Exercise training normalizes enhanced glutamate-mediated sympathetic activation from the PVN in heart failure

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Kleiber AC, Zheng H, Schultz HD, Peuler JD, Patel KP. Exercise training normalizes enhanced glutamate-mediated sympathetic activation from the PVN in heart failure. Am J Physiol Regul Integr Comp Physiol 294: R1863–R1872, 2008. First published April 2, 2008; doi:10.1152/ajpregu.00757.2007.—Exercise training (ExT) normalizes the increased sympathetic outflow in heart failure (HF), but the mechanisms are not known. We hypothesized that ExT would normalize the augmented glutamatergic mechanisms mediated by N-methyl-D-aspartic acid (NMDA) receptors within the paraventricular nucleus (PVN) that occur with HF. Four groups of rats were used: 1) sham-operated (Sham) sedentary (Sed), 2) Sham ExT, 3) HF Sed, and 4) HF ExT. HF was induced by left coronary artery ligation, and ExT consisted of 3 wk of treadmill running. In α-chloralose-urethane-anesthetized rats, the increase in renal sympathetic nerve activity in response to the highest dose of NMDA (200 pmol) injected into the PVN in the HF Sed group is approximately twice that of the Sham Sed group. In the HF ExT group the response was not different from the Sham Sed and Sham ExT groups. Relative NMDA NR1 receptor subunit mRNA expression was 63% higher in the HF Sed group compared with the Sham Sed group but in the HF ExT group was not different from the Sham Sed and Sham ExT groups. NR1 receptor subunit protein expression was increased 87% in the HF Sed group compared with the Sham Sed group but in the HF ExT group was not significantly different from the Sham Sed and Sham ExT groups. Thus one mechanism by which ExT alleviates elevated sympathetic outflow in HF may be through normalization of glutamatergic mechanisms within the PVN. A characteristic feature of heart failure (HF) is increased sympathoexcitation, which correlates with the severity of the disease as well as complications and mortality (28). The source of the increased sympathoexcitation associated with HF is not entirely understood, although several lines of evidence point to a role for the central nervous system (4, 16, 19). Sinoaortic (4) or cardiovagal (19) denervation does not normalize plasma norepinephrine (NE) levels in denervated compared with intact HF dogs. Plasma NE is also not altered in HF dogs by β-adrenergic blockade (19). Stimulation of the aortic depressor nerve produces altered lumbar sympathetic nerve activity in rats with HF compared with sham-treated rats (16). Together these data indicate an alteration within the central nervous system in HF.

The paraventricular nucleus (PVN) of the hypothalamus is reciprocally connected to other areas of the brain involved in control of cardiovascular function (18) and contains preautonomic neurons that project to sympathetic preganglionic neurons within the intermediolateral cell column of the spinal cord both directly and indirectly via the rostral ventrolateral medulla (40). Previous data from this laboratory suggest that the increased activity of PVN neurons associated with HF (32) is due to an increase in glutamatergic mechanisms within the PVN (21). The increase in renal sympathetic nerve activity (RSNA) in response to N-methyl-D-aspartic acid (NMDA) injected into the PVN is greater in HF rats than in sham-treated rats (21), which correlates with an increased expression of the NR1 subunit of the NMDA receptor within the PVN (21).

Exercise training (ExT) in HF patients increases survival, decreases complications, and decreases muscle sympathetic nerve activity (13). In a rapid pacing model of HF in rabbits, ExT decreases RSNA to that of normal rabbits (23). However, the mechanism by which ExT normalizes sympathetic outflow in HF is not known. In the present study, we hypothesized that one mechanism by which ExT restores sympathetic outflow to normal levels in HF is by normalizing the increased glutamatergic mechanisms within the PVN. Specifically, we wanted to determine whether ExT attenuated the increased RSNA response to NMDA injected into the PVN and whether ExT normalized the elevated expression of the NR1 subunit of the NMDA receptor in HF rats.

