High Salt Intake Increases Blood Pressure via BDNF-Mediated Downregulation of KCC2 and Impaired Baroreflex Inhibition of Vasopressin Neurons

Highlights

- Chronic high salt intake reduces chloride gradient in vasopressin neurons
- BDNF-TrkB activation causes KCC2 downregulation and collapse of chloride gradient
- High salt intake abolishes baroreceptor inhibition of vasopressin neurons
- Circulating vasopressin mediates high blood pressure during high salt intake

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In Brief

High salt consumption is causally linked to hypertension with unclear etiology. Choe et al. show that chronic high salt modifies a hypothalamic circuit, leading to excessive release of the antidiuretic hormone vasopressin. The resulting peripheral vasoconstriction increases blood pressure.
High Salt Intake Increases Blood Pressure via BDNF-Mediated Downregulation of KCC2 and Impaired Baroreflex Inhibition of Vasopressin Neurons

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SUMMARY

The mechanisms by which dietary salt promotes hypertension are unknown. Previous work established that plasma [Na+] and osmolality rise in proportion with salt intake and thus promote release of vasopressin (VP) from the neurohypophysis. Although high levels of circulating VP can increase blood pressure, this effect is normally prevented by a potent GABAergic inhibition of VP neurons by aortic baroreceptors. Here we show that chronic high salt intake impairs baroreceptor inhibition of rat VP neurons through a brain-derived neurotrophic factor (BDNF)-dependent activation of TrkB receptors and down-regulation of KCC2 expression, which prevents inhibitory GABAergic signaling. We show that high salt intake increases the spontaneous firing rate of VP neurons in vivo and that circulating VP contributes significantly to the elevation of arterial pressure under these conditions. These results provide the first demonstration that dietary salt can affect blood pressure through neurotrophin-induced plasticity in a central homeostatic circuit.

INTRODUCTION

High levels of dietary salt intake can significantly increase plasma sodium concentration and contribute to the development of salt-dependent hypertension (He et al., 2013; He and Macgregor, 2012; Schmidlin et al., 2007). However, the central mechanisms by which excess sodium can increase blood pressure (BP) remain poorly defined. Previous work has shown that a rise in plasma sodium can excite hypothalamic magnocellular neurosecretory cells (MNCs) that release the antidiuretic and vasoconstrictor hormone vasopressin (VP) (Bourque, 2008; Voisin and Bourque, 2002). Although VP can enhance BP when infused systemically (Fujisawa et al., 2012), increases in BP normally activate arterial baroreceptors (BR) that inhibit VP MNCs via GABA$_A$ receptors (GABA$_AR$) (Cunningham et al., 2002; Renaud et al., 1988). In principle, this negative feedback regulation of MNCs should mitigate the involvement of circulating VP in the development of hypertension. However, a recent study has shown that rat MNCs display a collapse in the transmembrane chloride (Cl$^-$) gradient required for inhibitory GABA$_AR$ signaling after chronic salt loading (Kim et al., 2011). A similar effect can be induced by a high-salt diet in uni-nephrectomized rats treated with deoxycorticosterone acetate (DOCA), where a weakening of BR-mediated inhibition is also associated with a VP-dependent increase in BP (Kim et al., 2013). These observations indicate that plastic changes in the BR-mediated control of MNCs can allow these cells to participate in the regulation of BP. However, it remains unknown if high dietary salt intake can by itself mediate a VP-dependent form of hypertension, and the signaling mechanisms responsible for state-dependent changes in the BR-mediated control of VP MNCs are unknown.

Recent studies have shown that a weakening of GABA$_AR$-mediated inhibition caused by a collapsed Cl$^-$ gradient can emerge under several pathological conditions, including chronic pain (Coull et al., 2003), epilepsy (Huberfeld et al., 2007), stress (Hewitt et al., 2009), and spasticity following spinal cord injury (Boulenguez et al., 2010). In each instance experiments revealed that the effect was caused by a downregulation of the expression or activity of the K$^+$/Cl$^-$ co-transporter 2 (KCC2), a molecule that maintains the low levels of intracellular [Cl$^-$] required for inhibitory GABA$_AR$ signaling (Ferrini and De Koninck, 2013). The expression of KCC2 is tightly linked to the activity of tropomyosin-related kinase B (TrkB) receptors, whose activation can suppress KCC2 transcription (Rivera et al., 2004). Although TrkB receptors can be activated by several ligands (McNamara and Scharfman, 2012; Yoshii and Constantine-Paton, 2010), recent studies have shown that the potent TrkB agonist brain-derived neurotrophic factor (BDNF) (Lu, 2003) is a common mediator of TrkB activation and KCC2 downregulation in central neurons (Boulenguez et al., 2010; Coull et al., 2005; Huang et al., 2013; Molinari et al., 2009). Moreover, BDNF is highly expressed in...
VP MNCs (Aliaga et al., 2002; Arancibia et al., 2007; Castren et al., 1995), and in vivo experiments have shown that the dendrites of these neurons can release BDNF in response to electrical activity induced by a systemic increase in plasma [Na⁺] (Arancibia et al., 2007). In this study we therefore investigated whether chronic high dietary salt intake can provoke VP-dependent hypertension due to a BDNF-TrkB-KCC2-mediated weakening of BR inhibition of VP MNCs.

RESULTS

SL Depolarizes EGABA and Eliminates Inhibitory Tone

To confirm that chronic high salt intake causes a collapse of Cl⁻ gradient in MNCs (Kim et al., 2011), we first examined the voltage dependence of GABAAR-mediated postsynaptic potentials (PSPs) using sharp electrode intracellular recordings in hypothalamic explants prepared from euhydrated (EU) rats or animals provided with 2% NaCl as drinking solution for 7 days (SL; salt loading). This treatment resulted in a significant increase in plasma osmolality (Figure S1), as reported previously (Kim et al., 2011). Recordings were obtained from the VP-rich zone of the supraoptic nucleus (see Experimental Procedures), and DNQX (20 μM) was present to block fast ionotropic glutamatergic transmission. The average input resistances of recorded MNCs from EU and SL rats were 167.6 ± 14.8 MΩ (n = 27) and 116.6 ± 7.9 MΩ (n = 33), respectively (p = 0.0016). Spontaneous PSPs (sPSPs) recorded under these conditions were completely eliminated by the addition of GABAAR blockers bicuculline (10 μM; not shown) or gabazine (1 μM; Figure S2), confirming their dependence on these receptors. As illustrated in Figure 1A, GABAAR-mediated sPSPs recorded in MNCs from EU preparations were consistently hyperpolarizing at voltages near action potential (AP) threshold (i.e., rheobase ~45 mV), reversing polarity at ~57.2 ± 1.5 mV (n = 14, six rats). In contrast, sPSPs recorded in MNCs from SL preparations were generally depolarizing at voltages near AP threshold and reversed at ~36.4 ± 3.9 mV; n = 10, three rats; p = 0.000012 versus EU control).

We next examined the value of E_GABA determined from the voltage dependence of PSPs evoked in MNCs by stimulation of the nucleus of the diagonal band of Broca (DBB), a site that relays the inhibitory effect of BRs onto VP MNCs in the supraoptic nucleus (Cunningham et al., 2002; Renaud et al., 1988). Electrical stimulation of the DBB caused prominent gabazine-sensitive PSPs in MNCs (Figure S2) and the average value of E_GABA determined from these responses was significantly more positive in explants prepared from SL rats (~35.3 ± 2.9 mV; n = 23, eight rats) than EU rats (~59.4 ± 1.6 mV; n = 13, seven rats; p = 0.0000015; Figure 1B). Identical results were obtained using gramicidin perforated patch recordings from identified VP neurons in angled hypothalamic slices prepared from transgenic Wistar rats expressing enhanced green fluorescent protein (GFP) in VP MNCs (Figures 2A–2D) and by intracellular recordings from the VP-rich zone of the supraoptic nucleus MNCs in Figure 1. SL Shifts E_GABA in MNCs

(A) Sweeps at left show sPSPs recorded using sharp electrodes at various voltages from MNCs in explants prepared from EU and SL rats (see Figures S1 and S2). Plots in the middle panel show mean (± SEM) sPSP amplitude for the cells at left against baseline voltage (arrows show EREV). Bar graphs at right represent the mean (± SEM) of all collected E_GABA values from sPSP analysis in all cells tested.

