The Anteroventral Bed Nucleus of the Stria Terminalis Differentially Regulates Hypothalamic-Pituitary-Adrenocortical Axis Responses to Acute and Chronic Stress

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The anteroventral region of the bed nucleus of the stria terminalis (BST) stimulates hypothalamic-pituitary-adrenocortical (HPA) axis responses to acute stress. However, the role of the anterior BST nuclei in chronic drive of the HPA axis has yet to be established. Therefore, this study tests the role of the anteroventral BST in physiological responses to chronic drive, using a chronic variable stress (CVS) model. Male Sprague-Dawley rats received either bilateral ibotenate lesions of the BSTdm/fu, or vehicle injection into the same region. Half of the lesion and control rats were exposed to a 14-d CVS paradigm consisting of twice-daily exposure to unpredictable, alternating stressors. The remaining rats were nonhandled control animals that remained in home cages. On the morning after the end of CVS exposure, all rats were exposed to a novel restraint stress challenge. CVS induced attenuated body weight gain, adrenal hypertrophy, thymic atrophy, and adrenal hypertrophy (2-5). Chronic stress-induced physiological changes are associated with increased activation of the PVN, including elevated expression of CRH mRNA and, in some stress models, AVP mRNA (4-6). Chronic stress models typically engender elevated pre-acute stress plasma corticosterone (2, 4, 5), and can also lead to a sensitization of the HPA axis to a novel stress challenge, resulting in a facilitation of ACTH and corticosterone secretion despite an enhanced feedback signal (7-9). Excessive exposure to glucocorticoids during chronic stress is thought to be involved in numerous stress-related pathologies, including depression, posttraumatic stress disorder, and other anxiety disorders (10).

Prolonged stress causes numerous neuroendocrine and physiological changes that are connected to enhanced HPA axis drive, including decreased body weight gain, thymic atrophy, and adrenal hypertrophy (2-5). Chronic stress-induced physiological changes are associated with increased activation of the PVN, including elevated expression of CRH mRNA and, in some stress models, AVP mRNA (4-6). Chronic stress models typically engender elevated pre-acute stress plasma corticosterone (2, 4, 5), and can also lead to a sensitization of the HPA axis to a novel stress challenge, resulting in a facilitation of ACTH and corticosterone secretion despite an enhanced feedback signal (7-9). Excessive exposure to glucocorticoids during chronic stress is thought to be involved in numerous stress-related pathologies, including depression, posttraumatic stress disorder, and other anxiety disorders (10).

Forebrain limbic regions such as the amygdala, hippocampus, and medial prefrontal cortex are responsible for tuning HPA axis responses to stress (see Ref. 11). These limbic regions have limited to no direct innervation of the PVN (12-15) and instead use synaptically or multisynaptic connections through intervening structures such as the bed nucleus of the stria terminalis (BST) (12, 15). The BST receives limbic input from the amygdala, hippocampus, and prefrontal cortex (16-18), and has abundant projections to the PVN (15, 19-23). The dorsomedial (dm) and fusiform (fu) subnuclei, which are...
located in the anterior division of the BST ventral to the anterior commissure, send heavy projections to the medial parvoocular PVN (20, 22). Stimulation of the anterior or lateral areas of the BST increases HPA axis activity, whereas lesions of these areas decrease stress-induced ACTH and corticosterone secretion (24–30). Combined, these data suggest that these anterior BST structures are positioned to relay stress-excitatory information to the PVN.

The contribution of the BST to chronic stress adaptation has yet to be assessed. Given the prominent role of this structure in acute stress responsiveness, it is important to determine whether this region plays a role in the development or maintenance of chronic stress symptomology. Therefore, this current study uses selective lesions targeting the PVN-projecting dm/fu nuclei to test the hypothesis that these nuclei are necessary for sensitization of HPA axis responsivity after chronic stress. To test our hypothesis, this study uses a chronic variable stress (CVS) paradigm, followed by a novel restraint challenge to assess CVS-induced adaptations in HPA axis reactivity.

