A single brain-derived neurotrophic factor infusion into the dorsomedial prefrontal cortex attenuates cocaine self-administration-induced phosphorylation of synapsin in the nucleus accumbens during early withdrawal

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RESEARCH ARTICLE

A Single Brain-Derived Neurotrophic Factor Infusion into the Dorsomedial Prefrontal Cortex Attenuates Cocaine Self-Administration-Induced Phosphorylation of Synapsin in the Nucleus Accumbens during Early Withdrawal

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Abstract

Background: Dysregulation in the prefrontal cortex-nucleus accumbens pathway has been implicated in cocaine addiction. We have previously demonstrated that one intra-dorsomedial prefrontal cortex brain-derived neurotrophic factor (BDNF) infusion immediately following the last cocaine self-administration session caused a long-lasting inhibition of cocaine-seeking and normalized the cocaine-induced disturbance of glutamate transmission in the nucleus accumbens after extinction and a cocaine prime. However, the molecular mechanism mediating the brain-derived neurotrophic factor effect on cocaine-induced alterations in extracellular glutamate levels is unknown.

Methods: In the present study, we determined the effects of brain-derived neurotrophic factor on cocaine-induced changes in the phosphorylation of synapsin (p-synapsin), a family of presynaptic proteins that mediate synaptic vesicle mobilization, in the nucleus accumbens during early withdrawal.

Results: Two hours after cocaine self-administration, p-synapsin Ser9 and p-synapsin Ser62/67, but not p-synapsin Ser603, were increased in the nucleus accumbens. At 22 hours, only p-synapsin Ser9 was still elevated. Elevations at both time points were attenuated by an intra-dorsomedial prefrontal cortex brain-derived neurotrophic factor infusion immediately after the end of cocaine self-administration. Brain-derived neurotrophic factor also reduced cocaine self-administration withdrawal-induced phosphorylation of the protein phosphatase 2A C-subunit, suggesting that brain-derived neurotrophic factor inhibits protein phosphatase 2A C-subunit, consistent with p-synapsin Ser9 dephosphorylation. Further, co-immunoprecipitation demonstrated that protein phosphatase 2A C-subunit and synapsin are associated in a protein-protein complex that was reduced after 2 hours of withdrawal from cocaine self-administration and reversed by brain-derived neurotrophic factor.

Conclusions: Taken together, these findings demonstrate that brain-derived neurotrophic factor normalizes the cocaine self-administration–induced elevation of p-synapsin in nucleus accumbens that may underlie a disturbance in the probability of...
neurotransmitter release or represent a compensatory neuroadaptation in response to the hypofunction within the prefrontal cortex-nucleus accumbens pathway during cocaine withdrawal.

**Keywords:** cocaine self-administration, immunoblotting, nucleus accumbens, phosphatase, synapsin

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**Introduction**

Susceptibility to drug relapse and other addictive behaviors is thought to depend on long-term neuroadaptations in the mesocortico–basal ganglia circuitry (Koob et al., 1997; Wang and McGinty, 1997; White and Kalivas, 2001). Within this network, attention has focused on the prefrontal cortex (PFC) projection to the nucleus accumbens (NAC), because disruption of glutamate neurotransmission in this pathway by repeated cocaine exposure triggers relapse to drug-seeking (McFarland et al., 2003). Brain-derived neurotrophic factor (BDNF) is expressed in PFC-NAC neurons wherein it regulates presynaptic glutamate release. We have reported that a single BDNF infusion into dorsomedial (dm) PFC immediately following the final session of cocaine self-administration (SA) restores normal extracellular levels of glutamate in the NAC after cocaine SA and suppresses persistent cocaine-seeking (Berglind et al., 2007, 2009). In searching for potential mechanisms that underlie the ability of BDNF in the PFC to normalize glutamate transmission in the NAC, our attention became focused on synapsins for the following reasons. First, synapsins are members of a family of BDNF-regulated presynaptic proteins that control neurotransmitter exocytosis by reversibly tethering synaptic vesicles to each other and to the actin cytoskeleton in a phosphorylation-dependent manner (Benfenati et al., 1985; Jovanovic et al., 1996). Second, after stimulation, the facilitatory effect of BDNF on glutamate release is attenuated in mice lacking synapsins I and II (Jovanovic et al., 2000). Third, BDNF increases the number of docked synaptic vesicles at the active zone of excitatory synapses, facilitating release probability (Tyler and Pozzo-Miller, 2001).