(B) Sweeps at left show PSPs evoked by electrical stimulation of the DBB at various voltages in MNCs recorded from EU and SL explants. Plots in the middle show mean (± SEM) evoked PSP (ePSP) amplitudes for cells at left against baseline voltage (arrows show EREV). Bar graphs at right represent the mean (± SEM) of all E_GABA values determined from ePSP analysis (n shown on bars; ***p < 0.001).
hypothalamic explants prepared from three different strains of rats (Long Evans, L-E; Sprague-Dawley, S-D; and Fischer 344 rats; Figure 2E).

To establish if differences in $E_{\text{GABA}}$ observed in MNCs from EU and SL rats have a functional impact on GABA AR-mediated inhibitory tone, we examined the effects of antagonizing GABAARs on spontaneous AP firing rate measured using non-invasive extracellular single-unit recordings. As shown in Figures 3A and 3B, bath application of 10 $\mu$M bicuculline caused a significant excitation of MNCs in EU explants (basal 1.9 ± 0.6 Hz versus bicuculline 2.6 ± 0.7 Hz; n = 10; p = 0.0032), consistent with the existence of a significant inhibitory tone under control conditions. Conversely, bath-application of bicuculline significantly inhibited the firing rate of MNCs in SL explants (basal 2.7 ± 0.7 Hz versus bicuculline 0.9 ± 0.4; n = 17; p = 0.0243; both paired t tests). These observations indicate that the depolarizing shift in $E_{\text{GABA}}$ induced by SL is sufficient to convert the inhibitory tone that prevails under normal conditions into an excitatory tone in the SL condition.

**SL Depolarizes $E_{\text{GABA}}$ via Downregulation of KCC2**

The depolarizing shift in $E_{\text{GABA}}$ observed in SL MNCs indicates that the intracellular concentration of chloride ($[\text{Cl}^-]$) is increased under these conditions. Because $[\text{Cl}^-]$ is determined by the relative activity of the Cl$^-$/C0$_2$ exporter KCC2 and the Cl$^-$/C0$_2$ importer NKCC1 (Chamma et al., 2012), an increase in $[\text{Cl}^-]$ could be mediated by either an increase in NKCC1 activity and/or a decrease in KCC2 activity. To clarify the mechanism mediating the depolarizing shift in $E_{\text{GABA}}$ observed in SL MNCs, we examined the functional contribution of these transporters under EU and SL conditions in L-E rats.

Application of the KCC1 antagonist bumetanide (10 $\mu$M) caused a small but significant hyperpolarization of the average value of $E_{\text{GABA}}$ in MNCs from EU rats ($-4.4 ± 1.9$ mV; n = 10; p = 0.047; Figures 4A and 4B). This observation indicates that KCC1-mediated Cl$^-$/C0$_2$ import plays a role in setting the $E_{\text{GABA}}$ of MNCs in these rats. If an increase in KCC1-mediated import was responsible for the positive shift in $E_{\text{GABA}}$ caused by SL, we would expect that the hyperpolarizing effect of bumetanide on $E_{\text{GABA}}$ would be enhanced in this condition. However, as illustrated in Figures 4A and 4B, application of bumetanide only had a small and non-significant effect on $E_{\text{GABA}}$ in SL MNCs ($-2.3 ± 2.9$ mV; n = 12; p = 0.45). Therefore the effect of SL on $E_{\text{GABA}}$ is not mediated by a change in the activity of KCC1.
Application of the KCC2 antagonist furosemide (100 μM) caused a significant and reversible depolarization of \( E_{GABA} \) in MNCs of EU rats (+9.5 ± 0.8 mV; \( n = 6; p < 0.001 \); Figures 4C and 4D), indicating that chloride extrusion contributes to the maintenance of a negative \( E_{GABA} \) in MNCs from EU rats. If a decrease in KCC2-mediated Cl" export was responsible for the depolarization of \( E_{GABA} \) in SL MNCs, we would expect that the effect of furosemide would be attenuated in this condition. Indeed, furosemide had no effect on the value of \( E_{GABA} \) in MNCs from SL rats (+0.1 ± 0.8 mV; \( n = 8; p = 0.86 \); Figures 4C and 4D; paired t test was applied to all bumetanide and furosemide experiments). Taken together, these results indicate that a reduction of KCC2 activity specifically mediates the depolarizing effect of SL on \( E_{GABA} \) in MNCs.

To determine if the reduced KCC2 activity caused by SL is associated with a decrease in transporter expression, we performed western blot analysis on lysates prepared from SL rats compared to EU (p = 0.026; n = 7 for both). These observations indicate that SL causes an increase in TrkB receptor activation without affecting the total amount of TrkB expressed in the supraoptic nucleus.

Figure 3. SL Eliminates Inhibitory Tone in Rat MNCs
(A) Excerpts of single-unit activity recorded from MNCs in EU and SL rats before (basal) and during bath applications of 10 μM bicuculline.
(B) Scatter plot displays absolute firing rates of individual MNCs with and without bicuculline (connected by lines) under EU (n = 10, two rats) and SL (n = 17, three rats) conditions. Bar graph overlays represent mean values (± SEM; *p < 0.05; **p < 0.01).

To determine if TrkB activation is required to mediate the effect of SL on \( E_{GABA} \), we examined the effect of reducing TrkB activity by scavenging endogenously released TrkB agonist molecules during the SL treatment. To this end, micro-catheters coupled to osmotic minipumps delivering either TrkB receptor body (TrkB-Fc) or vehicle were stereotaxically implanted unilaterally into the supraoptic nucleus. Delivery of TrkB-Fc using this approach caused a significant reduction in the density of p-TrkB-positive cells observed in the supraoptic nucleus of SL rats compared to the contralateral side (Figure S4), confirming that this approach reduces the extent of TrkB activation caused by SL.

Application of the TrkB antagonist to SL-TrkB-Fc rats caused a small but non-significant decrease in NKCC1 expression. Therefore the depolarization of \( E_{GABA} \) caused by SL in MNCs is due specifically to a reduction in Cl" extrusion mediated by a decrease in KCC2 expression and activity.

### SL Causes Activation of TrkB Receptors in the Supraoptic Nucleus

Previous work has shown that a decrease in the expression and functional activity of KCC2 can be mediated by the activation of TrkB upon phosphorylation at tyrosine residue 515 (YS15) (Rivera et al., 2004). We therefore examined if this mechanism was involved in the effect of SL on \( E_{GABA} \) in MNCs. We first performed western blot analysis on supraoptic nucleus lysates using antibodies directed against TrkB phosphorylated at YS15 TrkB (p-TrkB) or an unrelated TrkB epitope to measure total TrkB protein (tot-TrkB). As shown in Figures 5A and 5B, the average staining intensity of the p-TrkB protein band was significantly higher in lysates obtained from SL rats than EU rats, when normalized to either a loading control (p-TrkB/GADPH; p = 0.004) or tot-TrkB (p = 0.026; n = 7 for both). These observations indicate that SL causes an increase in TrkB receptor activation without affecting the total amount of TrkB expressed in the supraoptic nucleus.

To determine if TrkB activation is required to mediate the reduction in Cl" gradient induced by SL, we examined the effect of locally scavenging endogenously released TrkB agonist molecules during the SL treatment. As shown in Figure 5C, recordings from MNCs in explants prepared from SL rats in which the supraoptic nucleus was infused with vehicle showed an average value of \( E_{GABA} \) (−38.8 ± 4.06 mV; \( n = 13 \)) that was equivalent to that in non-cannulated SL rats (−33.9 ± 2.2 mV; \( n = 52; p = 0.246 \)) and significantly more depolarized than control (EU) rats (−57.4 ± 1.4 mV; \( n = 43; p < 0.001 \)). However, the average value of \( E_{GABA} \) measured in MNCs recorded from SL rats receiving TrkB-Fc into the supraoptic nucleus (SL-TrkB-Fc rats; −53.2 ± 3.0 mV; \( n = 21 \)) was significantly more hyperpolarized than vehicle-treated animals (p = 0.003) and equivalent to that observed in EU rats (p = 0.251; all comparisons made using one-way ANOVA followed by Student-Newman-Keuls post-hoc test). Furthermore, as illustrated in Figure 5D, DBB stimulation failed to significantly excite MNCs from SL-TrkB-Fc rats (evoked/baseline AP frequency = 1.63 ± 0.48; \( n = 6; p = 0.394 \)), whereas the same stimulation induced a robust excitation of MNCs from SL controls (3.37 ± 0.78; \( n = 16; p < 0.001 \); both Mann-Whitney rank-sum tests; Figure S5). These observations indicate that TrkB activation is required to mediate the depolarization of \( E_{GABA} \) induced by SL in MNCs.
BDNF Is Required for SL-Mediated Cl\(^{-}\) Gradient Collapse in MNCs

The activation of TrkB receptors can be mediated by ligands including brain-derived neurotrophic factor (BDNF), neurotrophin 4/5 (NT 4/5), and NT 3 (Boulle et al., 2012). However, previous work has shown that BDNF transcription is increased during physiological and pathological states under which Cl\(^{-}\)/C\(_0\) gradient collapse is evident in neurons (Gall, 1993; Rivera et al., 2002), and that the transcription and secretion of BDNF by VP MNCs increases during hyperosmotic stress (Arancibia et al., 2007). To determine if BDNF is required for SL-mediated collapse of the Cl\(^{-}\)/C\(_0\) gradient in MNCs, we examined the effect of knocking down the level of BDNF in vivo using a short-hairpin RNA (shRNA) that selectively inhibits BDNF synthesis (BDNF-shRNA; Figure S6).