Materials and Methods

Experimental animals

A total of 54 adult male Sprague-Dawley rats (275–300 g; Harlan, Indianapolis, IN) was used for this study. All rats were housed three per cage in conventional shoebox rat cages with food and water available ad libitum in a temperature and humidity controlled vivarium on a 12-h light, 12-h dark cycle (lights on at 0600 h). Upon arrival, all rats acclimated to the animal facility for at least 7 d before surgery. Animals were maintained in accordance with the Guide for the Care and Use of Laboratory Animals (National Institutes of Health, 1996). All animal protocols were approved by the Institutional Animal Care and Use Committee at the University of Cincinnati.

Ibotenate lesions

Rats were anesthetized by ip injection of a 87 mg/kg ketamine/13 mg/kg xylazine mixture. Preemptive analgesia was administered by sc injections of 260 μg/kg butorphanol (Torbugesic, Fort Dodge Animal Health, Fort Dodge, IA). Each rat was mounted in a Kopf stereotaxic apparatus (David Kopf Instruments, Tujunga, CA). Skulls were exposed, and burr holes were drilled at the calculated surface coordinates. Each rat received bilateral microinjections of ibotenate (0.5 μl/side, 5.0 μg/μl) in sterile PBS (pH 7.4) or 0.9% sterile saline into the anterior BST [anterior-posterior (AP) –10.0 mm, medial-lateral ±1.5 mm, dorsal-ventral (dura) –7.0 mm], with coordinates calculated from bregma (31). Each microinjection used a 26-gauge 1.5 mm, 1.5 mm, dorsal-ventral coordinate over a 1-min period and left in place for 1 min before injection. The ibotenate or saline sample was manually infused over 5 min at a rate of 0.05 μl/30 sec, followed by another 5-min waiting period to allow diffusion and minimize dorsal spread of injection up the needle track. The syringes were slowly lowered to the dorsal-ventral coordinate over a 1-min period and left in place for 1 min before injection. The ibotenate or saline was manually infused over 5 min at a rate of 0.05 μl/30 sec, followed by another 5-min waiting period to allow diffusion and minimize dorsal spread of injection up the needle track. The syringes were raised over the course of 1 min. Skull burl holes were sealed with sterile bone wax, and the skin was closed with wound clips. Animals recovered for at least 7 d after surgery before CVS exposure. Body weights were measured on the day of surgery, d-1 CVS exposure, d-8 CVS exposure, d 15 after restraint stress challenge, and just before sacrifice.

CVS protocol

Rats were randomly assigned to “sham non-CVS” (n = 12), “sham CVS” (n = 12), “lesion non-CVS” (n = 15), and “lesion CVS” (n = 15) groups, in which “CVS” groups were exposed to the CVS paradigm, whereas “non-CVS” rats remained in their home cages as unhandled controls. The CVS paradigm consisted of twice-daily exposure to altering stressors for 14 consecutive days (d 1–14). Morning stressors were administered between 0930 and 1030 h, whereas afternoon stressors were conducted between 1430 and 1530 h. Occasional overnight stressors began immediately after cessation of afternoon stressors and ended with initiation of the next day’s morning stressor. CVS stressors consisted of hypoxia (30 min in 8% oxygen), cold stress (1 h at 4°C, two rats per cage without bedding), rotation stress (1 h at 100 rpm on a platform orbital shaker), warm swim (20 min at 31–33°C), cold swim (10 min at 16–18°C), overnight social isolation (one rat per cage), and overnight social crowding (six rats per cage). Stressors were unpredictable for the rats by being presented in a semi-randomized order, with each stressor (except the overnight stressors) representing an equivalent number of times.

Acute novel restraint stress protocol

All rats received restraint stress between 0830 and 1030 h on d 15, the day after the cessation of CVS. Animals were placed in well-ventilated Plexiglas restraint tubes (Altuglas International, Philadelphia, PA), and a tail clip blood sample (250–300 μl) was immediately collected for the determination of plasma ACTH and corticosterone levels. All rats remained in the restrainers for 20 min, at which point another blood sample was collected. Animals were then released back into their home cages to recover. An additional blood sample was taken at 40 min from the onset of the restraint stress. At 60 min from the onset of restraint, the rats were immediately killed by decapitation, and trunk blood was collected. Brains were removed, flash frozen in isopentane on dry ice (−45°C), and stored at −80°C. Adrenal and thymus glands were also collected and weighed.