Synapsin I(a,b), II(a,b), and III regulate trafficking of synaptic vesicles between the reserve pool and the active pool in terminals that release different transmitters (Gitler et al., 2004a). Synapsin I has been associated with regulating synaptic vesicles trafficking in GABAergic neurons (Baldelli et al., 2007), whereas synapsin II has been associated with synaptic vesicles trafficking in glutamatergic terminals (Gitler et al., 2004b, 2008). Synapsin III has been associated with synaptic vesicles trafficking in dopamine terminals in the striatum (Kile et al., 2010). Further, in mice lacking all 3 synapsins, the ability to mobilize the reserve pool of synaptic vesicles and release dopamine from striatal terminals was diminished following a cocaine challenge (Venton et al., 2006). In synapsin I-II mutant mice, reductions in vGlut1, vGlut2, and glutamate reuptake were observed (Bogen et al., 2006, 2009), and in vGlut knockout mice, there was a decrease of synapsin I expression and fewer synaptic vesicles in several brain regions (Fremaux et al., 2004). Mice with decreased synapsin I and II expression are prone to epileptiform seizures due to depletion of the reserve synaptic vesicle pool and enhanced docking in the readily releasable pool of excitatory neurons (Rossahl et al., 1995). For these reasons and because of similar molecular weights of synapsins Ia, Ib, and Ila that make it difficult to discriminate them consistently by immunoblotting, we focused attention only on cocaine-induced changes in phosphorylation of synapsin I and Ila, collectively termed here p-synapsin.

Unphosphorylated synapsins bind to and tether small synaptic vesicles to the actin cytoskeleton in the preterminal reserve pool, whereas phosphorylation of synapsins at specific sites causes release of synaptic vesicles, enabling them to traffic from the reserve pool to the readily releasable pool, increasing the probability of action potential-dependent neurotransmitter release. For example, high-frequency stimulation that evokes calcium-dependent glutamate release is associated with protein kinase A/calcium/calmodulin-dependent kinase (CAMK-mediated) phosphorylation of Ser9 and CaMKII-mediated phosphorylation of Ser603 (Chi et al., 2001; Hilfiker et al., 1999, 2005). At the termination of exocytotic release, protein phosphatase 2A (PP2A)-mediated dephosphorylation of CAMKII sites restores synapsin binding to synaptic vesicles and tethering to the cytoskeleton in the reserve pool. In contrast, there is an inverse relationship between Ca++ influx and synapsin phosphorylation at Ser62/67 by extracellular signal-regulated kinase mitogen-activated protein (ERK MAP) kinase. High-frequency stimulation promotes transient ERK-mediated phosphorylation of synapsin at Ser62/67 sites that inhibits posttetanic potentiation and is terminated by PP2B (calcineurin)-dependent dephosphorylation (Jovanovic et al., 2001; Vara et al., 2009). However, low-frequency stimulation can promote phosphorylation of synapsin Ser62/67 (and Ser 9), facilitating dissociation of synaptic vesicles from the actin cytoskeleton (Chi et al., 2003). In addition, BDNF stimulates ERK/ERK-MAPK-mediated synapsin I Ser62/67 phosphorylation that augments glutamate release in cortical synaptosomes in a calcium-independent manner (Jovanovic et al., 2000).