Adeno-associated viruses (AAV) driving production of BDNF-shRNA or a scrambled sequence (scr-shRNA; used as control) were first administered by intracerebroventricular (i.c.v.) injection in anesthetized rats, and the animals were allowed to recover for 4–6 weeks prior to tissue collection for western blot analysis. As illustrated in Figures 6A and 6B, the average expression of BDNF in the anterior hypothalamus of BDNF-shRNA-treated rats was significantly lower than controls (−33.5 ± 19.3% relative to controls; n = four rats in each group; p = 0.044). We next examined the effects of delivering these AAVs by direct stereotaxic injection into the supraoptic nucleus. This procedure significantly reduced the average staining intensity of immunohistochemically detected BDNF in the supraoptic nucleus of rats having received BDNF-shRNA (60.1% ± 13.7%; n = 3 rats; p = 0.037) compared to scr-shRNA (100.0% ± 7.1%; n = four rats; Figure S7).

To determine if the TrkB-dependent collapse of Cl\(^{-}\) gradient and inhibitory signaling induced by SL is mediated by BDNF, we examined the effects of DBB stimulation on MNCs in hypothalamic explants from SL rats having normal and reduced expression of BDNF within the supraoptic nucleus. As found in otherwise intact SL rats, MNCs recorded from the supraoptic nucleus of SL rats having received scr-shRNA were commonly depolarized and excited by DBB-mediated activation of GABA\(_{\text{A}}\)Rs (Figure 6C) at voltages near threshold. In contrast, MNCs recorded from SL rats having received BDNF-shRNA displayed hyperpolarizing and inhibitory responses to DBB stimulation (Figure 6C). Moreover, the value of \(E_{\text{GABA}}\) measured in MNCs from SL rats subjected to BDNF knockdown (−56.2 ± 5.4 mV; n = 10) was significantly more hyperpolarized than in scr-shRNA animals (−38.3 ± 2.6 mV; n = 9; p = 0.005; one-way ANOVA and Student-Newman-Keuls post-hoc test; Figure 6D) and was equivalent to EU controls (p = 0.806; Figure 6D). These results indicate that BDNF is the endogenous TrkB agonist responsible for the collapse in Cl\(^{-}\) gradient observed in SL rats.

Activated Microglia Are Not Required for Cl\(^{-}\) Gradient Collapse

Previous studies have shown that BDNF can be released either by activated microglia (Coull et al., 2005) or by neuronal somata and dendrites (Kolarow et al., 2007; Kuczewski et al., 2009). A previous study has reported that microglia in the supraoptic...
nucleus become activated during SL (Ayoub and Salm, 2003). Therefore we investigated the possibility that these cells might provide the BDNF underlying the SL-mediated Cl⁻ gradient collapse. We first examined the morphology of microglia in the supraoptic nucleus using immunohistochemical detection of the microglial marker ionized calcium-binding adaptor molecule 1 (Iba1). In agreement with previous work, we found that microglia in the supraoptic nucleus of EU rats resemble those in resting states and that SL treatment leads to an activation of these cells displayed as a significant hypertrophy of their somata and processes (Figure 7A). Indeed, images of the supraoptic nucleus from SL rats showed a significantly greater Iba1-positive surface area than those from EU rats (p = 0.002 with t test; n = 12 sections from three rats in each group; Figure 7B).

In order to examine whether the activation of microglia is linked with the SL-induced Cl⁻ gradient collapse, we tested the effect of inhibiting microglial activation on the \( E_{\text{GABA}} \) of MNCs in the supraoptic nucleus. While undergoing the SL treatment, one group of rats concurrently received daily intraperitoneal (i.p.) injections of minocycline hydrochloride (50 mg/kg), a compound that crosses the blood-brain barrier and reduces microglial activation in the brain (Fan et al., 2007). Another group of rats receiving identical SL treatment received injections of equal amounts of saline as controls. As illustrated in Figure 7C, the mean value of \( E_{\text{GABA}} \) measured in MNCs from minocycline-treated SL rats (~37.3 ± 4.3 mV; n = 9) remained significantly more depolarized than EU rats (p < 0.001) and was equivalent to saline-injected SL rats (~30.4 ± 5.0 mV; n = 7; p = 0.560) or uninjected SL controls (p = 0.476; all one-way ANOVA followed by Student-Newman-Keuls post-hoc test). These results indicate that microglial activation is not required for SL-mediated collapse of the Cl⁻ gradient in MNCs.

**SL Impairs BR-Mediated Inhibition of MNCs**

To extend the functional significance of the findings reported above, we next determined if the loss of GABA\(_A\)-R-mediated inhibition caused by SL observed during in vitro recordings was sufficient to impair the BR-mediated inhibition of VP MNCs in vivo. Since it is well established that intravenous (i.v.) injection of \( \alpha \)-adrenoreceptor agonists such as phenylephrine (PE) inhibits firing in VP MNCs via BR activation (Cunningham et al., 2004; Renaud and Bourque, 1991; Renaud et al., 1988), we examined whether this response is diminished in SL rats. In single-unit extracellular recordings from VP MNCs in urethane-anaesthetized rats, i.v. infusion of PE (2.5 \( \mu \)g/kg) raised the mean arterial pressure by an equivalent amount in EU (+49.8 ± 5.3 mmHg, n = 14) and SL animals (+53.9 ± 5.9 mmHg, n = 16; p = 0.61). However, the significant inhibition of firing observed in neurons recorded in EU rats (~40.0 ± 10.1%; n = 10; p = 0.004; Wilcoxon signed-rank test) was eliminated in SL rats (~4.5 ± 26.6%; n = 17; p = 0.0784; paired t test; Figures 8A and 8B). Moreover, unlike in EU rats where VP MNCs were either inhibited (9/10 cells) or unaffected (1/10 cells), a significant proportion of VP MNCs in SL rats were excited by BR activation (6/17 cells; p = 0.033; \( \chi^2 \) test).

**Peripheral VP Receptors Contribute to Elevated BP in SL Rats**

The results described above indicate that BR inhibition is abolished in VP MNCs after SL. To determine if SL also causes an increase in the basal electrical activity of VP MNCs in vivo, we compared the spontaneous AP firing rate of these cells in EU and SL rats. Average firing rate of VP MNCs in SL rats was significantly higher (9.4 ± 1.2 Hz, n = 23) than that in EU rats (5.5 ± 0.8 Hz, n = 13; p < 0.05; data not shown), consistent with the increased VP secretion observed after SL (Ludwig et al., 1996).
To determine if an increase in firing and secretion by VP MNCs plays a role in the regulation of BP during SL, mean arterial pressure (MAP) was monitored by radio telemetry in freely moving rats. As illustrated in Figure 8C, MAP rose steadily during the course of the 7 day SL treatment (basal BP: 97.7 ± 1.9 mmHg; day 7: 113.5 ± 2.1 mmHg; n = 10; p < 0.001), whereas MAP remained stable in EU rats over the same time period (basal BP: 98.3 ± 2.5 mmHg; day 7: 100.6 ± 2.2 mmHg; n = 6; p = 0.473; one-way repeated-measures ANOVA with Student-Newman-Keuls post-hoc test). To determine if peripheral VP receptors contribute to the increase in BP induced by SL, we examined the effect of SL on MAP in rats receiving a continuous systemic infusion of dGly[Phaa1,d-tyr(et), Lys, Arg]VP, a VP receptor type 1 (V1R) antagonist. As shown in Figure 8C, the SL-mediated increase in MAP was significantly lower in rats treated with the V1R antagonist (Δ day 7: +8.8 ± 2.5 mmHg; n = 6) when compared to SL controls (+15.1 ± 1.0 mmHg; n = 10; p = 0.03).