METHODS

Animals. Male Sprague-Dawley rats weighing 220–280 g (Sasco Breeding Laboratories, Omaha, NE) were fed and housed according to institutional guidelines. Protocols were approved by the University of Nebraska Institutional Animal Care and Use Committee and were in accordance with the American Physiological Society’s “Guiding Principles in the Care and Use of Animals.” Rats were given rat chow and water ad libitum and were housed in a room with a 12:12-h light-dark cycle. Rats were allowed to acclimatize for 1 wk before cardiac surgery.

Induction of heart failure. Rats were randomly assigned to either the sham-operated control group or the HF group. HF was induced by ligation of the left coronary artery as described previously (32). Left ventricular dysfunction was assessed with hemodynamic and anatomic criteria. Echocardiograms were performed before and after the 3-wk ExT period. Left ventricular end-diastolic pressure (LVEDP) was measured after the ExT period with a Mikro-Tip catheter (Millar Instruments, Houston, TX) inserted into the left ventricle via the right carotid artery at the time of the terminal experiment. To measure infarct size, the heart was dissected free of adjacent tissues, and the atria were removed. The right ventricle was opened with a lengthwise incision such that the heart was flattened with the left ventricle lying...
in the middle and the right ventricle on either side of it. The right ventricle was removed and the remaining left ventricle laid flat. A digital image of the left ventricle was captured with a Kodak DC290 digital camera (Kodak, Rochester, NY), and the infarcted area and total left ventricle area were quantified with SigmaScan Pro. Infarct size (%) was found by dividing the size of the infarcted area by the total size of the left ventricle. Rats with elevated LVEDP (>15 mmHg) and infarct size >30% of total left ventricle wall were considered to be in HF. Urine NE concentration was measured by radioenzymatic assay according to previously described protocols (36). Sensitivity of the assay was 1–2 pg of NE per 50 ml of diluted urine (1:100). Plasma angiotensin II (ANG II) was measured with an RIA (Alpco Diagnostics, Salem, NH). Sensitivity of the ANG II assay was 0.68 pg/ml.

Exercise training. Three weeks after coronary artery ligation surgery or sham surgery, rats were randomly assigned to either ExT or sedentary (Sed) groups to produce four total experimental groups: sham-operated (Sham) Sed, HF Sed, Sham ExT, and HF ExT. For ExT, rats ran on a motor-driven treadmill (Columbus Instruments, Columbus, OH) for a period of 3 wk according to a modified protocol of Musch and Terrell (27). Renal nerve recording or NR1 expression experiments were performed 6–8 wk after coronary artery ligation surgery or sham surgery, and thus the ExT period took place during the final 3 wk of the progression of HF. Initially, low speed (10 m/min) and grade (0%) and short duration (10 min/day) were used to familiarize the rats with running on the treadmill. The speed, duration, and grade were gradually increased to 20–25 m/min, 60 min/day, and 5–10%, respectively, to ensure that a significant endurance effect was produced. This level of exercise is considered moderate for the Sham rats. Only rats that ran steadily with little or no prompting were used in the study. To ensure a similar level of ExT between groups, citrate synthase activity assays on the soleus muscle were performed according to the protocol of Serere (44).

General surgery for hemodynamic and RSNA measurements and microinjection. Experiments were performed 6–8 wk after HF or sham surgery. Rats were anesthetized with urethane (0.75 g/kg ip) and α-chloralose (70 mg/kg ip). The left femoral artery was cannulated with polyethylene tubing (PE-50) for injection of supplemental anesthesia. The left femoral arteriole was cannulated and connected via a pressure transducer (Gould P23 ID) to a computer data-recording and analyzing program (PowerLab) to record mean arterial blood pressure (MAP) and heart rate (HR).