Based on this background, we first evaluated whether cocaine, in the presence or absence of intra-dmPFC BDNF infusion at the end of SA, affects the phosphorylation of synapsins at the 3 major phosphorylation sites during early withdrawal (2 or 22 hours after the end of the last session of cocaine SA). These are the time points at which a normalizing effect of BDNF on cocaine SA-induced changes in phosphorylation is detected in both PFC and NAC (Berglind et al., 2007; Whitfield et al., 2011). Further, after determining that phosphorylation was altered by cocaine SA, we explored the possible association of PP2A with synapsin after cocaine SA and/or intra-dmPFC BDNF infusion.

**Methods**

**Animals**

Adult male Sprague-Dawley rats (Charles River, MA) weighing 301 to 325g at the beginning of each experiment were housed individually in a temperature- and humidity-controlled vivarium on a reversed 12-h–light–dark cycle (lights off at 7:00 AM). Animals were given 20 to 25g of standard rat chow (Harlan) with ad libitum access to water daily. Experimental procedures were performed during the dark cycle. All experimental protocols were approved by the Institutional Animal Care and Use Committee of the Medical University of South Carolina and complied with the National Institute of Health Guide for the Care and Use of Laboratory Animals (NIH publication no. 80–23, revised 1996). All efforts were made to minimize animal suffering and reduce the number of animals used.
Surgery

Animals were given at least a 5-day acclimation period. On the day of surgery, rats were anesthetized with a mixture of ketamine (66 mg/kg, intraperitoneally; Vedco Inc) and xylazine (1.33 mg/kg, intraperitoneally; Lloyd Laboratories), respectively, followed by equithesin (0.5 ml/kg) and ketorolac (2 mg/kg; Sigma-Aldrich). A silastic catheter was implanted into the right jugular vein with the distal end threaded subcutaneously and exiting on the back to connect to an infusion harness (Instech Solomon) for intravenous infusions. Immediately after catheterization, rats were implanted with 28-gauge bilateral stainless-steel guide cannulae (Plastics One) aimed 1 mm above the dmPFC infusion site. Coordinates used in the stereotaxic procedures were +3.0 mm anterior-posterior (AP), ±0.6 mm mediolateral, and −1.6 mm dorsal-ventral from the dural surface (Paxinos and Watson, 2005). Guide cannulae were secured to the skull with dental cement and steel screws. Two 10-mm stylets (Plastics One) were placed into the guide cannulae following surgery to prevent blockage throughout the experiment. After surgeries, daily intravenous infusions of 0.1 mL of the antibiotics, Cefazolin (10 mg/mL; Schein Pharmaceuticals) or Timentin (2.4 mg/mL), and 0.1 mL of heparinized saline (100 U/mL; Elkins-Sinn) were administered during surgical recovery and throughout SA to maintain catheter patency. Catheter patency was verified as needed with an intravenous infusion of 0.1 mL of methohexital sodium (10 mg/mL; Eli Lilly), which produces a rapid loss of muscle tone.

Cocaine SA and Intracranial Infusion

After 5 days of recovery, rats were food restricted to 90% of their body weight and subjected to 2-hour daily cocaine or yoked-saline SA sessions under a FR1 reinforcement schedule in standard operant conditioning chambers (MED Associates) for 14 days after they reached 10 infusions per session for 3 consecutive days. Each chamber was equipped with 2 retractable levers (7 cm above floor) and a circular stimulus light above each lever. Tygon infusion lines within the spring leash were mounted to a balanced metal arm and attached to a liquid swivel (Instech Solomon). Before each session, the catheter was secured to the infusion line. Presses on the active lever resulted in a 2-second cocaine hydrochloride (0.2 mg/50 µL/infusion; NIDA) infusion via a computer-controlled pump. Cocaine infusions were paired with a 5-second illumination of a white stimulus light above the active lever and a tone (2 kHz, 15 dB above ambient noise) followed by a 20-second timeout period, during which additional responses on the active lever resulted in no programmed consequences. Responses on the inactive lever were recorded but had no consequence. For each session, rats had to self-administer at least 10 infusions to fulfill the maintenance criterion. Immediately after the end of the last SA session, BDNF (0.75 µg/side; R&D Systems) or its vehicle, phosphate buffered saline (pH=7.4), was infused into the dmPFC. Each bilateral infusion was administered through an injector (33 gauge; Plastics One) inserted into the guide cannula so that 1 mm of the injector length extended beyond the guide cannula. After a 2-minute infusion (0.25 µL/min), the injectors remained in the cannulae for 1 minute to allow complete diffusion. Rats were then directly placed into their home cages.