RIA

Plasma corticosterone levels were measured by RIA using a kit from MP Biomedicals (Orangeburg, NY); plasma ACTH levels were measured by RIA using an antiserum donated by Dr. W. Engeland (University of Minnesota, Minneapolis, MN) and 125I-labeled ACTH (Amersham Biosciences, Piscataway, NJ) as tracer (22). For each hormone, all plasma samples were analyzed in duplicate within the same assay. Samples were randomized within assay runs. For the ACTH RIA, the intraassay coefficient of variation was 13% for plasma pools of 38 pg/ml. For the corticosterone RIA, the intraassay coefficient of variation was 8% for plasma pools of 120 ng/ml, which mirrors the manufacturer specifications.

Lesion verification

Brains were serially sectioned at 14 μm using a MICROM cryostat (MICROM International GmbH, Walldorf, Germany), mounted onto charged glass slides and stored at −20°C. Lesion sites were verified by Nissl staining of cells and neuronal nuclei (NeuN) immunolabeling as a neuronal marker. Sections were fixed in 4% paraformaldehyde and Nissl stained with cresyl violet, dehydrated through an ascending ethanol series, cleared in xylene, and coverslipped using DPX mountant. For NeuN immunohistochemistry, tissue sections were encircled by a hydrophobic slide marker (Super HT PAP pen; Research Products International Corp., Mt. Prospect, IL). The tissue sections were fixed in 4% paraformaldehyde and washed extensively with 50 mm potassium phosphate buffered-saline (KPBS) between each step. Subsequently, sections were incubated in 1% H2O2 for 10 min and blocked in incubation solution (4% normal goat serum and 0.3% Triton X-100 in KPBS) for 1 h. Sections were incubated overnight with a monoclonal antibody against NeuN (1:1000; CHEMICON International Inc., Temecula, CA) with Parafilm coverslips (American National Can Co., Greenwich, CT), followed by a 1-h incubation in biotinylated donkey antigoat IgG (1:500; Vector Laboratories, Burlingame, CA) and a 1-h incubation in avidin-horseradish peroxidase complex (1:500, ABC Elite Kit; Vector Laboratories), in which each step was in incubation solution. Finally, sections were incubated for 5 min in 0.02% diaminobenzidine (Sigma-Aldrich, St. Louis, MO), resulting in a brown reaction product. Slides were dehydrated through an ascending ethanol series, cleared in xylene, and coverslipped using DPX mountant.

Lesions were identified by the location of the needle track, loss of neurons, gliosis, and sparing damage to fibers of passage (33). Lesions targeting the anterior division of the BST were confirmed as “hits” when

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the primary damage included the fu and dm nuclei of the BST, with some occasional damage extending into the adjacent subcommissural zone of the BST, the ventral pallidum, and the parastriatal nucleus. Lesions that focused outside of the primary targeted dm/fu nuclei of the BST were considered misses. Rats with bilateral damage were included in the study, whereas rats with only partial unilateral lesions or missed lesions were removed from the analysis. The final "n" for experimental groups after lesion verification were: "sham non-CVS" (n = 12), "sham CVS" (n = 12), "lesion non-CVS" (n = 9), and "lesion CVS" (n = 10).

**In situ hybridization**

Antisense AVP, CRH, and c-fos riboprobes were generated by *in vitro* transcription using 35S-labeled uridine 5'-triphosphate (UTP). The rat AVP (exon C) cDNA construct is a 161-bp insert in a pCR4 TOPO vector, which was linearized with NotI restriction enzyme, and transcribed using T3 RNA polymerase. The rat c-fos cDNA construct (courtesy of Dr. T. Curran, Children’s Hospital of Philadelphia, PA) is a 587-bp fragment cloned into Bluescript SK vector, which was linearized with HindIII and transcribed with T3 RNA polymerase. The rat CRH cDNA construct is a 765-bp fragment cloned into pGEM3 vector (courtesy of Dr. Robert Thompson, University of Michigan, Ann Arbor, MI), which was linearized with HindIII and transcribed using T7 RNA polymerase. Each 15-μl riboprobe transcription reaction was made from 1.0- to 2.5-μg linearized DNA fragment, 62.5 μCi 35S-UTP, 330 μM ATP, 330 μM guanosine 5'-triphosphate, 330 μM cytidine 5'-triphosphate, 10 μM cold UTP, 1× transcription buffer, 66.6 μM dithiothreitol, 40 U RNase Inhibitor, and 20 U of the appropriate RNA polymerase.