The left kidney was exposed through a retroperitoneal flank incision. A branch of the renal nerve was isolated from fat and connective tissue. The central end of the nerve was placed on thin bipolar platinum electrodes. The nerve-electrode junction was fixed and electrically insulated from surrounding tissues with a Wacker Sigel mixture (604 and 601). The electrical signal was amplified with a Grass amplifier with high- and low-frequency cutoffs of 1,000 and 100 Hz, respectively. The rectified output from the amplifier was displayed, using the PowerLab system to record and integrate the raw nerve discharge. Basal nerve activity was determined at the beginning of the experiment, and background noise was determined by nerve activity recorded at the end of the experiment (after the rat was euthanized). The nerve activity during the experiment was calculated by subtracting the background noise from the recorded value. The RSNA response to injection of drugs into the PVN was expressed as a percent change from the basal value.

For placement of microinjection cannulas into the PVN, the anesthetized rat was placed in a stereotaxic apparatus (David Kopf Instruments, Tujunga, CA). A longitudinal incision was made on the head, and bregma was exposed. A small burr hole was made in the skull to allow access to the PVN. The coordinates for the PVN, determined with the Paxinos and Watson atlas (33), were 1.5 mm posterior to bregma, 0.4 mm lateral to midline, and 7.8 mm ventral to the dura. A thin needle (0.5-mm OD, 0.1-mm ID) connected to a 0.5-ml microsyringe (Hamilton, Reno, NV) was lowered into the PVN. NMDA (Calbiochem, La Jolla, CA) was injected into the PVN in three doses (50, 100, and 200 pmol in 50–200 nl in random order. Each animal received all three doses of NMDA. Subsequent injections were made at least 20 min after prior injections to allow MAP, HR, and RSNA to return to basal levels.

Brain histology. After microinjection experiments, Chicago blue dye (50 nl) was injected into the brain to histologically verify that the injection site was located within the PVN. The brains were removed, fixed in 10% formalin for at least 24 h, sectioned (30 μm) on a cryostat, and processed for histology. The sections were mounted on gel-coated microscope slides and stained with 1% neutral red. The location of the injection was visualized on a microscope, and injections with terminations within the boundaries of the PVN were considered to be appropriately targeted to the PVN. Injections located outside of the PVN were excluded from data for the PVN and were analyzed as anatomic controls.

Micropunch of PVN and isolation of mRNA for real-time RT-PCR and protein for Western blot measurements. The following experiments were performed in a group of animals separate from those used in the in vivo microinjection experiments described above. After the animal was euthanized, the brain was removed and quickly frozen on dry ice. Six serial coronal sections (100 μm) were cut through the hypothalamus at the level of the PVN with a cryostat, and, according to the Palkovits and Brownstein technique (29), the PVN was bilaterally punched with a diethyl pyrocarbonate (DEPC)-treated blunt 18-gauge needle attached to a syringe, such that there were 12 total punches per brain. For each brain, six of the punches were placed in 500 μl of Tri-Reagent (MRC, Cincinnati, OH), followed by sonication and extraction of mRNA according to the manufacturer’s instructions. Briefly, the homogenate was separated into organic and aqueous phases by the addition of bromochloroform and subsequent centrifugation. The RNA, contained in the aqueous phase, was precipitated with isopropanol, washed with ethanol, and solubilized in 10 μl of DEPC-treated water. The other six punches for each brain were placed in 100 μl of protein extraction buffer (1 M Tris, 0.5 M EDTA, 10% SDS, Triton-X-100, and 100 mM phenylmethylsulfonyl fluoride), sonicated, and incubated for 30 min at 37°C to extract the protein. The supraoptic nucleus (SON), lateral hypothalamus (LH), and dorsal cortex were punched at the same rostrocaudal level as the PVN to serve as anatomic control regions from the same cross-sectional slices.

