.

One intriguing question that arises from the current findings is how does excess dietary salt increase AT1 receptor activation in the RVLM? The ability of AT1 receptor blockade in the RVLM to lower SNA and ABP is reminiscent to those findings reported in water-deprived rats with elevated plasma sodium concentration.7 Although we did not observe differences in plasma sodium concentration between groups, excess dietary salt intake has been reported to increase plasma sodium concentration in rodents at night26 and humans.27 Interestingly, preliminary data from our laboratory indicate that lesion of osmosensitive sites in ventral forebrain lamina terminals prevents the salt-induced changes in the responsiveness of RVLM neurons.28 Yet, how does activation of osmotically sensitive sites in the forebrain increase AT1 receptor activation in the RVLM? First, forebrain osmosensitive regions densely innervate neurons of the hypothalamic paraventricular nucleus (PVH),29 and acute hyperosmolality increases the discharge of RVLM-projecting PVH neurons.30 Second, previous studies have reported that PVH neurons are Ang II immunoreactive,31 and the sympathoexcitatory response produced by disinhibition of the PVH is significantly attenuated by blockade of AT1 receptors in the RVLM.19 Thus, excess dietary salt may activate osmosensitive neurons in the forebrain lamina terminals to drive an angiotensinergic pathway from the PVH to the RVLM. At this time, we cannot exclude the possibility that dietary salt activates other afferent pathways or sources of Ang II input to the RVLM.

Previous studies have reported that sympathetic outflow is inversely related to the level of dietary salt intake,32–35 but the present study did not observe any difference in baseline renal and splanchnic SNA or ABP between groups. The reason for the differences between our findings and previous studies is unclear but may be related to the experimental conditions or species studied. However, our findings are consistent with a recent report in which chronic recordings of renal SNA and ABP in awake rabbits did not reveal any effect of dietary salt on baseline sympathetic vasomotor activity or ABP.36 Regardless of the effect of dietary salt intake on baseline SNA, the present findings together with previous reports11–13 suggest that dietary salt intake adjusts the gain or responsiveness of RVLM neurons. Therefore, a small increase in excitatory drive to the RVLM now produces an exaggerated increase in SNA and ABP. In fact, activation of somatic afferents produces a significantly greater increase in ABP of animals maintained on a high-salt diet;22 this response depends on glutamatergic neurotransmission in the RVLM.16 Similarly, dietary salt has been reported to potentiate the sympathoexcitatory response to hyperinsulinemia.37 These observations indicate that these functional changes in the responsiveness of RVLM sympathoexcitatory neurons have physiological significance in the regulation of SNA and ABP. Future studies will need to extend these observations and identify the level of dietary salt intake needed for such an effect; however, we suspect that the responsiveness of RVLM neurons is proportional to the level of dietary salt rather than a threshold because both dietary salt restriction and loading decreases or enhances, respectively, the pressor responses evoked from the RVLM.12
Perspectives
The present findings identify a new factor that influences the potency of the brain renin-angiotensin system in sympathetic-regulatory networks. This is particularly intriguing because excess dietary salt inhibits peripheral renin secretion, thereby suggesting that the excess dietary salt differentially modulates the activity of the peripheral versus brain renin-angiotensin system. Based on these findings, administration of AT1 receptor blockers at doses that functionally block RVLM AT1 receptors should reduce SNA and ABP in salt-sensitive hypertensive subjects and experimental models of hypertension. In fact, blockade of AT1 receptors in the RVLM of the Dahl salt–sensitive hypertensive rats lowered ABP. Subsequent identification of the factors that underlie the ability of dietary salt to alter the intrinsic excitability of RVLM sympathetic neurons during changes in dietary salt intake may represent a novel therapeutic target for the treatment of salt-sensitive hypertension.

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Disclosures
None.

