A Group of Descending Glutamatergic Neurons Activated by Stress in Corticolimbic Regions Project to the Nucleus Accumbens

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The nucleus accumbens (NAc) is the major component of the ventral striatum that regulates stress-induced depression. The NAc receives dopaminergic inputs from the ventral tegmental area (VTA), and the role of VTA-NAc neurons in stress response has been recently characterized. The NAc also receives glutamatergic inputs from various forebrain structures including the prelimbic cortex (PL), basolateral amygdala (BLA), and ventral hippocampus (vHIP), whereas the role of those glutamatergic afferents in stress response remains underscored. In the present study, we investigated the extent to which descending glutamatergic neurons activated by stress in the PL, BLA, and vHIP project to the NAc. To specifically label the input neurons into the NAc, fluorescent-tagged cholera toxin subunit B (CTB), which can be used as a retrograde neuronal tracer, was injected into the NAc. After two weeks, the mice were placed under restraint for 1 h. Subsequent histological analyses indicated that CTB-positive cells were detected in 170~680 cells/mm² in the PL, BLA, and vHIP, and those CTB-positive cells were mostly glutamatergic. In the PL, BLA, and vHIP regions analyzed, stress-induced c-Fos expression was found in 20~100 cells/mm². Among the CTB-positive cells, 2.6% in the PL, 4.2% in the BLA, and 1.1% in the vHIP were co-labeled by c-Fos, whereas among c-Fos-positive cells, 7.7% in the PL, 19.8% in the BLA, and 8.5% in the vHIP were co-labeled with CTB. These results suggest that the NAc receives a significant but differing proportion of glutamatergic inputs from the PL, BLA, and vHIP in stress response.

Key words: Stress, c-Fos, Retrograde tracer, Nucleus accumbens

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been characterized [1, 2, 10-12]. Chronic social defeat stress increased the firing rate of VTA-to-NAc dopaminergic neurons [13, 14]. Optogenetic phasic stimulation of VTA-to-NAc neurons, but not VTA-to-medial prefrontal cortex (mPFC) neurons, facilitated depressive-like behaviors. Conversely, optogenetically inhibiting the VTA-to-NAc circuit was anti-depressive, whereas inhibiting the VTA-to-mPFC projection produced depressive effects [15]. Stimulating mPFC-to-NAc neurons facilitated anti-depressive behaviors [16], whereas activating hippocampal or thalamic inputs [16], and ventral hippocampal inputs [17] to the NAc promoted depressive-like behaviors. Stimulating BLA-to-NAc glutamatergic neurons promoted cue-evoked sucrose intake, which required the activation of an intra-NAc dopamine D1 receptor signaling [18]. Glutamatergic synaptic strength in the NAc shell medium spiny neurons (MSNs) was increased after stress [19]. Overall, these studies suggest that the NAc receives not only dopaminergic VTA neurons, but also glutamatergic inputs from corticolimbic regions. However, the role of those glutamatergic neurons in regulating stress and depressive behaviors has only recently received attention [16, 17, 20-22].

Stimulation dependent c-Fos expression has been used to map brain regions that are activated under specific conditions [23-28]. For example, stress- or exercise-dependent c-Fos induction has been used to identify and characterize brain regions that are activated by stress [11, 29-31] or exercise [32-35]. Recently, functional labeling of specific neurons with a c-Fos promoter combined with sophisticated optogenetics or pharmacogenetics methods makes it possible to pinpoint the functional networks that mediate context-dependent behavioral changes in cognition and emotion [36-38].

In the present study, we investigated the extent to which descending glutamatergic neurons activated by stress in the mPFC, vHIP, and BLA innervate the NAc by using a combined method of staining neurons with activity-dependent c-Fos expression and a fluorescent retrograde marker, Alexa-conjugated cholera toxin subunit B (CTB). We identified a group of glutamatergic neurons in the mPFC, vHIP, and BLA that projected into the NAc and were activated by acute stress.

MATERIALS AND METHODS

Animals

Seven-week-old male C57BL/6J mice were purchased from a local supplier (Daehan Biolink Inc., Chungbuk, Korea). Upon arriving, they were grouped according to an experimental design and housed in pairs in regular plastic cages in a temperature- (23°C) and humidity-controlled (50–60%) room with a 12/12-hour light/dark cycle (light on at 7 A.M.). Mice were allowed access to food and water ad libitum in home cage, and were acclimated to the laboratory environment for 1 week before stereotaxic surgery. All procedures followed the animal care guidelines of Ewha Womans University School of Medicine (IACUC 16-018).