Real-time RT-PCR measurement of NR1 mRNA. After extraction of mRNA, samples underwent reverse transcription for 40 min at 37°C in the presence of 1.5 M random hexamers and 100 U MMLV-RTase. Real-time RT-PCR measurements were made with the iCycler iQ Multicolor Real-Time Detection System with output to a computer-based acquisition system (Bio-Rad). The protocol consisted of denaturation (95°C for 3 min), amplification and quantification repeated 50 times (95°C for 10 s, 55°C for 45 s), denaturation at 95°C for 1 min, reannealing at 55°C for 1 min, and a melt curve (55–95°C with a heating rate of 0.5°C per 10 s). The reaction mixture consisted of SYBR Green Supermix (Bio-Rad), 300 nM sense primer, 300 nM antisense primer, DEPC-treated H2O, and the cDNA template of interest. For NR1, the sense primer was 5′-ATAGTGCAATTCAC-CAAGAAGCC and the antisense primer was 5′-TAGCTGGCCCATCCTCCGT. For rp19, used as the reference gene, the sense primer was 5′-CCCCAATGGAAACCAACGAA and the antisense primer was 5′-AGTGACATGCAGGCTCT. Relative expression of NR1 was calculated with the Pfaffl equation, which relates expression of the target gene (NR1) to expression of a reference gene (rp19) (34).

Western blot measurement of NR1 protein. The total protein concentration from the extracted protein described above was measured with a bicinchoninic acid assay kit (Pierce, Rockford, IL). Samples were adjusted to contain the same concentration of total protein, and then equal volumes of 2× 4% SDS sample buffer were added. The samples were boiled for 3 min and then loaded onto a 7.5% SDS-PAGE gel (40 μg/20 μl per well). Gels were subjected to electro-
phoresis at 40 mA/gel for 60 min. The fractionated proteins on the gel were electrophoretically transferred to a polyvinylidene difluoride membrane (Millipore, Billerica, MA) at 300 mA for 90 min. The membrane was probed with primary antibody [rabbit anti-NR1 (1:500 dilution, Santa Cruz), rabbit anti-NR2B (1:500, Santa Cruz), or rabbit anti-β-tubulin (1:2,000, Santa Cruz)], washed with Tris-buffered saline-Tween 20, and then probed with secondary antibody (peroxidase-conjugated goat anti-rabbit IgG, 1:5,000 dilution; Pierce). An enhanced chemiluminescence substrate (Pierce) was applied to the membrane for 5 min, followed by a 30-s exposure within an Ep Chemi II Darkroom (UVP Bioimaging, Upland, CA) for visualization with the Worklab digital imaging system. Kodak ID software was used to highlight the bands and quantify the signal. The expression of NR1 or NR2B was calculated as the ratio of intensity of the NR1 or NR2B band, respectively, relative to the intensity of the β-tubulin band.

Data analysis. Data are presented as means ± SE. The data were subjected to two-way ANOVA followed by comparison for individual group differences with the Newman-Keuls test (7). Statistical significance was indicated by a value of \( P < 0.05 \).

RESULTS

General data. Table 1 presents baseline MAP and HR data as well as morphological characteristics and left ventricular function data among the four experimental groups. The data represent mean values from animals used for RSNA experiments and animals in which mRNA and protein expression within the PVN were measured. Baseline RSNA (integrated RSNA) was not different between groups. However, baseline RSNA varies from animal to animal and is therefore difficult to compare between animals. Only rats with ≥30% infarct of the left ventricular wall were included in the study. Five rats had infarct sizes <30% and were excluded from data analysis. Sham rats had no visible myocardial damage. LVEDP was significantly increased in the HF Sed rats compared with both Sham groups and the HF ExT group. HF Sed rats had significantly lower change in pressure over time (dP/dt) compared with Sham rats, which was partially improved by ExT. While Sham rats experienced cardiac dysfunction and that ExT did not normalize cardiac function.

Additionally, citrate synthase activity is presented in Table 1. Citrate synthase activity of the soleus muscle was significantly higher in both ExT groups compared with the Sed groups, demonstrating a significant effect of ExT. ExT had a similar effect in the two groups in that there was no significant difference between Sham ExT and HF ExT.

Figure 1 illustrates plasma ANG II and urinary NE data. Plasma ANG II levels were increased in HF Sed compared with Sham Sed rats but were not different in HF ExT compared with Sham Sed or Sham ExT rats, indicating that ExT normalized the elevated plasma ANG II associated with HF (Fig. 1A). Urinary NE levels were elevated in HF Sed rats compared with Sham Sed rats, indicative of increased overall sympathoexcitation. In HF ExT rats, urinary NE levels were not different from Sham Sed or Sham ExT rats, suggesting that ExT normalized the increased overall sympathetic outflow associated with HF (Fig. 1B).