Before hybridization, tissue slides were pretreated with 4% paraformaldehyde, washed in KPBS, acetylated, delipidized in chloroform, and dehydrated through a graded ethanol series. Each riboprobe was diluted (1.0 × 10^6 cpm/50 μl) in hybridization buffer (50% formamide, 1× Denhardt’s Solution, 10% dextran sulfate, 200 μg/ml fish sperm single-stranded DNA, 100 μg/ml yeast RNA, and 20 μM dithiothreitol), and applied to a one in 10 series of slides containing the mounted brain sections cut on the cryostat. Slides with hybridization media were then coverslipped, placed in hybridization chambers over blotting paper soaked in 50% formamide, and incubated overnight at 55°C. The next day, coverslips were removed and slides washed in 2× standard saline citrate (SSC). Slides were subsequently incubated in 100 μg/ml RNase A for 30 min at 37°C, washed numerous times in 0.2× SSC, once in 0.2× SSC for 1 h at 65°C, and finally dehydrated through a graded ethanol series.

**Image analysis**

Hybridized slides were exposed on Kodak Biomax MR film (Eastman Kodak Co., Rochester, NY) (5 h for AVP, 7 d for CRH, and 14 d for c-fos). Film images of brain sections were captured by a digital video camera. Anatomical brain regions were identified using Swanson (34), and Paxinos and Watson (31) rat brain atlases. Brain sections were matched for rostrocaudal level between rats for analyses. All brain regions and nuclei were clearly distinguishable by the specificity and intensity of AP, CRH, and c-fos expression patterns. The paraventricular and magnocellular aspects of the PVN were identified using the Nissl stained sections from a parallel series of tissue as a guide. Semiquantitative analyses of autoradiograph images were performed using Scion Image (Scion, Frederick, MD) software, and hybridization signal was expressed as gray level units. The gray level signal of a hybridized tissue region of interest was corrected by subtracting the gray level signal over a nonhybridized area of tissue (white matter, corpus callosum), and expressed as corrected gray level. 35S standards were also measured using Scion Image, and transferred to Assay Zap (Biosoft, Cambridge, UK, and P. L. Taylor) to generate a standard curve to verify that all measured gray levels were in the linear range of the film.

**Statistical analysis**

Data are expressed as mean ± SEM. To consider differences in body weight, adjusted thymus and adrenal gland weights were calculated as organ weight (mg) divided by final body weight (g) and multiplied by 100. AVP, CRH, and c-fos mRNA expression were expressed as corrected gray level. Integrated plasma ACTH and corticosterone responses were calculated as total area under the curve (AUC) for the restraint time course. AUC was calculated using the trapezoidal rule. The trapezoid area under each two adjacent hormonal data points (a and b) are calculated by (a + b)/2 t, where t = time between a and b. The total AUC is calculated by summing all the trapezoid areas under each two adjacent hormonal time points.

Organ weights, pre-acute stress plasma hormones, integrated plasma hormones, and *in situ* hybridization data were analyzed by two-way

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**FIG. 1.** Brain tissue sections were immunolabeled for NeuN for analyses of the extent of lesions of the anteroventral BST, targeting the dm and fu nuclei (BSTdm/fu) of the BST. A, Sham injection sites have intact neurons, minimal gliosis, and minimal needle track damage. B, In contrast, anteroventral BST lesions were identified by damage ventral to the anterior commissure (ac) at approximately AP − 0.15 to −0.3 mm from bregma. Scale bars, 300 μm. C, Schematic diagram illustrates the extent of the spread of damage of the smallest (black shaded area) to the largest lesions (cross-hatched shaded area) confirmed as hits of the anteroventral BST. Image adapted from Swanson (34). B, BSTdm/fu; ac, anterior commissure, olfactory limb; abst, BST anterodorsal area; ad, BST dorsolateral preoptic nucleus; APV, anteroventral periventricular nucleus hypothalamus; FS, fundus of the striatum; GPe, globus pallidus, lateral segment; Gpi, internal capsule; ju, BST juxtacapsular nucleus; LPO, lateral preoptic area; LSB, lateral septal nucleus, ventral part; MA, magnocellular preoptic nucleus; MEC, median eminence; Opf, optic radiation; ov, BST oval nucleus; SI, substantia innominata.
TABLE 1. Body weight after anterioventral BST lesion and chronic stress