Immunoblotting

Two or 22 hours after infusions, rats were rapidly decapitated and brains were removed and placed in a brain matrix (Braintree Scientific). Two-mm coronal slices at AP 2.7 to 0.7 rostral to Bregma were cut, and the NAc was bilaterally punched and frozen at −80°C. Coronal slices were also cut through the PFC between AP 2.7 and 3.7 mm rostral to Bregma to visually identify cannula placements in the dmPFC (all placements were verified to be in anterior cingulate/prelimbic cortex) before freezing and immunoblotting of other proteins (Sun et al., 2013). Protein from NAc samples was extracted by sonication in ice-cold RIPA lysis buffer and centrifuged at 10,000 × g for 20 minutes at 4°C. The resulting supernatants were collected and subjected to protein analysis with the Micro-Bicinchoninic Acid assay kit (Pierce, Rockford, IL). Equal amounts of protein extracts were run on SDS-PAGE gels and then transferred to PVDF membranes. Membranes were then blocked with 5% nonfat dry milk in Tris-buffered saline with Tween-20 (TBST; pH = 7.6) for 1 hour at room temperature and incubated with primary antibody against p-synapsin Ser9 recognizing all synapsin isoforms at the Ser9 site (1:2000; Cell Signaling), p-synapsin I Ser62/67 (1:1000), p-synapsin II Ser603 (1:3000; Millipore), or p-PP2Ac Tyr307 (1:5000; Abcam) overnight at 4°C. After 3 washes with TBST, membranes were incubated with their appropriate secondary antibodies for 1 hour at room temperature followed by 3 more washes with TBST. Antibody binding was detected by using an enhanced chemiluminescence kit (ECL Plus; GE Healthcare Bio-Sciences). Membranes were then stripped and reprobed for total protein by antisera against synapsin (1:2000; Cell Signaling) or PP2Ac (1:5000; Millipore) for quantitation. Equal loading proteins were further confirmed by probing with anti-calnexin antiserum (1:1000; Enzo Life Sciences). All Western-blot analyses were performed a minimum of twice and the data were averaged. The integrated density value of each phosphoprotein (phospho-synapsin Iα, Iβ, and IIα Ser9 measured together) and total protein band were measured using ImageJ software (NIH).

Co-Immunoprecipitation

The procedure of co-immunoprecipitation was previously described (Sun et al., 2009). Briefly, equal amounts of protein extracts (250 µg) were pooled from 2 to 3 rats per group and incubated in RIPA lysis buffer containing antibody against total synapsin (9 µg; Cell Signaling) or PP2Ac (8 µg; Millipore) overnight, followed by the addition of protein A/G agarose beads (30 µL; Santa Cruz Technologies) for 4 hours at 4°C. Beads were then washed 4 times in the RIPA lysis buffer. After final centrifugation, Lammeli buffer containing 1% β-mercaptoethanol was added, boiled for 5 minutes, loaded into gels, and analyzed by immunoblotting procedures as described above. Two percent of protein extract was used as input to normalize the protein band intensity.