Figure 2 is a schematic illustrating the location of injection sites of NMDA into the PVN. Of 30 microinjections, 24 were located within the PVN and 6 were outside of the PVN and therefore analyzed separately as anatomic controls. Administration of NMDA into these sites (anatomic controls) did not produce the typical increases in RSNA in response to NMDA injected into the PVN.

Microinjection of NMDA into PVN. Figure 3A illustrates examples of responses of RSNA, MAP, and HR to microinjection of the highest dose of NMDA (200 pmol) into the PVN in each of the four experimental groups. NMDA injected into the PVN elicited dose-dependent increases in RSNA in all groups, as shown in the group data in Fig. 3B. At the lowest dose of NMDA (50 pmol), there was a trend for the RSNA response to be potentiated in the HF Sed group. However, the mean response in that group was not significantly different from that in the Sham Sed group. Similarly, there was a trend for the RSNA response to 50 pmol of NMDA in the HF ExT group to be reduced compared with the HF Sed group, although there was not a statistically significant difference between the two groups. At the two higher doses of NMDA, the increase in RSNA was significantly potentiated in the HF Sed group consistent with previous work (21). At 100 pmol, the response in HF Sed rats was significantly higher than in Sham Sed and Sham ExT rats. The response to 100 pmol in the HF ExT group was significantly lower than in the HF Sed group and not different from either Sham group.

At the highest dose of NMDA (200 pmol), the response in the HF Sed group (93 ± 13%, \( n = 6 \)) was significantly higher \( (P < 0.05) \) than that in Sham Sed (45 ± 2%, \( n = 6 \)) and Sham ExT (33 ± 9%, \( n = 6 \)) groups. In the HF ExT group, this response (39 ± 5%, \( n = 6 \)) was significantly lower \( (P < 0.05) \) than in the HF Sed group and not different from either of the Sham groups. These data indicate that ExT normalizes the RSNA response to NMDA microinjection into the PVN in rats with HF.

Figure 4A illustrates the increase in MAP in response to NMDA injected into the PVN. At 100 pmol of NMDA, the increase in MAP in the HF Sed group was significantly higher compared with the Sham Sed and Sham ExT groups. The
increase in MAP in the HF ExT group was significantly attenuated compared with the HF Sed group and not different from either of the Sham groups. At the highest dose of NMDA (200 pmol), the increase in HR in the HF Sed group (31 ± 5 beats/min, n = 6) was significantly higher than the responses in the Sham ExT (10 ± 7 beats/min, n = 6) and HF ExT (9 ± 6 beats/min, n = 6) groups but was not statistically different from that of the Sham Sed group (18 ± 2 beats/min, n = 6).

Expression of NR1 subunit mRNA and protein within the PVN. NR1 mRNA expression, measured by real-time RT-PCR, is shown in Figs. 5 and 6. Examples of original real-time RT-PCR traces comparing Sham Sed and HF Sed groups (Fig. 5A) and HF Sed and HF ExT groups (Fig. 5B) are shown in Fig. 5. Traces for the reference gene rpl19 are shown in insets and traces for the target gene NR1 in main images in Fig. 5. Figure 6 shows the composite real-time RT-PCR data for the four experimental groups. Relative NR1 expression was significantly increased in the HF Sed group compared with the Sham Sed group. However, in the HF ExT group, relative NR1 expression was significantly lower than in the HF Sed group and not different from the Sham Sed or the Sham ExT group. These data indicate that ExT normalizes NR1 mRNA expression within the PVN in rats with HF.