Stereotaxic injection

Stereotaxic surgery was performed as described previously [39, 40] with minor modifications. In brief, mice were anesthetized by intraperitoneal (i.p.) injection of 3.5:1 mixture of ketamine (50 mg/ml) and xylazine hydrochloride (23.3 mg/ml) at 2 μl/g body weight. After 5 min, the mouse was staged on the surgical apparatus (Stoelting Co., Wood Dale, IL, USA). Cholera toxin subunit B (CTB) conjugated with Alexa Fluor® 488 (#C34755; Molecular Probes, USA) was delivered at 0.5 μl/min in 0.4 μl volume into each of both NAc regions (AP, +1.4 mm; ML, ±1.3 mm; DV, -5.0 mm from bregma) with a 30 G needle (Top Medical, Korea). Tygon® microbore tubing (#AAD04091; Saint-Gobain, USA), and a Hamilton microsyringe (#CAL80030; Hamilton, USA). CTB solution (0.1%, w/v) was prepared by dissolving CTB in sterile PBS (pH 7.28). After the injection finished, the needle was left in place for 3 min and removed with three intermediate steps over 3 min to minimize backflow. After the skin was sewn, mice were kept on a warm pad (30°C) until they awoke. To ensure retrograde transport, CTB-injected mice were kept for two weeks in their home cage before starting the experimental schedule.

Restraint stress

Mice were restrained as described previously [39-41]. Mice were individually placed in a 50-ml conical tube (#50050; SPL Life Science, Korea) that had many ventilation holes and were pushed toward the conical end. Tail side was plugged with a 3-cm piece from the middle of a 15-ml tube (#50015; SPL Life Science). The tube was closed with a cap that had a hole for the mouse’s tail. Tubes were placed in home cage for 1 h. Mice were restrained beginning at 10 A.M.

Immunohistochemistry

Immunohistochemistry was carried out as described previously [39, 40]. Mice were anesthetized with 2.5% (w/v) avertin (#T48402; Aldrich, USA) at 20 μl/g body weight, perfused with saline, and fixed with 4% (w/v) paraformaldehyde solution (#P6148; Sigma-Aldrich, USA). The fixed brain was cut into 40-μm-thick coronal sections using a vibratome (#VT1000S; Leica, German), and humidity-controlled (30 % glycerol) at -20°C. Brain sections at the levels of the PL (AP, +1.9 mm), NAc (AP, +1.3 mm), BLA (AP, -1.3 mm), and vHIP (AP, -3.1 mm) were used.
Fig. 1. Retrograde labeling of neurons projecting into the NAc. (A) Experimental design and time schedule for CTB injection (green arrows) into the NAc. After 14 days of the surgery, 1-h restraint was treated. Red arrow, time point for tissue preparation. CON, CTB-injected control mice; RST, CTB-injected and stress treated mice. (B, C) A fluorescence image (B) of Alexa-conjugated CTB (cholera toxin subunit B) expression in the lateral core and shell of the NAc 14 days after the surgery and a schematic diagram (C) of CTB injection (left panel). Projections from the prelimbic cortex (PL), ventral hippocampus (vHIP), ventral tegmental area (VTA), and basolateral amygdala (BLA) are indicated. acc, anterior commissures; core, NAc core; lSh, lateral shell of the NAc; mSh, medial shell of the NAc. Scale bar, 500 μm. (D) Schematic diagram illustrating endocytosis of CTB at the synaptic terminal and retrograde transport of fluorescent CTB along the axon and its translocation into the cell body region (adapted from [63, 64]). (E–G) CTB-positive cells in the PL (E), BLA (F), and vHIP (G). Schematic diagrams illustrating the coronal sections at the level of the PL (E), BLA (F), and vHIP (G). High magnification of the areas marked with a rectangle. Cg, cingulate cortex; fmi, anterior forceps minor of the corpus callosum; VI/V, cortical layers VI/V; II/III, cortical layers II/III; BLA, basolateral amygdala; CeA, central amygdala. Scale bars; 50 μm in the PL and BLA, and vHIP.
for immunohistochemical analysis. The sections were washed with PBS three times, blocked with 5% BSA in PBST (PBS containing 0.1% Triton X-100) for 1 h at 4°C, and incubated with monoclonal anti-glutamate antibody derived from the GLU-4 hybridoma clone (1:500; #G-9282; Sigma, USA) or c-Fos antibody (1:1000; #sc-52; Santa Cruz Biotechnology, USA) in 5% BSA blocking solution overnight. After washing, the sections were incubated with secondary antibody, DyLight® 549 horse anti-rabbit IgG (1:200; #DI-1094; Vector Laboratories, USA), for 2 h at room temperature. After washing, stained sections were mounted on a gelatin-coated glass slide with mounting solution (#S3023; Dako, Japan) and a cover slide.