Statistical Analysis

Two standard deviations from the mean were used to exclude potential outliers of each group. Independent t tests were used for the SA behavioral measurements. For biochemical data, 2-way analysis of variances (ANOVA) [treatment (saline vs cocaine) × infusion (vehicle vs BDNF)] were used followed by Newman–Keuls (SNK) multiple comparison tests when a significant interaction or main effect was found after ANOVA. Because of the small sample size (3–5 samples/group), Fisher’s Least Significant Difference test was used to analyze the co-immunoprecipitation data after ANOVA. The criterion for statistically significant differences was set at P<.05.
Results

During the last 3 SA sessions, cocaine self-administering rats exhibited significantly lower inactive lever pressing and higher active lever pressing than yoked-saline controls, indicating the acquisition of cocaine-associated reinforcing effects (Table 1; 2 hours: inactive lever: \( t_{29} = -3.31, P < .01 \); active lever: \( t_{29} = 10.53, P < .001 \); 22 hours: inactive lever: \( t_{29} = -4.10, P < .001 \); active lever: \( t_{29} = 7.66, P < .001 \)). Before receiving intra-dmPFC vehicle or BDNF infusion, rats with the cocaine SA history pressed a similar amount on the active levers and received a similar number of cocaine infusions.

Two hours after intra-dmPFC infusions, there was a significant treatment \( \times \) infusion interaction for p-synapsin Ser9 in the NAc (\( F_{1,24} = 5.20, P < .05 \)). A SNK test revealed greater p-synapsin Ser9 protein expression in the cocaine-/vehicle-treated group than in the saline-/vehicle- and cocaine-/BDNF-treated groups (\( P < .05 \)) (Figure 1a). Similarly, for the protein expression of p-synapsin Ser62/67, there was a trend toward a main effect of treatment (\( F_{1,24} = 3.68, P = .07 \)) with a significant treatment \( \times \) infusion interaction (\( F_{1,24} = 7.16, P < .05 \)). The cocaine-/vehicle-treated group alone had higher p-synapsin Ser62/67 in the NAc than in other groups (\( P < .05 \)) (Figure 1b). Neither cocaine nor BDNF altered p-synapsin Ser603 protein expression. Twenty-two hours after infusion, there was also a trend toward a main effect of infusion and a significant treatment \( \times \) infusion interaction for p-synapsin Ser9 (\( F_{1,24} = 3.49, P = .08 \) and \( F_{1,24} = 5.01, P < .05 \), respectively) (Figure 2a). The SNK multiple comparison revealed that the protein expression of p-synapsin Ser9 was significantly greater in the cocaine/vehicle group than in other groups (\( P < .05 \)). The protein levels of p-synapsin Ser62/67 and Ser603 were not changed 22 hours after infusion (Figure 2b and c). Further, there was no alteration in total synapsin expression at either time point examined.

PP2Ac is the protein phosphatase that dephosphorylates p-synapsin Ser9 (Jovanovic et al., 2001). We further evaluated p-PP2Ac Tyr307 (associated with inactivation of PP2A) protein levels due to the augmentation of p-synapsin Ser9 in the NAc and 22 hours after the end of cocaine SA. Two hours after infusion, 2-way ANOVA indicated that there was a significant main effect of treatment (\( F_{1,24} = 10.68, P < .01 \)) and a treatment \( \times \) infusion interaction (\( F_{1,24} = 10.03, P < .01 \)). SNK multiple comparisons showed a higher p-PP2Ac Tyr307 protein induction in the cocaine/vehicle group than in other groups (\( P < .01 \)) (Figure 3a). Similarly, 22 hours after infusion, there was a significant main effect of treatment (\( F_{1,24} = 9.60, P < .05 \)) with a treatment \( \times \) infusion interaction (\( F_{1,24} = 5.28, P < .05 \)). Further analysis indicated that the protein expression of p-PP2Ac in the cocaine-/vehicle-treated group was significantly increased compared with other groups (\( P < .01 \)) (Figure 3b). The total PP2Ac protein levels were not altered at either time point examined.