<table>
<thead>
<tr>
<th></th>
<th>No CVS</th>
<th></th>
<th>CVS</th>
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<tbody>
<tr>
<td></td>
<td>Sham</td>
<td>Lesion</td>
<td>Sham</td>
<td>Lesion</td>
</tr>
<tr>
<td>d − 6 (Surgery) % body weight gain</td>
<td>100 ± 0.0</td>
<td>100 ± 0.0</td>
<td>100 ± 0.0</td>
<td>100 ± 0.0</td>
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<tr>
<td>d 1 CVS % body weight gain</td>
<td>104 ± 0.6</td>
<td>104 ± 1.3</td>
<td>106 ± 0.5</td>
<td>104 ± 1.0</td>
</tr>
<tr>
<td>d 8 CVS % body weight gain</td>
<td>112 ± 1.3</td>
<td>113 ± 1.9</td>
<td>103 ± 1.1*</td>
<td>105 ± 1.1*</td>
</tr>
<tr>
<td>d 16 (End of CVS) % body weight gain</td>
<td>120 ± 1.1</td>
<td>118 ± 2.3</td>
<td>110 ± 1.0*</td>
<td>109 ± 1.2*</td>
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</table>

Data are shown as mean ± SEM (n = 9–12).
* P < 0.05 vs. respective “No CVS.”

Results

Lesions of the BSTdm/fu nuclei

The role of the dm/fu nuclei of the BST in HPA axis regulation after acute and/or chronic stress exposure was assessed using bilateral ibotenate lesions targeting these nuclei. Lesion location and extent were verified by Nissl stain and NeuN immunolabeling (Fig. 1). Confirmed anterior BST lesions were centered ventral to the anterior commissure at approximately AP −10.0 mm from bregma, and were relatively consistent in the extent of damage. For all the data in this study, the bilateral lesions were confirmed as “hits” when the damage primarily included the fu and dm nuclei of the BST, with limited damage extending into the adjacent subcommissural zone. The rats with lesion damage localized outside of the targeted BST nuclei were not included in the statistical analyses. The majority of damage in the anterior BST nuclei remained ventral to the anterior commissure, minimal ventral damage in the medial preoptic area, and no caudal damage in the posterior division of the BST.

Body and organ weights

There were main effects of CVS (F1,165 = 33.25; P < 0.05) and day (F3,165 = 273.47; P < 0.05), and interactions between CVS x day (F3,165 = 64.16; P < 0.05), and day x day (F3,165 = 3.21; P < 0.05) on body weight (Table 1). However, there was no effect of lesion on body weight gain, no lesion x day interaction, and no lesion x CVS x day interaction, indicating that destruction of the BSTdm/fu is not sufficient to alter the consequences of chronic stress on body weight.

There were main effects of CVS on raw adrenal weights (Table 2) (F1,38 = 18.60; P < 0.05) and adjusted adrenal weights (F1,38 = 71.85; P < 0.05). However, there were no effects of lesion on raw or adjusted adrenal weights, and there was no lesion x CVS interaction. More specifically, CVS groups had greater raw and adjusted adrenal weights (P < 0.05) than their respective non-CVS groups, suggesting that CVS induced adrenal hypertrophy/hyperplasia regardless of lesion.

There was a main effect of CVS on raw thymus weights (Table 2) (F1,38 = 9.73; P < 0.05), but no effect of CVS on thymus weight adjusted for body weight. In addition, there were no effects of lesion on raw or adjusted thymus weights, and there was no lesion x CVS interaction. Overall, the data indicate that CVS decreases thymic weight to the same extent in lesion and control animals.

Pre-acute stress plasma ACTH levels

For pre-acute stress morning plasma ACTH (Fig. 2A), there was a main effect of CVS (F1,38 = 15.46; P < 0.05) and lesion (F1,38 = 7.03; P < 0.05). Post hoc analyses revealed that CVS exposure elevated pre-acute stress plasma ACTH levels only in rats with BST lesions.