Stained sections were examined with a fluorescence microscope (#BX51; Olympus, Japan) and filters specific for Alexa Fluor™ 488 (#39002; Chroma Technology, USA) and DyLight® 594 (#39010; Chroma Technology, USA). Image analysis was performed with MetaMorph® software (Molecular Devices, USA).

Statistical analysis

Two-sample comparisons were performed with Student’s t-test. PRISM software 6 (GraphPad Software, USA) was used for statistical analyses. All data are presented as the mean±SEM, and statistical differences were accepted at the 5% level.

RESULTS

To specifically label a group of neurons in the limbic system that projected into the NAc, a fluorescent retrograde neuronal tracer, Alexa-conjugated cholera toxin subunit B (CTB) [42-44], was stereotaxically injected into the NAc lateral core and shell on both

Fig. 2. Most of CTB-positive cells in PL, BLA, and vHIP regions were stained by GLU-4, a glutamatergic neuron marker. (A–I) Schematic diagrams illustrating the coronal section of the PL (A), BLA (D), and vHIP (G), and the areas (filled blue area) analyzed for cells labeled by fluorescent-CTB (green) and anti-glutamate antibody (GLU-4, red). Fluorescence images showing cells labeled with CTB (green), or GLU-4 (red), and high magnification of an overlay (boxed areas, merged) in the PL (B), BLA (E), and vHIP (H). Quantification of CTB labeled cells, GLU-4 labeled cells, and CTB plus GLU-4 double-labeled cells in the PL (C), BLA (F), and vHIP (I). Venn diagrams showing the summed numbers of cells stained with CTB only (green) or CTB+GLU-4 (yellow) counted in the sections of PL (C), BLA (F), and vHIP (I) regions. Bar graphs showing the normalized mean number in mm² of cells stained with CTB only (green) and double-labeled with CTB+GLU-4 (yellow) in the PL (C), BLA (F), and vHIP (I). n=4 animals for each region. *indicates cells stained with CTB or GLU-4. Arrow heads indicate cells stained with CTB and GLU-4. Scale bars; 50 μm in the PL and BLA, and 100 μm in the vHIP. All data are mean±SEM. Student’s t-test.
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sides (Fig. 1A, B). After surgery, mice were placed in their home cages for 2 weeks, allowing that the fluorescent tagged CTB could be transported to the cell body of neurons in the PL, BLA, and vHIP (Fig. 1C). Then, CTB-injected mice were then restrained for 1 h (Fig. 1A). Immediately after restraint, mice were sacrificed, and brains were histologically examined. CTB-injected unrestrained mice were used as a control. We previously demonstrated that restraint for 1 h induced significant c-Fos levels in most forebrain regions without saturation [11]. We confirmed that fluorescent-labeled CTB was detected in various brain regions including the PL, BLA, and vHIP (Fig. 1D–G).

Next, we examined whether CTB-labeled neurons in the PL, BLA, and vHIP are glutamatergic using monoclonal anti-glutamate antibody (GLU-4 for short), which has been used for labeling glutamatergic neurons [45, 46]. In the PL region examined (AP, 1.98±0 mm), 89.8% of CTB-labeled neurons cells (743/829) were stained by GLU-4 antibody, a specific marker for labeling glutamatergic neurons, whereas 10.2% (86/829) were not labeled by GLU-4 (Fig. 2A–C). In the BLA region examined (AP, 1.31±0.03 mm), 93.0% of CTB-labeled neurons (610/657) were positive to GLU-4, whereas 7.0% (47/657) were not labeled by GLU-4 antibody (Fig. 2D–F). In the vHIP region examined (AP, -3.19±0.03 mm), 90.5% of CTB-labeled neurons (883/980) were positive to GLU-4, whereas 9.5% (97/980) were not stained by GLU-4 (Fig. 2G–I).