Table 1. Average Number of Lever Presses and Cocaine Infusions during the Last 3 Sessions of SA

<table>
<thead>
<tr>
<th>Group</th>
<th>Treatment</th>
<th>Inactive Lever</th>
<th>Active Lever</th>
<th>Cocaine Infusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 h</td>
<td>Yoked-saline</td>
<td>4.19 ± 1.40</td>
<td>5.50 ± 1.23</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>Cocaine SA</td>
<td>0.40 ± 0.25*</td>
<td>35.31 ± 2.10*</td>
<td>27.49 ± 1.35</td>
</tr>
<tr>
<td>22 h</td>
<td>Yoked-saline</td>
<td>3.81 ± 0.88</td>
<td>6.83 ± 1.45</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>Cocaine SA</td>
<td>0.26 ± 0.18*</td>
<td>35.13 ± 3.69*</td>
<td>28.92 ± 3.69</td>
</tr>
</tbody>
</table>

Abbreviation: SA, self-administration.

Discussion

In the present study, we demonstrated that p-synapsin Ser9 was increased in the NAc after 2 and 22 hours of withdrawal from cocaine SA. Elevation of p-synapsin Ser62/67 was also observed in cocaine-treated rats, but only at the 2-hour time point. Phosphorylation of PP2Ac was significantly elevated (indicating inactivation) after cocaine SA at the 2 time points examined. This elevation and that of p-synapsin at both the Ser9 and Ser62/67 sites was normalized by intra-dmPFC BDNF infusion. Further, co-immunoprecipitation revealed a protein-protein interaction between synapsin and PP2Ac in presynaptic terminals that cocaine reduced and intra-dmPFC BDNF infusion reversed.

These data indicate that during early withdrawal from cocaine SA, p-synapsin Ser9 is elevated due to a dissociation of PP2Ac from synapsin and a decrease in PP2A activity in presynaptic terminals in the NAc. Intra-dmPFC BDNF reactivates PP2A, resulting in enhanced binding of PP2A to, and dephosphorylation of, p-synapsin Ser9 in NAc terminals. Although these terminals likely originate from PFC neurons, indirect effects via the amygdala, for example, cannot be ruled out. Since PP2A also mediates the deactivation of CaMKII (DeRemer et al., 1992) and CAMKII is localized in the neuropil (Picciotto et al., 1995), future studies can focus on whether elevation of p-synapsin Ser9 is associated with a reduction of PP2A’s inhibitory effect on CaMKII in PFC-NAc terminals. Elevated p-synapsin Ser9 in NAc immediately after the last cocaine SA session using a different regimen (4-hour daily sessions) has been previously detected (Edwards et al., 2007), suggesting that changes in p-synapsin Ser9 expression are common time-dependent monitors of presynaptic plasticity in NAc in response to cocaine SA.

In contrast to the elevated p-synapsin Ser9 after 2 and 22 hours of withdrawal from cocaine, an increase in ERK-mediated p-synapsin Ser62/67 was observed only at the 2-hour time point.
The phosphorylation of ERK-mediated p-synapsin in response to low activity at 2 hours may be reversed as neuronal activity is restored and more Ca^{2+} influx triggers PP2B activation with time (22 hours). A similar upregulation of p-synapsin Ser9 and downregulation of ERK-activated p-synapsin Ser62/67 has been documented after 4-aminopyridine induced Ca^{2+} influx (Jovanovic et al., 2001). Moreover, a decrease in phosphorylation of ERK and CREB in dmPFC was observed at 2 hours, but not after...
22 hours, of abstinence from cocaine SA (Whitfield et al., 2011), suggesting a gradual recovery of neuronal activity in the PFC as the phase of withdrawal changes. Therefore, the dynamic alterations of p-synapsin 62/67 in the NAc may represent a sensitive biomarker of PFC neuronal activity changes. Because CaMK- and its upstream activators as well as other possible alternative intracellular mechanisms.