Plasma ACTH responses to novel stress

Time-course analysis of ACTH responses to restraint (Fig. 2, B and C) indicated main effects of time (F3,163 = 52.26; P < 0.05) and CVS (F1,163 = 10.04; P < 0.05), and a significant CVS x time interaction (F3,163 = 5.77; P < 0.05). However, there was no main effect of lesion on ACTH secretion, and there were no other interactions. Post hoc analyses indicated that in CVS rats, lesions augmented plasma ACTH levels at 20 min after the onset of restraint (P < 0.05). There was also a main effect of CVS on integrated plasma ACTH response to restraint (Fig. 2D) (F1,36 = 10.33; P < 0.05), but no effect of lesion, and no lesion x CVS interaction. Post hoc analyses revealed that CVS elevated the integrated ACTH levels only in rats with lesions (P < 0.05).

Pre-acute stress plasma corticosterone levels

There was a main effect of CVS on pre-acute stress AM plasma corticosterone levels (Fig. 3A) (F1,38 = 27.22; P < 0.05), but no effect of lesion, and no lesion x CVS interaction. Post hoc analyses indicated that CVS exposure increased pre-acute stress corticosterone levels regardless of lesion (P < 0.05). These data indicate that chronic stress increases resting corticosterone levels to an equal extent in lesion and control groups.
There was a significant effect of CVS (F_{1,167} = 24.23; \textit{P} < 0.05) and time (F_{3,167} = 196.30; \textit{P} < 0.05) on the plasma corticosterone response to a 20-min novel restraint (Fig. 3, B and C), but no main effect of lesion. However, both lesion \times time (F_{3,167} = 4.98; \textit{P} < 0.05) and lesion \times CVS (F_{3,167} = 5.75; \textit{P} < 0.05) interactions were significant, indicative of differential effects of lesions on corticosterone secretion. Post hoc analyses revealed that in rats with lesions, prior CVS exposure elevated plasma corticosterone levels at 20 (\textit{P} < 0.05), 40 (\textit{P} < 0.05), and 60 min (\textit{P} < 0.05) after the onset of restraint relative to rats with lesions and no prior exposure to CVS. In addition, in animals with no prior exposure to CVS, lesions significantly decreased plasma corticosterone response at 60 min after the onset of restraint (\textit{P} < 0.05). There was also a main effect of CVS on integrated plasma corticosterone response to restraint (Fig. 3D) (F_{1,38} = 15.75; \textit{P} < 0.05), but there was no effect of lesion, and no lesion \times CVS interaction. Post hoc analyses indicated that in rats with lesions, prior CVS exposure elevated the integrated corticosterone levels (\textit{P} < 0.05). Together, these data are consistent with previous lesion studies indicating an excitatory role for the anterior BST in acute stress excitation (30) but suggest that this region plays a role in inhibition of acute stress responses after prolonged stress.

### c-fos mRNA expression in the PVN

Induction of a PVN c-fos mRNA response to novel restraint stress was assessed as an indirect indicator of neuronal activation (Fig. 4). There was a main effect of lesion on c-fos mRNA in the PVN (F_{1,37} = 5.94; \textit{P} < 0.05), but no effect of CVS, and there was no lesion \times CVS interaction. Post hoc analysis revealed that in the rats with no prior exposure to CVS, lesions reduced PVN c-fos mRNA expression (\textit{P} < 0.05), as previously documented (30).

### Table 2: Adrenal and thymus weight after anteroventral BST lesion and chronic stress

<table>
<thead>
<tr>
<th></th>
<th>Sham No CVS</th>
<th>Sham Lesion</th>
<th>CVS No CVS</th>
<th>CVS Lesion</th>
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<tr>
<td>Raw adrenal (mg)</td>
<td>51.2 ± 1.2</td>
<td>49.6 ± 1.0</td>
<td>56.9 ± 1.2</td>
<td>56.6 ± 2.2</td>
</tr>
<tr>
<td>Adjusted adrenal (mg/100 g BW)</td>
<td>13.6 ± 0.3</td>
<td>13.6 ± 0.4</td>
<td>17.2 ± 0.3</td>
<td>17.3 ± 0.6</td>
</tr>
<tr>
<td>Raw thymus (mg)</td>
<td>363 ± 17</td>
<td>375 ± 31</td>
<td>314 ± 13</td>
<td>301 ± 17</td>
</tr>
<tr>
<td>Adjusted thymus (mg/100g BW)</td>
<td>98 ± 5.1</td>
<td>102 ± 7.1</td>
<td>96 ± 4.3</td>
<td>92 ± 4.6</td>
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</table>

Data are shown as mean ± SEM (n = 9–12). BW, Body weight. * \textit{P} < 0.05 vs. respective "No CVS." 