**DISCUSSION**

GABA Hyperpolarizes and Inhibits MNCs in EU Rats

Previous extracellular recordings in vivo have shown that local application of GABA inhibits the electrical activity of MNCs in the rat supraoptic nucleus (Arnauld et al., 1983) and that inhibition of VP MNCs due to BR activation requires local GABA_ARs (Jhamandas and Renaud, 1986). In agreement with these observations, a previous study reported that the value of E_{GABA} measured by perforated patch recording in hypothalamic slices under control conditions (Haam et al., 2012). Because this is a critical issue for our understanding of the network basis for regulation of MNCs, we re-examined this issue using a variety of approaches.
As shown in Figure 2, gramicidin-based perforated patch recordings from identified VP MNCs in the supraoptic nucleus of hypothalamic slices showed that EGABA is consistently and significantly more hyperpolarized than rheobase ($E_{\text{GABA}}$) in EU rats. Identical results were obtained during intracellular recordings from the VP-rich (ventral-caudal) zone of the supraoptic nucleus in hypothalamic explants prepared from three different strains of rats (Figure 2E). Our findings are therefore in agreement with previous studies indicating that MNCs in EU rats maintain a hyperpolarized EGABA that allows these cells to be potently inhibited by GABAergic synaptic inputs, including those activated by BR activation. Although we have no explanation for the discordant results reported by Haam and colleagues, the present work and studies by others have shown that the neuronal Cl$^-$ gradient is a labile parameter that can be rapidly collapsed as a result of stress (Hewitt et al., 2009) or experimental procedures (Dzhala et al., 2012).

Reduced KCC2 Expression Depolarizes EGABA during SL

A previous study has reported that the depolarizing shift in $E_{\text{GABA}}$ observed in MNCs from SL S-D rats is caused by an increase in the activity and expression of the Cl$^-$ importer NKCC1 (Kim et al., 2011). Although our experiments and those of Kim and colleagues revealed an equivalent SL-induced shift in $E_{\text{GABA}}$, our electrophysiological analysis with blockers of KCC2 and NKCC1 indicated that this effect is due to an SL-mediated reduction in KCC2 activity, rather than an increase in NKCC1 activity. Our findings were corroborated by western blot analysis showing that KCC2 expression is significantly reduced by SL in L-E rats, whereas NKCC1 levels are not significantly increased (Figure 4E). The absence of increased NKCC1-mediated Cl$^-$ import in SL rats may seem surprising since [Cl$^-$/C0]i levels are increased in this condition. However it remains possible that a small residual NKCC1-mediated import could be sufficient to raise [Cl$^-$/C0]i under conditions where KCC2 activity is depressed. Alternately, the increased value of [Cl$^-$/C0]i that prevails in MNCs from SL rats could simply result from the cumulative effect of Cl$^-$ influx mediated by basal synaptic and extrasynaptic GABA$_{AR}$ activity and weakened Cl$^-$ extrusion due to downregulated KCC2. The basis for differences between our results and those of Kim (Kim et al., 2011) are unknown, but two possible explanations should be considered.

First, previous work has shown that the pattern of gene expression is different in the supraoptic nucleus of different rat strains (Hindmarch et al., 2007). Although SL has been found...
to cause a depolarization of $E_{\text{GABA}}$ in all strains examined so far (Fischer 344, L-E, S-D, and Wistar), it is possible that Cl\(^{-}\) transport is mediated more strongly by NKCC1 in S-D rats and more strongly by KCC2 in L-E rats. Second, the tissue preparation methods used in the two studies were slightly different. Notably, whereas our samples were collected by rapid microdissection of the supraoptic nucleus immediately following removal of the brain, samples obtained by Kim and colleagues were collected by excision of the supraoptic nucleus from hypothalamic slices. Given the labile nature of KCC2 expression in slices (Dzhala et al., 2012), it is possible that this procedure can mask the true contribution of KCC2 to Cl\(^{-}\) homeostasis in the supraoptic nucleus. Given the ubiquitous involvement of changes in KCC2 expression as a mediator of altered Cl\(^{-}\) in various pathophysiological states, and the lesser degree of trauma applied to the tissue used in our western blot and electrophysiological analysis (i.e., acute explants), we believe that KCC2 serves as the predominant regulator of [Cl\(^{-}\)] in the supraoptic nucleus and that its modulation is a key contributor to the depolarizing shift in $E_{\text{GABA}}$ induced by SL in vivo.

**BDNF-Activated TrkB Depolarizes $E_{\text{GABA}}$ during SL**

Our results show that SL causes the functional activation of TrkB receptors in MNCs of the supraoptic nucleus and that activation of these receptors is required to mediate the depolarization of $E_{\text{GABA}}$ in response to this stimulus. Previous studies have shown that KCC2 activity can be regulated by several types of TrkB-dependent mechanisms. For example, TrkB activation has been shown to suppress KCC2 transcription through a cAMP response element-binding protein (CREB)-dependent pathway (Rivera et al., 2004). Our observation that KCC2 protein expression is significantly reduced in SL rats is consistent with this mechanism and suggests that a decrease in KCC2 synthesis may contribute significantly to the decrease in KCC2 activity associated with this condition. However, the involvement of post-translational mechanisms cannot be excluded. For example, TrkB activation has been shown to activate calpain (Zadran et al., 2010), which can reduce KCC2 activity by proteolytic cleavage (Puskarjov et al., 2012). Moreover, TrkB activation can increase NMDA receptor activity by phosphorylation (Carreno et al., 2011), and Ca\(^{2+}\) influx through NMDA receptors has been shown to reduce KCC2 activity via endocytosis (Lee et al., 2011). The specific mechanism(s) by which TrkB activation downregulates KCC2 in SL MNCs remain to be determined.

Our shRNA-mediated knockdown experiments revealed that BDNF is the agonist responsible for the activation of TrkB in response to SL in MNCs. BDNF has also been identified as a mediator of altered neuronal Cl\(^{-}\) homeostasis in neuropathic pain (Coull et al., 2005) and morphine hyperalgesia (Ferrini et al., 2013). In the latter models, experiments revealed that activated microglia secrete the BDNF required for TrkB activation and downregulation of KCC2 (Ferrini and De Koninck, 2013). Although SL causes microglial activation in the rat supraoptic nucleus (Figure 7) (Ayoub and Salm, 2003), the effect of SL on $E_{\text{GABA}}$ in MNCs was not prevented by inhibiting microglial activation in vivo. Therefore, an alternate possibility is that BDNF is released by the MNCs themselves. Indeed, BDNF is known to be released by many types of neurons in an activity-dependent manner (Kuczewski et al., 2009), and previous work has shown that BDNF is synthesized by MNCs (Carreno et al., 2011). Moreover, acute hyperosmotic conditions, which excite MNCs (Bourque, 2008), have been shown to enhance the transcription of BDNF by MNC somata and to stimulate BDNF release by the dendrites of these neurons in the supraoptic nucleus in vivo (Aliaga et al., 2002; Arancibia et al., 2007). Therefore SL is likely to trigger the activation of TrkB receptors as an autocrine or paracrine response to BDNF release by the MNCs themselves.

**High Salt Intake Promotes a VP-Dependent Increase in BP**

Increases in plasma sodium associated with a high level of dietary salt intake are linked to elevated BP in salt-sensitive hypertensive patients (He et al., 2013; He and Macgregor, 2012; Schmidlin et al., 2007), but the mechanisms underlying this effect are unclear. Hyperosmolality is a necessary consequence of hypernatremia and thus activates central osmoreceptors (Bourque, 2008). Moreover, the recruitment of central osmoreceptor pathways has been shown to enhance sympathetic tone through an excitation of preautonomic neurons (Toney and Stocker, 2010) and VP release through glutamatergic excitation of MNCs (Trudel and Bourque, 2010). Although both factors could potentially contribute to increases in BP, the activation of BR should normally counteract increases in BP by opposing both mechanisms. However, in vitro and in vivo results strongly indicate that collapse of the Cl\(^{-}\) gradient eliminates BR inhibition of VP-MNCs during SL, suggesting that this negative feedback mechanism is impaired under these conditions. Moreover, in a significant proportion of MNCs, the depolarizing response to $\text{GABA}_A$ receptor activation appeared to be sufficient to produce an excitatory response to BR activation. By promoting a further excitation of MNCs, this functional switch in the polarity of BR input would eventually result in enhanced VP secretion and thus amplify the contribution of this hormone to the elevation in BP as SL progresses. Indeed, data shown in Figure 8C indicate that circulating VP mediates a significant proportion of the SL-mediated increase in BP and that V1R-mediated contribution to BP increased as a function of time during the SL protocol.