Figure 4. Intra-dorsomedial (dm) prefrontal cortex (PFC) brain-derived neurotrophic factor (BDNF) infusion normalizes the reduction of PP2A:synapsin I binding in the nucleus accumbens (NAc) after cocaine self-administration (SA). (a) A control experiment shows images of specificity for PP2A and synapsin binding in the NAc from drug naive rats. Left: Representative images of NAc homogenates immunoprecipitated by t-synapsin followed by probing with t-synapsin and t-PP2Ac antibodies. Right: Immunoblots illustrating the bands of t-synapsin and t-PP2Ac after t-PP2Ac antibody immunoprecipitation. (b) Top: Quantitative analysis of PP2Ac-synapsin protein binding levels in the NAc 2 hours after the end of cocaine SA. Bottom: Representative bands of an immunoblot of t-PP2Ac and its input protein levels after immunoprecipitation by t-synapsin antibody from each group (SV: N = 3; SB: N = 3; CV: N = 4; CB: N = 5). Histograms show the mean ± SEM of each group. *P < .05 vs SV; #P < .05 vs CB, 2-way analysis of variance (ANOVA) followed by Fisher’s Least Significant Difference (LSD) multiple comparison. B, BDNF; C, cocaine; CNXN, calnexin; IB, immunoblotting; IP, immunoprecipitation; S, saline; V, vehicle.

In conclusion, we demonstrated both p-synapsin Ser9 and Ser62/67 phosphorylation in the NAc during early withdrawal is likely related to BDNF’s ability to increase synaptic activity. Emerging evidence indicates that inhibition of synaptic activity in the PFC blocks the ability of intra-PFC BDNF to suppress cocaine-seeking (Go and McGinty, 2013). In addition, BDNF has been demonstrated to increase excitatory glutamatergic neuronal activity (Haubensak et al., 1998) as well as PP2A activity in cortical neurons (Jovanovic et al., 2004). Sustained depolarization also increases PP2A activity in midbrain dopamine neurons (Padmanabhan and Prasad, 2009). Thus, it is possible that intra-dmPFC BDNF infusion activates PP2A/PP2B and normalizes p-synapsin Ser9 and Ser62/67 levels and restores the cocaine-depleted reserve pool of synaptic vesicles in NAc terminals as a function of restoring PFC neuronal activity during early withdrawal. Future studies will attempt to extend and confirm this hypothesis by establishing whether cocaine SA depletes, and BDNF restores, the reserve pool of SVs in the NAc and the possible underlying mechanisms. Surprisingly, in the NAc of yoked-saline animals, we did not observe any p-synapsin changes after intra-dmPFC BDNF infusion, which is in contrast to a previous study demonstrating the elevation of p-synapsin in the synaptosomal fraction induced by local BDNF application (Jovanovic et al., 2010). Because total homogenates of NAc were used and BDNF was infused in the dmPFC in the present study, the underlying mechanism of BDNF effects on the local axon terminals in the NAc could not be examined in detail, which may partially explain the discrepancy. Further experiments using synaptosomal and/or presynaptic fractions are needed to elucidate BDNF’s effects on p-synapsin and its upstream activators as well as other possible alternative intracellular mechanisms.