**Fig. 2.** A, In sham rats, prior CVS exposure did not affect the plasma ACTH response to 20-min restraint stress. B, In rats with lesions, prior CVS exposure elevated the plasma ACTH response to an acute 20-min restraint at the 20-min time point compared with rats with lesions and no prior exposure to CVS. C, In rats with lesions, CVS significantly elevated pre-acute stress AM plasma ACTH levels. Anteroventral BST lesions potentiated pre-acute stress AM plasma ACTH levels only in the rats exposed to CVS. D, In rats with prior exposure to CVS, anteroventral BST lesions elevated the integrated ACTH response. Data are presented as mean ± SEM (n = 9–12). *, \textit{P} < 0.05 vs. respective "No CVS." #, \textit{P} < 0.05 vs. respective Sham.
CRH and vasopressin mRNA expression in the PVN

CRH mRNA expression in the PVN (Fig. 5) and AVP mRNA expression in the medial parvocellular aspect of the PVN (Fig. 6) were assessed as indicators of central HPA axis tone. There was a significant effect of CVS on CRH mRNA in the PVN ($F_{1,37} = 14.13; P < 0.05$), but there was no effect of lesion, and there was no lesion x CVS interaction. Post hoc analyses revealed that CVS significantly elevated CRH mRNA expression in the PVN regardless of lesion ($P < 0.05$ vs. respective “No CVS.”)

There were no significant effects of lesion or CVS on AVP mRNA expression in the parvocellular PVN, and there was no lesion x CVS interaction. Similarly, there were no differences among experimental groups for magnocellular AVP expression in the posterior magnocellular PVN or in the supraoptic nucleus (data not shown).

Discussion

The current study demonstrates that damage to the anteroventral BST, encompassing the PVN-projecting BSTdm/fu subnuclei, has divergent effects on HPA axis responses to acute vs. chronic stress. Anteroventral BST lesions reduced c-fos activation of the PVN and corticosterone secretion in response to acute restraint, whereas corticosterone responses to novel restraint were potentiated in rats with lesions that had prior CVS exposure. Similarly, pre-acute stress plasma ACTH, integrated ACTH, and corticosterone responses were elevated only in the CVS-exposed rats with anteroventral BST damage. Overall, the data suggest that the BSTdm/fu act as activators of the HPA axis response to acute stress challenges but are also involved in inhibitory regulation of HPA axis reactivity after chronic stress. The data are consistent with a chronic stress-induced functional reassignment of the anterior BST-PVN circuitry, occurring at or above the level of the BST.

This study used a 2-wk exposure to chronic unpredictable stress that elicited consistent physiological and neuroendocrine adaptations. CVS evoked decreased body weight gain, adrenal hypertrophy/hyperplasia, thymic involution, elevated AM resting plasma corticosterone levels, and potentiated CRH mRNA in the PVN as previously shown by our group (9, 35), suggestive of increased HPA tone. None of these endpoints was significantly affected by anterior BST lesions, suggesting that damage to this region does not alter...
the development of the major physiological indices of chronic stress.

The novel acute restraint challenge evoked a significant increase in plasma ACTH levels at 20 min in all groups. Moreover, this response to the restraint was further potentiated by prior CVS exposure only in rats with anteroventral BST lesions, indicating damage to a circuit that would have otherwise constrained the response. Notably, this increase can be attributed at least in part to the markedly lower ACTH response observed in rats with lesions and no prior exposure to CVS, compared with sham rats with no prior exposure to CVS. When analyzing the integrated ACTH response, the AUC was also significantly elevated by CVS only in rats with lesions, suggesting that the anteroventral BST inhibits CVS-induced sensitization of ACTH secretion. In addition, resting AM plasma ACTH levels were also augmented only in the CVS-exposed rats with lesions, suggesting enhanced pre-acute stress activation at the level of the pituitary. Overall, contrary to our a priori hypothesis, the dm and fu nuclei provide inhibitory drive on HPA activity after chronic stress exposure, acting to inhibit both resting AM plasma ACTH secretion and ACTH responses to stress, in contrast to a role in excitation of the HPA axis to an acute stress without prior CVS exposure.