There are three known mechanisms through which circulating VP can modulate BP. First, increases in systemic VP have been shown to enhance BR reflex-mediated decreases in sympathetic output though a V1R-dependent action in the area postrema (Hasser et al., 1997). Second, VP has been shown to activate V1R expressed at the subfornical organ to reduce BP, possibly via a reduction of sympathetic output (Smith and Ferguson, 1997). However, in both cases V1R activation promotes a lowering of BP. It is therefore unlikely that these mechanisms contribute to the elevated BP found under SL. Third, V1Rs are widely expressed in vascular smooth muscle where they mediate a potent vasoconstrictor effect (Henderson and Byron, 2007). Moreover, VP levels reportedly associated with SL (Ludwig et al., 1996) appear to be sufficient to induce vasoconstriction (Henderson and Byron, 2007), suggesting that increases in circulating VP associated with SL could contribute to the elevation of BP observed under such conditions. This hypothesis is directly supported by our finding that antagonism of V1R partially reversed the SL-mediated elevations in BP. While the nature of
the residual component in SL-induced BP increase after V1 blockade remains undetermined, we suspect a possible role of hyperosmolality-induced increase in sympathetic outflow (Toney and Stocker, 2010).

Concluding Remarks
Our results show that high salt intake can cause a collapse of the Cl− gradient across VP neurons and abolish BR-mediated negative feedback inhibition of these cells due to the activity-dependent release of BDNF and downregulation of KC22 that results from the autocrine activation of TrkB receptors. The elevated firing rate of MNCs associated with this condition leads to an increase in VP secretion and a significant V1R-dependent increase in BP. These findings show that state-dependent changes in neurotrophin signaling can mediate pathological consequences by promoting plastic changes in hypothalamic homeostatic networks.

EXPERIMENTAL PROCEDURES

Drugs and Antibodies
Kynurenic acid was purchased from Sigma. DNXQ (Tocris) was kept at a stock concentration of 20 mM in dimethyl sulfoxide (DMSO) (Sigma). Bicuculline methochloride, furosemide, bumetanide, and gabazine were obtained from Tocris. V1R antagonist dGlu[Phaa1,d-tyr(et), Lys, Arg]VP was purchased from Bachem. TrkB-Fc chimera was purchased from R&D Systems. Antibodies from commercial sources: KC22 (1:500), pan-TrkB (1:1,000), and GAPDH (1:5,000) from Millipore; BDNF (Santa Cruz Biotechnology; 1:100 for WB, 1:300 for IHC); Y515 phosphorylated TrkB from Abcam (1:100); and Iba1 from Wako chemicals (1:1,000). NKCC1 antibody (1:500) was generously provided by Dr. R. James Turner (NIH). VP-neurophysin antibody (mouse monoclonal, 1:100) was kindly provided by Dr. Hal Gainer (NIH).

Animals
Adult male rats were maintained on a 12:12 hr light cycle and provided with ad libitum access to food and water except where indicated in specific protocols. All procedures involving animals were conducted according to protocols approved by the Facility Animal Care Committee of McGill University, University of Otago Animal Ethics Committee, and the Institutional Animal Care and Use Committee of the UNT Health Science Center.

In Vitro Electrophysiological Recordings
Acute rat hypothalamic explants prepared as previously (Ghamari-Langroudi and Bourque, 2001) were superfused (~1–1.5 ml/min) with warm (31°C–33°C) oxygenated (95% O2, 5% CO2, pH 7.35) artificial cerebrospinal fluid (ACSF) comprising NaCl (104 mM), NaHCO3 (26 mM), NaHPO4 (1.23 mM), KCl (2 mM), MgCl2 (1 mM), CaCl2 (2 mM), and D-glucose (10 mM), and mannitol was added to the desired osmolality. For extracellular recordings, KCl was increased to 4 mM and CaCl was reduced to 1 mM. Recordings were made using micropipettes (extracellular, 15–20 MΩ; intracellular, 80–110 MΩ) filled with 2 M K-acetate.

Acute hypothalamic slices 400 μm thick were cut at an angle of 38° relative to the surface of the cortex as described previously (Trudel and Bourque, 2010) and submerged in warm oxygenated ACSF (same as above). Granimicin (Sigma) was dissolved in DMSO at a stock concentration of 0.05 mg/ml and diluted 1:500 into K-glutamate-based internal pipette solution. Borosilicate glass patch pipettes (3–6 MΩ) were pulled and filled with the granicin-containing internal solution. Recordings from identified VP neurons were made by targeting eGFP-expressing neurons in slices prepared from transgenic VP-eGFP Wistar rats. After establishing gigahm seals with target cells, measurements were made after the pipette resistance dropped to values between 40 and 80 MΩ. For both preparations, electrical stimulation was performed using a DS2 Digitimer coupled to a concentric bipolar electrode (FHC, Inc.) placed within the DBB.

In Vivo Electrophysiological Recordings
The pituitary stalk and right SON of urethane-anaesthetized rats (1.25 g/kg, Sigma) were exposed by a ventral approach through the oral cavity. Extracellular single unit recordings were made using micropipettes (15–40 MΩ) filled with 0.9% saline. MNCs were identified by antidromic spikes elicited from pituitary stalk stimulation with a bipolar electrode (Science Products GmBH). VP-MNCs were characterized by their spontaneous phasic activity or by a lack of excitation following intravenous (i.v.) cholecystokinin (CCK) injection (20 μg/kg, 0.5 ml/kg in 0.9% saline), or as oxytocin-MNCs by transient excitation following CCK injection (Sabatier et al., 2004). At the end of the experiment, the rats were killed by anesthetic overdose (60 mg/kg pentobarbital or 1 g/kg urethane, i.v.).

SON Tissue Lysate Preparation
Brains of age-matched EU and 7 day SL rats were rapidly removed and submerged into oxygenated isosmotic ACSF, and then SON tissue blocks (1 mm3) or the entire anterior hypothalami were excised using a pair of angled spring scissors. Tissue lysates were then prepared by trituration in HEPES-based buffer followed by centrifugation. Western blotting experiments were performed using standard procedures and intensities of bands were quantified using ImageJ (NIH).

Immunohistochemistry
Brains of age-matched EU and 7 day SL Long Evans rats perfused with 4% paraformaldehyde (Sigma) were sectioned in the coronal plane (50 μm thick) using a vibratome and stained immunohistochemically using various primary antibodies listed above. Once prepared into slides, images of immunostained structures were captured as continuous stacks of confocal images (1 μm thick) using an Olympus FV1000 scanning laser confocal microscope equipped with a krypton/argon mixed gas laser. All of the analysis was performed with ImageJ (NIH).

Drug Infusion of Unilateral SON
Rats were instrumented with unilateral cannula targeted at the SON as previously described (Carreño et al., 2011). Under isoflurane anesthesia (2%), the rats were placed in a stereotaxic frame (Kopf Instruments), and a 28 gauge cannula (Plastics One) was placed in the right SON. Each cannula was connected to an osmotic pump implanted subcutaneously around the neck region of rats (ALZET) filled with either saline or TrkB-Fc. After 7–10 days of recovery, rats underwent a 7 day SL protocol and then were prepared into hypothalamic explant preparations for in vitro electrophysiological experiments. The efficiency of TrkB-Fc infusion was tested by post-hoc immunohistochemistry.

