Chronic excitotoxic lesion of the dorsal raphe nucleus induces sodium appetite

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Abstract

We determined if the dorsal raphe nucleus (DRN) exerts tonic control of basal and stimulated sodium and water intake. Male Wistar rats weighing 300-350 g were microinjected with phosphate buffer (PB-DRN, N = 11) or 1 µg/0.2 µl, in a single dose, ibotenic acid (IBO-DRN, N = 9 to 10) through a guide cannula into the DRN and were observed for 21 days in order to measure basal sodium appetite and water intake and in the following situations: furosemide-induced sodium depletion (20 mg/kg, sc, 24 h before the experiment) and a low dose of dietary captopril (1 mg/g chow). From the 6th day after ibotenic acid injection IBO-DRN rats showed an increase in sodium appetite (12.0 ± 2.3 to 22.3 ± 4.6 ml 0.3 M NaCl intake) whereas PB-DRN did not exceed 2 ml (P < 0.001). Water intake was comparable in both groups. In addition to a higher dipsogenic response, sodium-depleted IBO-DRN animals displayed an increase of 0.3 M NaCl intake compared to PB-DRN (37.4 ± 3.8 vs 21.6 ± 3.9 ml 300 min after fluid offer, P < 0.001). Captopril added to chow caused an increase of 0.3 M NaCl intake during the first 2 days (IBO-DRN, 33.8 ± 4.3 and 32.5 ± 3.4 ml on day 1 and day 2, respectively, vs 20.2 ± 2.8 ml on day 0, P < 0.001). These data support the view that DRN, probably via ascending serotonergic system, tonically modulates sodium appetite under basal and sodium depletion conditions and/or after an increase in peripheral or brain angiotensin II.

Key words
- Sodium appetite
- Water intake
- Dorsal raphe nucleus
- Serotonergic system
- Ibotenic acid

Introduction

The organum vasculosum laminae terminalis, median preoptic nucleus and subfornical organ (SFO), structures participating in hydroelectrolytic and cardiovascular homeostasis, receive input from serotonergic neurons of the dorsal raphe nucleus (DRN) (1-6). There is a reciprocal connection linking DRN and SFO (7-9) which suggests that the DRN participates in cardiovascular as well as in extracellular fluid (ECF) volume and electrolyte homeostasis. Angiotensinergic stimulation of the SFO elicits electrophysiological alterations in DRN serotonergic neurons (9). On the other hand, isosmotic
hypovolemia induced by hemorrhage evokes changes in the electrical activity of DRN serotonergic neurons connected with the SFO (10). In agreement with this, Vivas’ group (11) reported c-Fos expression in DRN serotonergic neurons and laminae terminalis structures in response to salt intake under basal conditions or after sodium depletion induced by peritoneal dialysis. Furthermore, the intracerebroventricular (icv) administration of serotonin (5-HT) and of the 5-HT2A/C receptor agonists induces an increase in renal sodium excretion (12). In addition, electrolytic lesions of the DRN drastically reduce the plasma levels of atrial natriuretic peptide (ANP), both in basal and ECF volume expansion situations (6,12,13). In this context, it is important to note the very important role of ANP in sodium excretion and salt appetite (5,6,13-19).

Renal afferent information about tubular sodium load and plasma volume is transmitted toward forebrain areas related to angiotensin conversion (ANG I to ANG II) and subsequent angiotensin receptor activation (20). DRN afferent fibers arise from these areas (especially the SFO), probably involved in a feedback loop regulating sodium satiety and contributing to ECF volume and cardiovascular homeostasis (7,9,21).

Taken together, this evidence supports our view that central control of sodium satiety and excretion is regulated by the serotonergic system that originates from the midbrain, but only one study acknowledged that the suppression of ascending serotonergic pathways through DRN electrolytic lesion evokes an increase in sodium appetite (21). Corroborating these findings, other investigators showed that basal c-Fos expression in serotonergic neurons of the DRN decreased after sodium depletion induced by peritoneal dialysis and increased after spontaneous and induced sodium intake, suggesting that there is a tonic inhibition of sodium appetite by serotonergic cells of this nucleus (11). Although we hypothesized the same, when working with DRN electrolytic lesion it is necessary to limit the lesion to the neuronal cell body located in the ventromedial portion of the DRN and to perform longer observations (21).

Thus, anatomic and physiological evidence clearly shows the role of the DRN in integrating viscerosensory input and forebrain signals concerning cardiovascular and hydroelectrolyte homeostasis. In this context, the objective of the present study was to investigate the influence of excitotoxic DRN lesion by local ibotenic acid microinjection on sodium appetite both in the basal condition and in the presence of natriorexigenic paradigms, such as furosemide-induced sodium depletion and low-dose dietary supplementation with captopril.

Material and Methods

Animals

Male Wistar rats weighing 300-350 g were maintained in a room with lights on from 7:00 to 19:00 h, controlled temperature at 25°C and free access to Purina chow and water. Before brain stereotaxic surgery the rats were housed in metabolic cages for at least 5 days. All the experimental protocols and animal procedures were carried out in accordance with current Brazilian legislation.