In the present work, the corticosterone response to novel restraint was significantly diminished at 60 min by anteroventral BST lesions in rats with no prior exposure to CVS exposure. This replicates our previous findings indicating that the anteroventral BST stimulates HPA reactivity to an acute stress (30). However, in CVS-exposed rats, lesions of the anteroventral BST potentiated the plasma corticosterone response at 40 min after onset of restraint. CVS also elevated the total integrated plasma corticosterone response and corticosterone levels at 20, 40, and 60 min after novel restraint, but only in rats with lesions of the anteroventral BST. Interestingly, we failed to see a sensitization of the corticosterone response in CVS sham rats, although this is not always observed after CVS or other chronic stress paradigms (9, 36). These data are consistent with the ACTH data and, together, indicate that the anteroventral BST clearly has opposing roles in stress responsivity, likely providing excitation during an acute stress challenge but driving inhibition after adaptations to chronic stress exposure.

At the level of the PVN, c-fos mRNA expression was decreased by lesions of the anteroventral BST in the absence of CVS exposure, further indicating the BSTdm/fu provides activation of the PVN in response to a single acute stressor, consistent with our previous findings (30). In contrast, the lesion-induced decrease in c-fos induction was not observed in rats exposed to CVS. Although there was no main effect of CVS on c-fos mRNA, there was a trend for CVS to decrease c-fos mRNA in the PVN (P = 0.065), similar to previous studies in our group that have reported CVS-induced de-
increases in c-fos mRNA in the PVN at 60 min after onset of restraint stress (9, 35). Overall, we interpret these data to suggest that a prior history of chronic stress potentially reverses the lesion-induced decrease in PVN activation and may be related to the CVS-induced up-regulated hormonal responses to restraint by CVS exposure in rats with lesions only.

In this study other indices of chronic stress were measured after CVS, including increased adrenal weight, suggestive of a history of elevated ACTH exposure (3), and thymic involution, suggestive of a history of increased circulating glucocorticoids. In addition, a rapid decline in body weight gain was observed after onset of CVS, as shown by others (37). Chronic stress is also known to up-regulate CRH and AVP mRNA expression in the PVN. However, none of these physiological or neuroendocrine adaptations to chronic stress was affected by lesions of the anteroventral BST, suggesting that disruption of these BST nuclei does not affect the development of steady-state changes in HPA function engendered by chronic stress.

There are several possible, and perhaps overlapping, mechanisms that may explain the ability of anteroventral BST lesions to inhibit acute responses in chronic stress. We observed CVS-induced increases in CRH mRNA but not AVP mRNA expression in the PVN. However, none of these physiological or neuroendocrine adaptations to chronic stress was affected by lesions of the anteroventral BST, suggesting that disruption of these BST nuclei does not affect the development of steady-state changes in HPA function engendered by chronic stress.

Second, the BSTdm/fu receives heavy inputs from many other known stress-regulatory sites, including abundant noradrenergic inputs from the brainstem (40, 42) and limbic input from the central nucleus of the amygdala and the infralimbic cortex (12, 18). It is yet to be determined whether specific circuits from different brainstem and limbic regions converge on the same type of HPA-projecting neurons, but the dissociation between chronic and acute stress regulation by the BSTdm/fu nuclei suggests that there may be separate pools of neurons within these upstream nuclei that provide very different information or feedback about stressful stimuli.

Finally, it is possible that removal of BSTdm/fu can elicit downstream neuroplastic changes favoring PVN excitation. Thus, loss of BSTdm/fu input to the PVN or PVN-projecting structures (such as the nucleus of the solitary tract) may permit a compensatory enhancement of inputs that are preferentially sensitive to the effects of chronic stress.

In conclusion, the current study documents that the dm/fu nuclei of the BST differentially regulate HPA responses by activating acute stress responses while attenuating HPA responses after chronic unpredictable stress adaptations. We infer that the anteroventral BST contains heterogeneous populations of neurons that differentially regulate the capacity or efficiency of HPA activation, dependent on prior experience. Given that lesions of this region do not affect steady-state changes in HPA function after stress, it is likely that the anteroventral BST is involved in stress plasticity. Inappropriate limbic processing of stressful information is implicated in numerous chronic stress-related disease states (43, 44) and may involve functional changes in circuitry funneling through these important integrative BST nuclei.

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