In conclusion, we demonstrated both p-synapsin Ser9 and Ser62/67 in the NAc were induced during early withdrawal (Greenfield et al., 2011) is accompanied by the attenuated phosphorylation of striatal-enriched protein tyrosine phosphatase and its targets, GluN1/2B, in the dmPFC after 2 hours of abstinence from cocaine SA (Sun et al., 2013). Increased phosphatase activity and decreased glutamatergic transmission may indicate synaptic depression in the PFC as a critical part of the withdrawal cascade. Second, basal extracellular glutamate levels in the NAc are similarly reduced after 1, 21, or 60 days of cocaine SA abstinence (Lutgen et al., 2014; Wydra et al., 2013), but a specific enhancement of evoked release probability at PFC-NAc synapses occurs after 1 day of abstinence (Suska et al., 2015) and excessive evoked glutamate release occurs in response to cocaine prime-induced reinstatement after extinction (McFarland et al., 2003). Third, basal electrophysiological activity in the PFC of rats is suppressed, but stimulus-evoked burst duration and firing rate are enhanced within 24 hours after the end of cocaine SA (Sun and Rebec, 2006). Fourth, downregulation of immediate early gene and Bdnf mRNA occurs within 22 hours of cocaine withdrawal, but robust upregulation occurs in response to cue-induced relapse to drug-seeking (Hearing et al., 2008; McGinty et al., 2010). Finally, basal hypoactivity punctuated by hyper-responsiveness of PFC to cues associated with cocaine are critical disturbances in abstinent human cocaine addicts that are associated with craving and relapse (Childress et al., 1999; Goldstein and Volkow, 2011). Thus, it is possible that by repeatedly driving synaptic activity and excessive glutamate release in NAc (Migliens et al., 2008), cocaine SA causes hyper-phosphorylation of synapsins and depletion of the reserve SV pool along with increased evoked release probability that extends into withdrawal.

The mechanism underlying the ability of a BDNF infusion into dmPFC to normalize synapsin and PP2A phosphorylation in the NAc during early withdrawal is likely related to BDNF’s ability to increase synaptic activity. Emerging evidence indicates that inhibition of synaptic activity in the PFC blocks the ability of intra-PFC BDNF to suppress cocaine-seeking (Go and McGinty, 2013). In addition, BDNF has been demonstrated to increase excitatory glutamatergic neuronal activity (Haubensak et al., 1998) as well as PP2A activity in cortical neurons (Jovanovic et al., 2004). Sustained depolarization also increases PP2A activity in midbrain dopamine neurons (Padmanabhan and Prasad, 2009). Thus, it is possible that intra-dmPFC BDNF infusion activates PP2A/PP2B and normalizes p-synapsin Ser9 and Ser62/67 levels and restores the cocaine-depleted reserve pool of synaptic vesicles in NAc terminals as a function of restoring PFC neuronal activity during early withdrawal. Future studies will attempt to extend and confirm this hypothesis by establishing whether cocaine SA depletes, and BDNF restores, the reserve pool of SVs in the NAc and the possible underlying mechanisms. Surprisingly, in the NAc of yoked-saline animals, we did not observe any p-synapsin changes after intra-dmPFC BDNF infusion, which is in contrast to a previous study demonstrating the elevation of p-synapsin in the synaptosomal fraction induced by local BDNF application (Jovanovic et al., 2010). Because total homogenates of NAc were used and BDNF was infused in the dmPFC in the present study, the underlying mechanism of BDNF effects on the local axon terminals in the NAc could not be examined in detail, which may partially explain the discrepancy. Further experiments using synaptosomal and/or presynaptic fractions are needed to elucidate BDNF’s effects on p-synapsin and its upstream activators as well as other possible alternative intracellular mechanisms.

In conclusion, we demonstrated both p-synapsin Ser9 and Ser62/67 in the NAc were induced during early withdrawal...
from cocaine SA. This hyper-phosphorylation of synapsins may represent a molecular disturbance of the reserve pool of synaptic vesicles that contributes to synaptic depression and alters release probability in the PFC-NAC pathway during withdrawal from cocaine SA. However, the possible contribution of PFC-amygdala-NAC input on synapsin phosphorylation cannot be excluded, since BDNF was elevated in the amygdala after intra-dmPFC BDNF infusion (McIntyre et al., 2010). Nevertheless, together with our previous findings (Berglind et al., 2007, 2009; Whitfield et al., 2011), the ability of intra-dmPFC BDNF infusion to rescue p-synapsin in the NAC and related signaling cascades in the PFC suggests that the restoration of PFC function during early withdrawal may abrogate maladaptive responses that disturb the PFC output to the NAC. Therefore, the BDNF/TrkB system in the PFC-NAC pathway remains a potential therapeutic target for early intervention in cocaine addiction.

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Statement of Interest
None.

References


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