Ibotenic lesions

Rats were anesthetized with sodium pentobarbital (Hypnol 3%, Fontoveter, Itapira, SP, Brazil, 50 mg/kg body weight, ip) and a stainless steel guide cannula (0.6-mm external diameter) was stereotactically implanted 0.2 mm above the DRN. The following coordinates were used: anteroposterior, 7.6-7.8 mm posterior to bregma; lateral, 0.0 mm from midline; vertical, 6.4-6.6 mm from skull calvaria (22). Ibotenic acid microinjections (1 µg/0.2 µl dissolved in sterile 0.2 M so-
Sodium phosphate buffer, pH 7.4) were performed during a period of 1 min. Sham-lesioned rats were microinjected with 0.2 µl phosphate buffer. A prophylactic dose of penicillin was administered to operated rats (30,000 IU, im; Fort Dodge Saúde Animal Ltda., Campinas, SP, Brazil). Data from rats with lesions outside the dorsoventromedial region of the DRN were not included in the statistical analysis.

Experimental procedures

DRN-excitotoxic-lesioned rats microinjected with ibotenic acid (IBO-DRN) or sham-lesioned rats microinjected with sodium phosphate buffer (PB-DRN) were used in three experimental protocols:

Group 1. IBO-DRN (N = 10) and PB-DRN (N = 11) rats were maintained in metabolic cages under basal conditions to measure water and 0.3 M NaCl consumption for 21 days following surgery.

Group 2. On day 21 after microinjection, IBO-DRN (N = 9) and PB-DRN (N = 11) rats were submitted to sodium and water depletion induced by 20 mg/kg furosemide, sc (Lasix, Aventis Pharma, Suzano, SP, Brazil). During the 24 h following furosemide administration a low-sodium diet was offered to the animals (23). Distilled water and 0.3 M NaCl were then supplied and the consumption of these fluids was recorded hourly for 5 h and at the end of 24 h (Figure 1).

Group 3. A low dose (1 mg/g of chow) of dietary captopril (Bristol-Myers Squibb, São Paulo, SP, Brazil) was added every day to the chow from the 6th to the 11th days after surgery (PB-DRN, N = 11 and IBO-DRN, N = 10). This model promotes a systemic rise in plasma ANG I levels, which results in an elevated ANG II synthesis in circumventricular forebrain structures underlying the subsequent natriorexigenic response (24-29). The consumption of 0.3 M NaCl and water was determined before (3-day pretreatment period), during (captopril treatment) and after treatment (9-day recovery period).

Histological analysis

Rats were anesthetized and transcardially perfused with 4% paraformaldehyde to identify the lesion site. Brains were removed and serial coronal sections were examined after staining with crystal violet by the Holzer method.

Statistical analysis

Differences between the IBO-DRN and PB-DRN groups were analyzed by repeated measures two-way ANOVA followed by the Newman-Keuls test, with the level of significance set at 5%.

Results

The experimental protocol is presented schematically in Figure 1. Midbrain histological analyses of IBO-DRN rats revealed typical lesions in the ventromedial portion of the DRN. Fibrillar astrocytes were evident along the anteroposterior axis of this nucleus (Figure 2).

Sodium appetite was increased in IBO-DRN rats compared to PB-DRN rats throughout the experimental period, especially after 6 days post-lesion (P < 0.01). From day 11 on, 0.3 M NaCl intake reached a level always higher than 18 ml (Figure 3A), what persisted up to the 35th day post-lesion (data...
not shown). In addition, most of the IBO-DRN rats showed a drive to eat salt, which was evident when a salt lump was offered through the metabolic cage door. This phenomenon was frequently observed in rats which had been previously offered the salt lump. Water intake did not differ between groups during the experimental period (Figure 3B).

Furosemide-induced sodium and water depletion enhanced the natriorexigenic response of IBO-DRN, which became significantly higher than that of PB-DRN only after 240 min of fluid offer, remaining at this level for 24 h (9.8 ± 1.0, 10.2 ± 1.0, and 21.6 ± 3.9 vs 13.1 ± 0.98, 15.3 ± 1.1, and 37.4 ± 3.8 ml, at 240, 300, and 1440 min in PB-DRN and IBO-DRN, respectively, P < 0.04; Figure 4A). Renal sodium and water loss-induced volume depletion evoked an increased water intake in IBO-DRN animals during the experimental period up to 24 h after fluid presentation (P < 0.002; Figure 4B).