AAV-Mediated Knockdown of BDNF in the Supraoptic Nucleus
Adeno-associated virus (AAV) serotype 2 conjugated with shRNA directed against BDNF was custom-generated (Vector Biolabs). To test the effectiveness of the shRNA’s ability to induce a specific knockdown of BDNF in vitro, HEK293T cells were transfected with 2 μg BDNF with or without BDNF-shRNA. Twenty-four hours later, the conditioned media was collected and concentrated and then run in a western blot. AAV conjugated with either the BDNF- or scrambled (scr)-shRNA (Vector Biolabs) were injected intracerebroventricularly (i.c.v.) into the third ventricle (2 μl) to test the in vivo effectiveness of knockdown in hypothalamic tissue. After 4 weeks the rat brains were harvested and the anterior hypothalami were microdissected and processed for western blotting. To induce a specific knockdown of BDNF in the supraoptic nucleus, rats were stereotaxically injected with AAVs conjugated with either BDNF- or scr-shRNA (1 μl from Bregma, X: 1.5 mm, Y: −0.5 mm, Z: 7.7 mm). The virus was allowed to express in vivo for 5–6 weeks, after which the rats underwent a 7 day SL treatment. Their brains were harvested and prepared for hypothalamic slices as described above. After patch-clamp recordings, the slices were immersion-fixed in 4% paraformaldehyde overnight and processed for post-hoc immunohistochemistry to confirm the success of knockdown.

Inhibition of Microglial Activation via Intraperitoneal Injections of Minocycline
Minocycline hydrochloride (Sigma) was dissolved in 0.9% saline, and appropriate amounts of NaOH were added to adjust the pH to ~7.4. Each rat
received a daily dose of 50 mg/kg i.p., for 7 days, during which they also underwent SL treatment. A control group of rats received injections of equal amounts of saline. On the last day of SL, the rat brains were harvested and made into hypothalamic slice preparations, and patch-clamp recordings were performed.

**Radio Telemetry Measurement of BP and In Vivo Infusion of VP Antagonist**

As previously described (Cunningham et al., 2012), rats were implanted with an abdominal aortic catheter attached to a TA11PA-C40 radio telemetry transmitter under isoflurane anesthesia (2%). The transmitter was secured to the abdominal muscle and remained in the abdominal cavity for the duration of the experiment. After a 2 week recovery period, MAP signals from the telemetry device were recorded during a 7 day SL protocol using a Dataquest IV radio telemetry system (Data Sciences Inc.). Prior to SL treatment, a subgroup of these rats was also implanted with an osmotic minipump (ALZET) in a subcutaneous pouch made between the scapulae under isoflurane anesthesia (2%) for subcutaneous drug infusion.

**Statistics**

All values in this study are reported as mean ± the standard error of the mean (SEM). Unless explicitly stated, statistical differences between mean values were tested using Student’s two-tailed t test. All statistical tests were performed with Sigmaplot 12.3 (SPSS Inc.) or Prism 5 (Graphpad Software Inc.). Data points above or below twice the standard deviation from the mean were excluded from statistical comparisons. Differences between values were considered to be significant when p < 0.05.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures and seven figures and can be found with this article online at http://dx.doi.org/10.1016/j.neuron.2014.12.048.

**AUTHOR CONTRIBUTIONS**


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**REFERENCES**


Supplemental Information

High Salt Intake Increases Blood Pressure via BDNF-Mediated Downregulation of KCC2 and Impaired Baroreflex Inhibition of Vasopressin Neurons
Katrina Y. Choe, Su Y. Han, Perrine Gaub, Brent Shell, Daniel L. Voisin, Blayne A. Knapp, Philip A. Barker, Colin H. Brown, J. Thomas Cunningham, and Charles W. Bourque
**Figure S1.** 7 day SL significantly increases plasma osmolality in rats. Bar graphs show the significant increase in the mean (± s.e.m) plasma osmolality of rats that drank 2% NaCl solution as their only source of water for 7 days (n shown on bars, p < 0.001, Mann-Whitney rank sum test).
Figure S2. DBB stimulations produce PSPs in MNCs which is fully mediated by GABA$_A$Rs. Electrophysiological trace indicates the elimination of sPSP (A) and ePSP (B) in MNCs evoked by DBB stimulation under 20 μM DNQX (glutamate receptor blocker; black trace), which is completely abolished by the addition of 1 μM gabazine (GBZ, GABA$_A$R antagonist; red trace).
Figure S3. SL induces decreased KCC2 expression in VP MNCs. The robust expression of KCC2 (red) in VP-expressing MNCs (green) in EU rats is observed in confocal micrographs of immunohistochemically stained brain sections containing the supraoptic nucleus. After SL, KCC2 expression is markedly decreased in these neurons. Scale bar represents 20 μm.
Figure S4. TrkB-Fc infusion during SL significantly reduces the number of MNCs expressing p-TrkB. Bar graphs compare the average (± s.e.m) number of p-TrkB positive neurons in SL rats between right supraoptic nucleus which received TrkB-Fc infusion (SL+TrkB-Fc) and the contralateral side which was uninfused (n shown on bars, p<0.05; Student’s t-test).
Figure S5. TrkB-Fc infusion during SL eliminates DBB-induced excitation. Scatter plots indicate the ratio between DBB-stimulation evoked and baseline AP frequency for each cell and bar graph overlays indicate the mean (± s.e.m). MNCs in SL rats are significantly excited by DBB stimulation ($p<0.001$ marked by *, Mann-Whitney rank sum test) whereas TrkB-Fc infused SL rats (SL+TrkB-Fc) were not affected by the stimulation ($p=0.394$, n.s. represents not significant).
Figure S6. BDNF-shRNA abolishes release BDNF release in HEK293T cells. A representative Western blot shows that extracellular BDNF is detected in the media around HEK293T cells transfected with plasmids containing the BDNF gene, but co-transfection with plasmids containing BDNF-shRNA abolishes BDNF release into the cell media.
**Figure S7.** Local infusion of AAV-BDNF-shRNA reduces BDNF-immunopositive signal in the supraoptic nucleus. Confocal images show post-hoc immunohistochemical staining against BDNF in the supraoptic nucleus of hypothalamic slices used in electrophysiological experiments. Note the downregulation of BDNF protein in the AAV-BDNF-shRNA injected supraoptic nucleus compared to the AAV-scr-shRNA injected supraoptic nucleus (area ato the left of the optic tract (OT), where recordings from MNCs were obtained). Scale bar represents 100 μm.
SUPPLEMENTAL EXPERIMENTAL PROCEDURES

Animals

Adult male Sprague-Dawley, Fischer 344, Long-Evans rats (all from Charles River Laboratories, Wilmington, MA, 100-250 g), VP-eGFP Wistar transgenic (Ueta laboratory, Kitakyushu, Japan) were maintained on a 12:12 h light cycle and provided with ad libitum access to food and water except where indicated in specific protocols. SL was achieved by providing adult male rats with 2% NaCl solution as drinking fluid instead of tap water for 7 days. Rat plasma samples of EU or SL rats were prepared by collecting the trunk blood after decapitation and isolating the plasma using a centrifuge. Osmolality measurements were made using a freezing-point microosmometer (Advanced Instruments). All rats were acclimated to laboratory conditions for at least 1 week prior to being used in experimental protocols. All procedures involving animals were conducted according to protocols approved by the Faculty Animal Care Committee of McGill University following the standards of Canadian Council on Animal Care (Bourque Laboratory); the University of Otago Animal Ethics Committee in accordance with the recommendations of the Australian and New Zealand Council for the Care of Animals in Research and Teaching (Brown laboratory, University of Otago), and the Institutional Animal Care and Use Committee of the University of North Texas Health Science Center, following the NIH Guide for the Care and Use of Laboratory Animals (Cunningham laboratory, UNT Health Science Center).