Further analyses indicated that in the PL region of control mice, CTB-positive cells were counted in 175±54 cells/mm², and c-Fos-

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**Fig. 3.** Analyses of cells stained by the stimulation-dependent neural activity marker c-Fos and the retrograde tracer CTB in PL, BLA, and vHIP neurons. (A–I) Schematic diagrams illustrating the coronal section of the PL (A), BLA (D), and vHIP (G), the areas (filled blue area) analyzed for cells stained by c-Fos and/or fluorescent CTB, and high magnification (boxed areas). Fluorescence images showing cells labeled with CTB (green) or c-Fos (red), and high magnification of an overlay of CTB and c-Fos labeled cells (boxed area) in the PL (B), BLA (E), and vHIP (H). Arrow heads indicate CTB-positive cells and arrows indicate c-Fos-positive cells. Scale bars: Scale bars, 50 μm in the PL and BLA, and 100 μm in the vHIP. The examined sections of the PL were at the level of 1.95±0.01 mm (AP) from the bregma, those of the BLA were at -1.39±0.03 mm, and those of the vHIP were at -3.22±0.04 mm. The numbers of animals examined were n=3–4 animals for the PL, n=4–5 animals for the BLA, and n=3 animals each for the vHIP. Quantification of CTB labeled cells, c-Fos labeled cells, and CTB and c-Fos double-labeled cells in the PL (C), BLA (F), and vHIP (I) of control (CON) and restrained (RST) mice. All data are mean±SEM. *p<0.05, **p<0.01 (Student’s t-test).

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positive cells were detected in 1±0 cells/mm² (Fig. 3A–C; CON). In contrast, in the PL region of mice treated with restraint, CTB-positive cells were found in 167±36 cells/mm², and c-Fos-positive cells were counted in 58±15 cells/mm² (Fig. 3A–C; RST). Double labeling analysis indicated that among the CTB-positive cells in the PL, 2.7% (23/837) were co-labeled by stress-induced c-Fos, whereas of the stress-induced c-Fos-positive cells, 8.0% (23/289) were co-labeled by CTB (Fig. 4).

In the BLA region of control mice, CTB-positive cells were detected in 428±91 cells/mm², and c-Fos-positive cells were found in 3±1 cells/mm² (Fig. 3D–F; CON). Whereas in the BLA region of mice treated with restraint, CTB-labeled cells were counted in 488±74 cells/mm², and c-Fos-positive cells were found in 98±20 cells/mm² (Fig. 3D–F). Double labeling analysis indicated that among the CTB-positive cells in the BLA, 4.0% (59/1464) were co-labeled by stress-induced c-Fos, whereas of the stress-induced c-Fos-positive cells, 20.1% (59/294) were co-labeled by CTB (Fig. 4).

In the vHIP region of control mice, CTB-positive cells were found in 684±105 cells/mm², and c-Fos-positive cells were detected in 0±0 cells/mm² (Fig. 3G–I; CON). In contrast, in the vHIP region of mice treated with restraint, CTB-labeled cells were counted in 608±32 cells/mm², and c-Fos-positive cells were detected in 105±21 cells/mm² (Fig. 3G–I). Double labeling analysis indicated that among the CTB-positive cells in the vHIP, 1.1% (8/726) were co-labeled by stress-induced c-Fos, whereas of the stress-induced c-Fos-positive cells in the vHIP, 6.3% (8/128) were co-labeled by CTB (Fig. 4).

Overall, these results suggest that a group of glutamatergic neurons in the PL, BLA, and vHIP were projecting to the NAc, among which 2.7% (23/837) in the PL, 4.0% (59/1464) in the BLA and 1.1% (8/726) in the vHIP were activated by stress. Among the c-Fos-positive cells counted, 8.0% (23/289) of neurons in the PL, 20.1% (59/294) of neurons in the BLA, and 6.3% (8/128) of neurons in the vHIP projected to the NAc (Fig. 4).

**DISCUSSION**

The NAc is the core of the ventral striatum that integrates various excitatory inputs from the medial prefrontal cortex (mPFC), vHIP and amygdala (Amy), and it regulates emotionally motivated behaviors and action-related behaviors [2, 7]. The NAc receives not only dopaminergic inputs, but also glutamatergic inputs from forebrain structures [5, 6]. Our results also indicate that the NAc region receives significant levels of glutamatergic inputs from the PL, BLA, and vHIP. Approximately 167±36 cells/mm² in the PL, 488±74 cells/mm² in the BLA, and 608±32 cells/mm² in the vHIP were labeled by fluorescent CTB. However, among the CTB-positive cells, only 2.7% of cells in the PL, 4.0% of cells in the BLA, and 1.1% of cells in the vHIP were co-labeled with c-Fos. These results suggesting that most NAc-projecting cells in the PL, BLA, and vHIP project to the NAc. In contrast, most cells in the PL, BLA, and vHIP that express c-Fos were not co-labeled by CTB.
vHIP appear to be silent or not directly related to stress response, whereas there are a group of glutamatergic neurons projecting to the NAc in the PL, BLA, and vHIP that are activated by stress.