Low-dose captopril treatment promoted a still higher increase in 0.3 M NaCl intake in IBO-DRN animals and was significantly different compared to the PB-DRN group on the first 2 days of treatment (13.6 ± 1.7 and 20.8 ± 2.2 vs 33.8 ± 4.3 and 32.5 ± 3.4 ml on the 1st and 2nd days in PB-DRN and IBO-DRN, respectively, P < 0.002; Figure 5A). On the first day after the end of captopril treatment there was an abrupt fall of 0.3 M NaCl intake in the IBO-DRN group from 30.1 ± 3.3 to 7.9 ± 3.3 ml. In contrast, PB-DRN rats showed a slight reduction compared to IBO-DRN rats (from 27.5 ± 3.0 to 19.4 ± 4.1 ml), with higher levels of 0.3 M NaCl intake than IBO-DRN at this time point. The 0.3 M NaCl intake was gradually recovered after day 12 (6 days after the end of captopril treatment) in both groups, returning to levels similar to those observed before captopril treatment. There was no difference in water intake between the groups during 14 days, except on the first two days after captopril treatment interruption when an increase in this parameter was observed in IBO-DRN rats (Figure 5B).

Discussion

Our results show that DRN lesions disrupt the functional integrity of central neural pathways that normally exert a negative drive leading to salt intake inhibition. It has been postulated that the DRN serotonergic neurons receive viscerosensory inputs concerning renal sodium load and/or ECF volume variations to trigger homeostatic regulation of sodium appetite (6,21,30). This view has
been shared by other investigators who have shown increased c-Fos expression in DRN serotonergic neurons after salt intake under both basal and sodium depletion conditions (11).

Low-dose captopril treatment resulted in a higher saline intake by ibotenic DRN-lesioned rats, suggesting that there is either an enhanced capacity of ANG I conversion to ANG II in forebrain structures, or an increased sensitivity to ANG II-evoked sodium appetite (24,25,27-29). On this basis and regarding reciprocal relationships between SFO and DRN, we postulated that the presumptive angiotensinergic sensitivity of the SFO is enhanced following the failure of inhibitory signals arising from the midbrain (7,10,28).

Although we did not investigate the forebrain angiotensinergic sensitivity, we recently demonstrated that experiments involving an increase in plasma ANG II levels exacerbate the sodium appetite of ibotenic-lesioned rats (31). Hence, the view that suppression of neural pathways originating in the DRN reduces the threshold of the sodium appetite response through an increase in angiotensinergic sensitivity of SFO neurons should be considered. That is, lesioned rats putatively respond to the natriorexigenic stimuli (induced by high plasma levels of ANG I or ANG II) which would be subthreshold for sham-lesioned rats.

Also, we did not rule out the possibility that the deficit in the ascending regulation of sodium appetite is due, at least in part, to disruption of interconnections between the DRN and lateral parabrachial nucleus (LPBN) (32,33). This presumptive interaction was suggested by Lind’s work (7), which presupposed that viscerosensory information relayed from the pontine nucleus towards the raphe would be transmitted to the SFO through ascending serotonergic pathway from the DRN. The LPBN is an important pontine circuit that receives homeostatic viscerosensory signals from the right atrium and vena cava stretch receptors and arterial and atrial baroreceptors implicated in the control of extracellular volume and blood pressure (34).
The blockade of 5-HT2A/C receptors in the LPBN exacerbates the ingestion of hypertonic saline induced by icv administration of ANG II or by a combination of furosemide + low dose of captopril (23). Microinjection of the AT1 antagonist losartan into the SFO prevented the additional salt intake induced by methysergide administration in the LPBN of rats submitted to furosemide + captopril treatment (35). Later, Menani and colleagues (36,37) showed a widespread involvement of the inhibitory serotonergic mechanism in the LPBN. Thus, bilateral injections of serotonergic receptor antagonist methysergide amplified hypertonic saline intake after furosemide-induced sodium depletion, water deprivation or central cholinergic stimulation.

Corroborating data obtained by electrolytic DRN lesion (21), our results reveal that excitotoxic damage of DRN neurons induces a rise in basal NaCl intake. In that study (21), Olivares and co-workers made short-term observations (3 days only) and their findings correlated with reduced ANP level in electrolytic DRN-lesioned rats under basal and volume expansion conditions (13). Although the reduced ANP levels could explain the increase in sodium appetite, it has been demonstrated that renal sodium excretion normalizes 7 days after electrolytic lesion of the DRN, possibly concomitant with a restoration of ANP secretion (6,13).

Therefore, on the basis of the present data, the above-mentioned observation conflicts with the hypothesis that DRN lesion-evoked sodium appetite occurs as a result of reduced ANP secretion. We suggested that the suppression of ascending serotonergic pathways exacerbates sodium appetite through a disinhibitory mechanism in the forebrain, possibly within the SFO, whose target neurons would be ANG II-sensitive. However, the precise mechanism that underlies forebrain mediation of serotonergic sodium satiety is unknown.

Our data suggest that an ascending circuit, possibly serotonergic, originating in the DRN, integrates viscerosensory and forebrain signals for sodium appetite homeostasis. The results obtained under basal conditions allow us to postulate that the DRN neural circuit constitutes an integrative system that tonically triggers modulation of the outputs to the forebrain to induce salt satiety. Furthermore, disruption of the reciprocal connection between DRN and LPBN could contribute to the deficit in forebrain signaling for sodium appetite modulation.

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References