Preparation of in vitro hypothalamic explants and solutions

Acute hypothalamic explants were prepared from adult male Long Evans, Fischer 344, and Sprague-Dawley rats (80-160 g) as described previously (Ghamari-Langroudi and Bourque, 2001) and superfused (~1-1.5 ml/min) with warm (31-33 °C) oxygenated (95% O2; 5% CO2, pH 7.35) ACSF comprising (in mM): NaCl (104), NaHCO3 (26), NaH2PO4 (1.23), KCl (3), MgCl2 (1), CaCl2 (2), D-glucose (10) and mannitol added to the desired osmolality. For extracellular recordings, KCl was increased to 4 mM and CaCl2 was reduced to 1 mM to promote spontaneous electrical activity. GABA_A activity in MNCs were successfully isolated from other types of synaptic activity by adding appropriate amounts of 200 mM stock solution of DNXQ, a blocker of AMPARs and kainate receptors (Tocris Bioscience, Bristol, United Kingdom) dissolved in dimethyl sulfoxide (DMSO) (Sigma Aldrich Co., St-Louis, MO; kept at 4 °C), to ACSF at a final concentration of 20 μM. All remaining synaptic activity was abolished by adding GABA_A blockers gabazine (1 μM) or bicuculline (10 μM; both from Tocris) to the bath.
**Electrophysiological recordings: intracellular**

Intracellular recordings (low pass filtered at 3 kHz) were made via high resistance pipettes (80-110 MΩ) pulled with P-97 micropipette puller (Sutter Instrument, Novato, CA) using 1.2 x 0.68 mm (outer diameter x inner diameter) filaments glass capillary, filled with 2M K Acetate (100-130 MΩ). Electrical stimulation of the DBB was performed using a DS2 Digitimer coupled to a concentric bipolar electrode (FHC, Bowdoinham, MC) placed within the DBB. The rat supracoaptic nucleus is comprised of two distinct populations of MNCs: oxytocin-releasing (~35% of the cells) and VP-releasing neurons (~65% of the cells), which are each grouped at opposite poles of the nucleus (Rhodes et al., 1981). While cells were not specifically identified as VP MNCs in our experiments performed with the hypothalamic explant preparations, the electrodes were guided to the caudal and ventral (VP-enriched) part of the nucleus thus most recordings were likely from VP secreting neurons. For each neuron tested, a linear fit between the amplitude of DBB-evoked or spontaneous PSPs and the baseline membrane voltage was obtained, and reversal potential was determined to be the membrane voltage where the PSP amplitude becomes 0 mV. For the post-stimulus analysis of excitability, baseline activity was defined to be the average AP frequency during the 100 ms time period preceding the electrical stimulus and DBB-evoked changes in activity was measured as the average AP frequency measured during the 40 ms period that followed stimulation. DBB-induced changes in probability of firing were calculated evoked/basal frequency. All of the above analyses were performed using pClamp 10.0 and Sigmaplot 10.0 software (Molecular Devices Corp., Sunnyvale CA).

**Electrophysiological recordings: extracellular**

Extracellular recordings of AP firing were made using micropipettes with resistances at 15-20 MΩ using the same glass and puller as intracellular recording electrodes again filled with 2 M K Acetate. The voltage signal was amplified 100x and band-pass filtered between 600 and 1500 Hz. Bicuculline-induced changes in steady-state firing measured by extracellular recording were quantified as the difference in mean firing rate observed during the last 30s of a 3-5 min application of bicuculline (Tocris; 10 μM) compared to the average rate of firing observed during the 3 minutes preceding drug application. Neurons with unstable baseline activity were excluded from the analysis. All of the above analyses were performed using pClamp 10.0 and Sigmaplot 10.0 software.

**Preparation of angled hypothalamic slices and Gramicidin perforated patch clamp recordings**

Hypothalamic slices 400 μm thick were cut at an angle of 38° relative to the surface of the cortex as described previously (Trudel and Bourque, 2010) and incubated in warm (31-33
oxygenated (95% O₂; 5% CO₂, pH 7.35) ACSF comprising (in mM): NaCl (104), NaHCO₃ (26), NaH₂PO₄ (1.23), KCl (3), MgCl₂ (1), CaCl₂ (2), D-glucose (10) for at least 1 hour before recordings began. Prior to recording, 20 μM DNQX (Tocris) was added to the ACSF in order to isolate GABAₐ receptor activity. Some recordings were obtained in the presence of 3 mM kynurenic instead of DNQX, with no visible difference in the results. Gramicidin (Sigma) was dissolved in DMSO at a stock concentration of 0.05mg/μL and diluted 1:500 into internal pipette solution (in mM): K-gluconate (140), MgCl₂ (2), HEPES (10), ATP (2) and GTP (0.4), pH adjusted to 7.25 with NaOH). Borosilicate glass patch pipettes (3-6 MΩ) were pulled using P-97 micropipette puller and filled with normal internal solution at the tip, and then with the gramicidin containing internal solution in order to allow initial seal formation. Recordings from identified VP neurons were made by targeting eGFP-expressing neurons in slices prepared from transgenic VP-eGFP Wistar rats. Gigaohm-seals were quickly established with target cells, then E_GABA measurements were made after the pipette resistance dropped to values between 40-80 MΩ, which typically took 40-60 minutes. Once these resistance values were reached, pipette resistances remained stable throughout the recordings. Cells were voltage-clamped at -70 mV, and 10 mV voltage steps were applied ranging from -110 to -20 mV during which electrical stimulations of the DBB were performed with a bipolar electrode coupled to a DS2 isolated stimulator. The average amplitude of the evoked or spontaneous postsynaptic currents per voltage step were measured from 2-3 consecutive trials and plotted into a graph, then a linear fit was established in order to determine the value of E_GABA for each cell tested. To confirm the perforated patch configuration negative pressure was applied at the end of the recording to rupture the seal and establish the whole cell configuration. Equilibration of [Cl⁻], with the pipette solution (dialysis) occurred within a few minutes of seal rupture, whereupon the value of E_GABA reached the value predicted by the Nerst equation (-70 mV). All of the above analyses were performed using pClamp 10.0 and Sigmaplot 10.0 software.

In vivo extracellular recordings

Male F344 rats (250 – 350g) were anaesthetized by i.p. injection of urethane (1.25 g kg⁻¹, Sigma). A catheter was inserted into the left femoral vein for drug injection. A Telemetry Research-Millar BP transducer (Telemetry Research, Auckland, New Zealand) was inserted into the ascending aorta through the right femoral artery to monitor BP. The pituitary stalk and right supraoptic nucleus were exposed by a ventral approach through the oral cavity. Extracellular single-unit recordings were made using a glass recording microelectrode (15-40 MΩ) filled with 0.9% saline. A side-by-side RMI SNEX-200 stimulating electrode (Science products GmbH, Hofheim, Germany) was placed on the stalk of the pituitary gland and used to elicit antidromic spikes in neurons that project an axon to the pituitary gland, thus characterizing them as MNCs. BP and MNC activity were recorded via a CED 1401 ADC interface using Spike2 software.
(Cambridge Electronic Design, Cambridge, UK) and analyzed offline. MNCs were characterized as VP neurons by their spontaneous phasic activity or by a lack of excitation following IV injection of cholecystokinin (CCK; 20 µg kg\(^{-1}\), 0.5 ml kg\(^{-1}\) in 0.9% saline), or as OT neurons by transient excitation following CCK injection (Sabatier et al., 2004). At the end of the experiments, the rats were killed by anaesthetic overdose (60 mg kg\(^{-1}\) pentobarbitone or 1 g kg\(^{-1}\) urethane, IV).

**Sodium Dodecyl Sulfate (SDS) Polyacrylamide Gel Electrophoresis (SDS/PAGE) and Western Blotting**

Brains of age-matched EU and 7-day SL Long Evans and Sprague-Dawley rats were rapidly removed and submerged into oxygenated isosmotic ACSF, and then supraoptic tissue blocks (1 mm\(^3\)) were excised using a pair of angled spring scissors. Each sample consisted of 4 supraoptic nuclei blocks collected from 2 rats. For experiments involving anterior hypothamic tissue, each sample consisted of one block of anterior hypothalamus from a single rat. Tissue blocks were then triturated and homogenized in a HEPES-based buffer containing protease inhibitor (Roche) and 1.0% Triton-x-100 (Sigma) and incubated at 4 °C for 30-60 minutes. Following centrifugation at 13,000 r.p.m. for 15 min, protein concentrations of the supernatant was measured using Bio-rad Dc protein assay and Ultrospec 2100 pro UV/visible spectrophotometer. DDT (Sigma) and loading buffer containing Bromophenyl blue (Sigma) was added to the supernatant and was incubated at 80 °C for 5 minutes. Samples were then loaded onto a NuPAGE 4-12 % Bis-Tris gel (Invitrogen) and proteins were separated by electrophoresis at 150 V for 1 h. The proteins were transferred to a Hybond-LFP PVDF membrane (GE healthcare, Piscataway, NJ, USA) in transfer buffer (192 mM glycine, 25 mM Tris-HCl, 20 % methanol, pH 8.3) at 70 V for 1 h. Membranes were blocked with Tris buffer containing 1% skim milk at 4 °C overnight, then were incubated with rabbit polyclonal antibodies against KCC2 (Milipore, Billerica, MA, USA; 1:500 for 1 h), BDNF (Santa Cruz biotechnology, Dallas, TX, USA; 1:1,000), pan-TrkB (Milipore; 1:1,000), or pTrkB (phospho Y515; Abcam, Cambridge, MA, USA; 1:100), followed by incubation with HRP goat anti-rabbit secondary antibody (Jackson ImmunoResearch, West Grove, PA, USA; 1:5,000). Incubation with the NKCC1 antibody (mouse monoclonal; generous gift from Dr. R. James Turner, NIH) was performed at the dilution of 1:500 for 1 h, and then with HRP goat anti-mouse secondary antibody (Jackson ImmunoResearch; 1:5,000). Detection was performed using an enhanced chemiluminescence kit (ECL; Perkin Elmer). Membranes were stripped by incubation with 100 mM glycine at pH 2.8 for 30 minutes and sequentially incubated with mouse monoclonal GAPDH antibody (Millipore; 1:5,000) for 1 h then HRP goat anti-mouse secondary antibody (Jackson ImmunoResearch; 1:5,000) for 30 minutes. ECL detection was again performed. Intensity of the bands was quantified using the gel analysis protocol in ImageJ software (NIH Scion Image) and the
intensities of KCC2, NKCC1, BDNF, tot-TrkB, and p-TrkB bands were quantified as ratios of GAPDH staining intensity.