NR1 protein expression, measured by Western blot, is shown in Fig. 7. Sample gels showing NR1 and β-tubulin protein in the four experimental groups are presented in Fig. 7A. Figure 7B shows the composite data for NR1 expression within the PVN, calculated as the ratio of the density of the NR1 band to the density of the β-tubulin band. The level of NR1 protein expression in the HF Sed group was significantly higher than in the Sham Sed group. In the HF ExT group, NR1 protein expression was significantly lower than in the HF Sed group and was not different from either the Sham Sed group or the Sham ExT group.

Figure 8 shows NR1 protein expression within the SON, LH, and cortex and NR2B protein expression within the PVN. SON, LH, and cortex samples were punched from the same coronal sections as the PVN to be used as control areas. There were no significant differences in NR1 protein expression within the SON, LH, or cortex among all of the experimental groups (Fig. 8A), indicating that the changes in NR1 expression by HF and ExT are specific to the region of the PVN. There were also no significant differences in PVN NR2B protein expression among the four experimental groups (Fig. 8B),
suggesting that the changes in expression within the PVN are specific for the NR1 subunit. Overall, these data show that ExT normalizes the increased expression of NR1 within the PVN in rats with HF.

**DISCUSSION**

In the present study we found that ExT normalizes the potentiated increase in RSNA in response to NMDA microinjected into the PVN in rats with HF. Further, while expression of the NR1 subunit of the NMDA receptor within the PVN was increased in HF rats in agreement with previous data, the present results demonstrate that NR1 expression in the PVN of HF ExT rats was not different from that in Sham Sed or Sham ExT rats. This normalization of NR1 expression within the PVN suggests a mechanism by which the RSNA responses to NMDA injected into the PVN are normalized by ExT in HF rats. Together, these results indicate that one mechanism by which ExT normalizes sympathetic outflow in HF is normalization of glutamatergic mechanisms within the PVN.

The coronary artery ligation model of HF has been used extensively by this laboratory (51) and others (35). The utility of this model as a simulation of HF is demonstrated by increased LVEDP, decreased dP/dt of the left ventricle, decreased ejection fraction, and 30% infarct size at the time of the experiment, 6–8 wk after the coronary artery ligation procedure. The advantage of using this model as opposed to other models of HF, such as ventricular pacing, is that ligation of the coronary artery mimics blockage of the artery, commonly seen in patients with HF. Additionally, in the present study, we found that ExT only partially improved the increased LVEDP and the decreased dP/dt and did not improve the decreased ejection fraction associated with HF, indicating that ExT did not normalize cardiac function per se in this model of HF.

We also analyzed the efficacy of our regimen of ExT. Citrate synthase activity is commonly used as a marker for the metabolic capacity of muscle (43) and is increased in skeletal muscle by exercise (11). We measured citrate synthase activity

![Graph showing the effect of NMDA injections on RSNA, MAP, and HR.](http://ajpregu.physiology.org/)
in the soleus muscle in all four groups and found that it was increased in both of the ExT groups compared with the Sed groups, indicating not only an effect of the ExT itself but also a similar level of ExT in the Sham ExT and HF ExT groups.

The PVN has a major role within the forebrain, with a role in direct regulation of sympathetic outflow (48). Preautonomic neurons originating in the dorsal and lateral parvocellular divisions of the PVN project to sympathetic preganglionic neurons in the intermediolateral cell column of the spinal cord directly (40) and indirectly via the rostral ventrolateral medulla (40). Importantly, spinally projecting preautonomic PVN neurons have been shown to be sensitive to NMDA (5, 37). The PVN is involved in fluid balance and vasopressin release (48), whereas the activity of PVN neurons is increased in HF rats (32). Given these characteristics of the PVN, as well as the increase in sympathetic outflow (10, 31) and salt and water retention in HF (8), there is good evidence that the PVN is involved in the increased sympathetic activity associated with HF (30). The results of the present study are in agreement with this idea: the increase in sympathetic nerve activity in response to NMDA injected into the PVN, as well as the expression of NR1 within the PVN, is increased in HF rats compared with sham-operated rats.