On the other hand, concerning the c-Fos-positive cells, 8.0% of cells in the PL, 6.3% of cells in the vHIP, and 20.1% of cells in the BLA were co-labeled by CTB. Thus, among the stress-activated cells, a relatively high portion of cells in the PL, vHIP, and BLA are projecting to the NAc. In particular, 20.1% of c-Fos-positive neurons in the BLA projected into the NAc, raising the possibility that there might be a possible role of BLA-to-NAc neurons in regulating stress responses. Optogenetic stimulation of glutamatergic BLA-to-NAc neurons suppressed reward seeking behaviors, which required intra-NAc dopamine D1 receptor signaling [18]. Following repeated restraint, the vHIP was inappropriately activated, which indirectly reduced the neural activity of the BLA-to-NAc pathway [47]. Optogenetic stimulation of vHIP-to-NAc projections promoted depressive-like behaviors [48], whereas optogenetically stimulating mPFC-to-NAc neurons facilitated anti-depressive behaviors [16]. Collectively, these results suggest that the NAc receives, and thereby is regulated by, various descending glutamatergic neurons in the PL, vHIP, and BLA. Of the descending glutamatergic neurons projecting to the NAc, a relatively small group of neurons are activated in stress responses, but their relative percentages among the stress-activated cells were 8.0% in the PL, 6.3% in the vHIP, and 20.1% in the BLA, and thus they are not low.

Mice subjected to chronic social defeat stress had an increased firing rate of VTA-to-NAc dopaminergic neurons, and this change was responsible for depressive behavior [13, 14]. Consistently, optogenetic phasic stimulation of VTA-to-NAc neurons, but not VTA-to-mPFC neurons, facilitated depressive behaviors [15]. Mice treated with chronic, unpredictable, mild stress (CUMS) or chronic restraint stress (CRST) exhibited depressive behaviors [41, 49-52], but mice treated with CUMS or CRST had reduced excitability of VTA-to-NAc neurons [53-55]. These studies suggest that the activities of VTA dopaminergic neurons and the NAc are regulated differently in different stress models, even though they all result in depressive behavior. Concerning this complexity, it will be worthy to test whether descending glutamatergic inputs to the NAc and their downstream neural circuits differ depending on the stress model for the expression of depressive behaviors.

Our results indicate that a group of glutamatergic neurons activated by stress in the PL, vHIP, and BLA project into the NAc. However, this result needs to be interpreted in line with the following points. CTB-labeled NAc-projecting neurons in the PL, BLA, and vHIP are likely excitatory glutamatergic neurons [5, 56]. In fact, most of CTB-labeled neurons were positive to the staining with GLU-4 antibody (Fig. 2). However, 10.2% of the CTB-labeled neurons in the PL, 7.0% in the BLA, and 9.5% in the vHIP were not stained by GLU-4 antibody (Fig. 2). It is possible that those GLU-4-negative cells are long-range GABA neurons [57, 58]. However, considering that many neurons in the brain previously thought to release glutamate as neurotransmitter also release the inhibitory neurotransmitter GABA [59], more sophisticated tools, such as a genetic marker for GABAergic neurons, might be useful to precisely define the identity of those cells. The CTB-viral vector injection labeled the lateral core and shell on both sides of the NAc (Fig. 1), but it could not label all NAc neurons nor label NAc neurons at the same efficacy. Projections to more medial parts of the NAc were not effectively labeled in the present study. Nonetheless, our results present the relative proportion of PL, vHIP, and BLA non-dopaminergic neurons that projected to the injected sites of the NAc and how they were activated by stress.

Our results might be used for the following type of experiments. A transgenic strategy carrying an optogenetic tool [60] or a viral vector-mediated Gq-DREADD (designer receptor exclusively activated by the designer drug: hM3Dq) system [61], when combined with activity-dependent labeling of neurons with specific tools, such as cre-recombinase or transcription factors [62], might be used to investigate whether the cell groups activated by stress in the PL, vHIP, or BLA have a modulatory role in stress-dependent behavioral changes or depressive behaviors.

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