Post-hoc immunohistochemistry of hypothalamic explant preparations

Hypothalamic explants were fixed by immersion in 4% paraformaldehyde in 0.1 M PBS for 7-9 days once electrophysiology was completed in order to test the efficiency of TrkB-Fc infusion. After fixation they were then incubated in 20% sucrose in PBS for 5 days. The explants were sectioned coronally on a Leica cryostat at 30 µm and direct mounted onto gelatin coated slides. The sections were then permeabilized and blocked with 3% horse serum and 0.25% Triton-X in 0.1 M PBS for 2 hours at room temperature. This was followed by 24 h incubation with rabbit p-TrkB antibody (TrkB515 ab51187; Abcam; 1:500) and guinea pig VP antibody (Peninsula Laboratories San Carlos, CA, USA; 1:10,000), at 4°C. After washing, the sections were washed and incubated with biotinylated goat anti-rabbit secondary antibody (Vector Labs, Burlingame, CA, USA; 1:100) and donkey anti-guinea pig CY3 secondary (Jackson ImmunoResearch; 1:1,250). Finally the slides were incubated with streptavidin conjugated with Dylight 488 (Vector Laboratories; 1:1,250) at room temperature. Slides were coverslipped with Vectashield Hardset media (Vector Labs). Sections were analyzed using an Olympus IX-2 DSU confocal microscope equipped with epifluorescence and excitation/emission filter sets to visualize Cy3 and Dylight 488 labeled secondary antibodies. Images were captured using a Q-imaging Retiga-SRV camera. Images were pseudocolored and the numbers of p-TrkB positive profiles were counted for each side on each image using ImageJ software, and then average cell counts were calculated for each rat.

AAV-mediated knockdown of BDNF in the supraoptic nucleus

Adeno-associated virus serotype 2 conjugated with shRNA directed against BDNF were custom-generated at high titer (1.0 x 10^{13} GC/ml; Vector Biolabs, Philadelphia, PA, USA). The targeting sequence against rat BDNF was ACCATAAGGACGCGGACTTGT (Sadri-Vakili et al., 2010) and the expression was driven by the U6 promoter. To test the effectiveness of the shRNA’s ability to induce a specific knockdown of BDNF in vitro, HEK 293T cells were transfected with 2 µg of BDNF with or without BDNF-shRNA 24h after plating. The media was changed 16h after transfection then collected 24h later. The conditioned media was then concentrated in an Amicon centrifuge tube and 2x laemli sample buffer was added. The samples were then run in a Western blot to detect whether the amount of BDNF secreted in the conditioned media is reduced when co-expressed with BDNF shRNA. The effectiveness of AAV-BDNF-shRNA to induce a knockdown of BDNF in hypothalamic tissue was tested in a group of rats by making intracerebroventricular (i.c.v.) injections of the virus into the third ventricle (2 µL). Briefly,
Isoflurane-anaesthetised rats were placed in a stereotaxic frame where rectal temperature was maintained at 37 °C by a thermostatically controlled electric blanket. Through a craniotomy, the tip of a 10 μL Hamilton syringe was lowered into the third ventricle stereotaxically (AP: -0.5 mm from Bregma, ML: 0 mm, P: 5 mm). Another group of rats received i.c.v. injections of equal amounts of high-titer AAVs conjugated with a scrambled (scr) sequence of shRNA (Vector Biolabs) as controls. After allowing in vivo expression for 3 weeks, last of which the rats also underwent the SL treatment, the rat brains were harvested and the anterior hypothalami were microdissected. The tissue blocks were homogenized and processed for Western blotting as described in the above section. To induce a specific knockdown of BDNF shRNA in the supraoptic nucleus, rats received 1 μL of buffer solution containing AAVs conjugated with either BDNF- or scr-shRNA was loaded onto a glass pipette and injected into the supraoptic nucleus using stereotaxic coordinates AP: -0.5 mm from Bregma, ML: 1.5 mm, and P: 7.7 mm. The virus was allowed to express in vivo for 5-6 weeks, after which the rats underwent a 7-day SL treatment. Their brains were harvested and prepared into hypothalamic slices as described above. After patch-clamp recordings, the slices were immersion-fixed in 4% paraformaldehyde overnight and processed for post-hoc immunohistochemistry to confirm the success of knockdown.

**Immunohistochemistry**

Age-matched EU and 7-day SL Long Evans rats were deeply anesthetised by IP injection (0.7 ml/100 g) of a solution containing urethane (0.25 g/ml) and perfused via the left ventricle with 500 ml (~ 5ml/min) of PBS containing 4% paraformaldehyde (Sigma). Coronal brain sections (50 μm thick) were made in PBS buffer using a vibratome. Sections were stained using antibodies against BDNF (rabbit polyclonal; Santa Cruz; 1:300), KCC2 (rabbit polyclonal; Millipore, USA; 1:400), VP-neurophysin (mouse monoclonal; kindly provided by Dr. Hal Gainer, NIH; 1:100), or Iba1 (rabbit polyclonal; Wako chemicals, Japan; 1:1,000). Sections were then incubated in secondary antibodies against rabbit IgG conjugated with Alexa Fluor 568 (Molecular Probes, Eugene, OR, USA; 1:500) or goat anti-mouse conjugated with Alexa Fluor 488 (Molecular Probes, Eugene, OR, USA; 1:500). The sections were then mounted in SlowFade® Gold Antifade reagent (Invitrogen, USA) and observed under Olympus FV1000 scanning laser confocal microscope equipped with a krypton/argon mixed gas laser. Laser lines emitting at 488 and 568 nm were used to excite the Alexa 488- and 568-conjugated secondary antibodies. Images of immunostained structures were captured as continuous stacks of confocal images (1 μm thick) using a custom acquisition software (Olympus). For analysis of Iba1 positive microglial structures, each confocal image was thresholded to isolate the immunopositive signals, and average staining intensities in circled areas containing the supraoptic nucleus were calculated from 4 sections per rat. For analysis of BDNF staining, circles
were drawn over the supraoptic nucleus in each confocal image, and average staining intensities were quantified. The values were then averaged between 5 continuous sections, each 5 μm apart. All of the analysis was performed with ImageJ software (NIH Scion Image).

*Inhibition of microglial activation via intraperitoneal injections of minocycline*

Minocycline hydrochloride (Sigma) was dissolved in 0.9% saline and appropriate amounts of NaOH were added to adjust the pH to ~7.4. Each rat received a daily dose of 50mg/kg i.p. for 7 days, during which they also underwent SL treatment. A control group of rats received injections of equal amounts of saline. On the last day of SL, the rat brains were harvested and made into hypothalamic slice preparations, and patch-clamp recordings were performed.

*Statistics*

All values in this study are reported as mean plus or minus the standard error of the mean (± s.e.m.). Unless explicitly stated, statistical differences between mean values were tested using Student’s two-tailed t-test. Mann-Whitney rank sum test, Wilcoxon Signed Rank Test, Student’s paired t-test, chi-square test and ANOVA with Student-Newman-Keuls posthoc test were performed with datasets where appropriate. All of the above statistical tests were performed with Sigmaplot 12.3 software (SPSS Inc., Chicago IL) or Prism 5 software (Graphpad Software Inc., La Jolla, CA, USA). Data points above or below the twice the standard deviation from the mean were excluded from statistical comparisons. Differences between values were considered to be significant when \( p < 0.05 \).
SUPPLEMENTAL REFERENCES