References

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Julye M. Adams, John J. McCarthy, and Sean D. Stocker
Department of Physiology, University of Kentucky, Lexington, KY 40536-0298 USA

Address correspondence to:
Sean D. Stocker, Ph.D.
Assistant Professor
Department of Physiology, University of Kentucky
800 Rose St. MS-508
Lexington, KY 40536-0298
Email: sean.stocker@uky.edu
Phone: 859-323-4344
Fax: 859-323-1070
Western blot analysis

Protein was isolated from RVLM, an area immediately dorsolateral to the RVLM, and renal cortex using TRIzol according to manufacturer’s directions. These samples were identical to those used for RT-PCR analysis of AT1a, AT1b, and AT2 receptors in the manuscript. The protein concentration of each sample was determined using the Bio-Rad DC protein assay (Bio-Rad Laboratories, Hercules, CA) with bovine serum albumin as a standard. Ten micrograms of each sample was separated by SDS-PAGE (10% gel) and then transferred to nitrocellulose membrane (0.2 m) (Bio-Rad, Hercules, CA). The membrane was incubated in blocking buffer (5% nonfat dry milk in TBS plus 0.1% Tween-20 [TBS-T]) for 1 hr at room temperature and then incubated in blocking buffer overnight at 4°C with a 1:500 dilution of rabbit polyclonal AT1 antibody (N-10; sc-1173; Santa Cruz Biotechnology, Inc. Santa Cruz, CA.). To determine specificity of the AT1 antibody, a second membrane was incubated with the AT1 antibody pre-absorbed with five-fold excess of a blocking peptide. After the overnight incubation, each membrane was washed for 5 min four times in TBS-T and then incubated with anti-rabbit horseradish peroxidase (HRP)-conjugated secondary antibody (10 ng/mL) for 1 hr at room temperature in blocking buffer. Following this incubation, the membranes were washed again in TBS-T as described above and then incubated for 5 min in ECL Plus (GE Healthcare, Piscataway, NJ) and exposed to X-ray film. As an additional control, protein was isolated from the renal cortex of wild-type and AT1a-/-mice using TRIzol and subject to Western blot analysis as described above. Kidneys from wild-type and
AT1a-/- mice were generously provided by Drs. Lisa Cassis and Alan Daugherty (University of Kentucky). Results are presented in Supplemental Figure S4.
Figure S1. Individual examples of ABP, mean ABP, and renal SNA after microinjection of (A) 1.0 nmol L-glutamate, (B) 0.6 pmol ANG II, (C) 60 pmol ANG II into the RVLM of rats drinking water (LEFT) or 0.9% NaCl (RIGHT). Summary data presented in Figure 1. ▼, microinjection.
Figure S2. Mean ± SE of mean ABP, renal SNA, and heart rate of rats drinking water (○, n=3-4) or 0.9% NaCl (●, n=3-4) for 14 days during (A) bilateral injection of isotonic saline into the RVLM or (B) intravenous injection of losartan (2nmol / 0.2 mL). Injection was performed at time = 0 min. Mean ABP, renal SNA or heart rate of both groups was not altered by either injection.
Figure S3. Schematic drawings of RVLM injection sites for rats drinking water (○) or 0.9% NaCl (●) and receiving an injection of (A) ANG II and (B) losartan or kynurenic acid. Kynurenic acid injection sites are denoted by ▽ or ▼. Sections represent -11.6 (top) and -11.9 (bottom) in reference to bregma. IO indicates inferior olive; p, pyramidal tracts; NA, nucleus ambiguus; ST, spinal trigeminal nucleus.
Figure S4. Western blot analysis of AT1 receptor protein in the RVLM (R), area that is dorsolateral to the RVLM (A), and renal cortex (C). Lanes 1-3 represent bands visualized from each of the 3 different samples in control rats using a rabbit anti-AT1 receptor antibody (1: 500). Note each of these bands were eliminated by preabsorption of the antibody with the blocking peptide. Lanes 7-10 represent bands observed with the same antibody from the renal cortex of wild-type (WT) and AT1a -/- mice (KO). Note that the bands are at the same relative intensity regardless of the tissue examined, and bands were present in tissue samples from the area dorsolateral to the RVLM where no ANG II binding has been reported. Similar bands were observed in WT versus KO mice. Detailed methods are described in the online supplement at “Western Blot Analysis”.