In our NMDA microinjection experiments, it is possible that the spread of NMDA reached areas surrounding the PVN. Histological examination of the spread of dye injected at the end of each experiment revealed that the injections were limited to the PVN, although it is possible that the areas of spread of NMDA and dye did not precisely overlap. Thus, while the spread of dye was limited to the PVN, it is possible that NMDA affected regions outside the PVN, notably the dorsomedial nucleus of the hypothalamus, which is located rostrally to the PVN (33) and also has a significant role in modulation of sympathetic nerve activity, MAP, and HR (6). In our experiments in which the injection was found to be located outside of the PVN, the typical increase in RSNA, MAP, and HR in response to NMDA was not observed, indicating that the areas affected by NMDA in these animals were not involved in control of RSNA, MAP, or HR. This evidence indicates that the increases in RSNA, MAP, and HR in response to NMDA microinjection were most likely mediated by neurons located within the PVN.

A functional NMDA receptor comprises the NR1 subunit and at least one NR2 subunit (9, 45). In the present study we showed that expression of the NR1 subunit increases in the PVN in HF, and this increase is prevented with ExT. We also measured protein expression of an NR2 subunit that is abundant in the PVN, NR2B. Previous data from our laboratory (21) indicate that expression of NR2B within the PVN is not altered by HF. In the present experiments, our data indicate that NR2B also is not altered by ExT. This suggests that changes in PVN subunit expression, with implications for NMDA receptor function, may be limited to NR1. It is possible, however, that expression of at least one other NR2 subunit may be altered by HF and/or ExT. We did not examine expression of other NR subunits (NR2A, NR2C, NR2D, NR3A, or NR3B). The NR2A subunit is also expressed within the PVN, albeit less densely than NR2B (14). While the NR2C and NR2D subunits are also expressed within the PVN, the expression of the NR2B subunit is considerably greater, particularly in neurons within the parvocellular subdivisions (14). Nonetheless, it is possible that HF and/or ExT alter the expression of one or more of these subunits to affect the function of NMDA receptors, which remains to be examined.

In addition, while we did not measure glutamate levels in the present study, we previously demonstrated (21) that glutamate levels within the PVN are unaltered by HF. Thus the increased glutamatergic response within the PVN in HF is due to increased expression of NR1 and not to a change in concentration of the receptor agonist.

We also measured NR1 protein expression within the SON, another area within the hypothalamus important in cardiovascular control, and found that it was not significantly changed in HF [in agreement with previous data (21)] or as a result of ExT. This suggests that the changes in NR1 expression in hypothalamic cardiovascular control areas are selective for specific areas such as the PVN. Additionally, we measured NR1 protein expression within the lateral hypothalamus, which is adjacent to the PVN, and in the cortex, to serve as anatomic

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**Fig. 4.** A: mean changes in MAP following injections of NMDA into the PVN. B: mean changes in HR following injections of NMDA into the PVN. *P < 0.05 vs. Sham. #P < 0.05 vs. Sed.
controls and found that NR1 expression was not changed in either region as a result of HF or ExT, further lending support to our conclusion that changes in NR1 expression in HF and/or with ExT are anatomically specific to the PVN. Furthermore, when NMDA was injected into an area adjacent to the PVN, little or no change in RSNA was observed, indicating that the alterations in NMDA-mediated RSNA responses are specific and limited to the PVN.

The source(s) for the changes in NR1 expression within the PVN in HF and ExT is not entirely clear since there are multiple possibilities. Decreased cardiac output as a result of HF may increase stimulation of cardiopulmonary receptors, which send afferent neural input to the PVN. Humoral factors, which are both central and peripheral, are also changed in HF. For example, in the present study plasma ANG II concentration was increased in HF Sed rats compared with Sham Sed rats, but in the HF ExT rats the level of plasma ANG II was not different from that in the Sham Sed or Sham ExT groups. The ANG II type 1 (AT₁) receptor has been shown to participate in a variety of neuronal intracellular signaling pathways (for review see Ref. 46) and thus may play a role in modulation of NR1 gene expression. Although neurons within the PVN cannot directly detect the concentration of ANG II in the plasma, these neurons receive input from circumventricular organs, notably the subfornical organ (SFO) (26). PVN-projecting SFO neurons increase firing in response to ANG II (1), and the SFO-PVN projection is angiotensinergic (2). It is possible that the augmented plasma ANG II associated with HF chronically increases firing of these PVN-projecting SFO neurons, and this increased angiotensinergic input from the SFO to the PVN might alter expression of NR1 in PVN neurons. ExT, in turn, normalizes plasma ANG II and thus may normalize firing of PVN-projecting SFO neurons, which would restore the enhanced glutamatergic mechanisms within the PVN to control levels.

Another factor that may be involved in the alteration of glutamatergic mechanisms within the PVN is nitric oxide (NO), which acts as an unconventional neurotransmitter in the
central nervous system (39) and has been shown to regulate gene expression (3). We previously demonstrated (22) that prior injection of \( \text{NG} \)-monomethyl-L-arginine into the PVN significantly augments the increase in RSNA in response to NMDA injected into the PVN, suggesting negative feedback from NO on the NMDA system within the PVN. We have also shown that the inhibitory effect of NO within the PVN is blunted in HF, as is neuronal nitric oxide synthase (nNOS) expression (50), but these are normalized by ExT (52). The loss of negative feedback by NO due to the downregulation of nNOS within the PVN in the HF state may remove an inhibitory modulation of NO on the expression of NR1. Thus it is possible that the decrease in NO mechanisms in HF may contribute to the increase in glutamatergic mechanisms within the PVN. The restoration of nNOS within the PVN by ExT may restore the NO-mediated negative feedback on glutamatergic mechanisms, normalizing NR1 expression and NMDA-mediated changes in RSNA. NMDA receptor activation in the nucleus tractus solitarii has been shown to increase NO release (25). Thus, if this system also exists within the PVN, another possibility is that the release of NO by NMDA receptor activation is altered by HF. However, to our knowledge, NMDA receptor activation has not been shown to regulate NO release within the PVN in HF.

It is also possible that the effects of ExT on the PVN in HF are mediated by other mechanisms. For example, the mechanisms by which the exercise pressor reflex activates PVN neurons (20) and by which the reflex itself is altered in HF (42) are not understood, but it is possible that HF alters PVN neurons involved in the exercise pressor reflex. Other mechanisms that are altered by HF and ExT, such as baroreceptor (15), cardiopulmonary (36), chemoreceptor (47), or cardiac sympathetic afferent (49) reflexes, as well as input from central command (12) or from humoral factors such as atrial natriuretic factor (41) or vasopressin (38) may be involved as well.

In conclusion, the results of the present study demonstrate that ExT attenuates the potentiated increase in RSNA in response to NMDA injected into the PVN in HF rats. Normalization of expression of the NR1 subunit of the NMDA receptor may underlie this effect. Therefore, one mechanism by which ExT restores sympathetic outflow in HF is through normalization of glutamatergic mechanisms within the PVN.
Perspectives and Significance

HF is commonly characterized by an overall increase in sympathetic activation, an effect intended to compensate for the decrease in cardiac output but that ultimately exacerbates the condition. Uncovering the mechanism(s) involved in this increased sympathetic outflow may lead to improved treatment of this particular symptom of the disease, improving quality of life and survival. We have demonstrated, both previously and in the present work, that alterations within the glutamatergic system within the PVN play a role in the hyperactivity of the renal nerves in HF, thus identifying this system as a potential therapeutic target. Furthermore, among the numerous beneficial effects of ExT on the HF state is alleviation of the increased sympathetic activation. In the present set of experiments, we have shown that one of the ways ExT improves sympathetic outflow is through normalization of the enhanced glutamatergic mechanisms within the PVN. This understanding of the mechanisms of ExT-induced normalization of sympathoexcitation furthers our knowledge of the mechanisms underlying some of the symptoms of HF, offers a target in care of the disease, and emphasizes the importance of ExT in treatment of HF.

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