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Differential Regulation of MeCP2 Phosphorylation in the CNS by Dopamine and Serotonin

Ashley N Hutchinson¹, Jie V Deng¹, Dipendra K Aryal², William C Wetsel^{1,2,3,4} and Anne E West^{*,1}

¹Department of Neurobiology, Duke University Medical Center, Durham, NC, USA; ²Department of Psychiatry and Behavioral Sciences, Duke University Medical Center, Durham, NC, USA; ³Mouse Behavioral and Neuroendocrine Analysis Core Facility, Duke University Medical Center, Durham, NC, USA; ⁴Department of Cell Biology, Duke University Medical Center, Durham, NC, USA

Systemic administration of amphetamine (AMPH) induces phosphorylation of MeCP2 at Ser421 (pMeCP2) in select populations of neurons in the mesolimbocortical brain regions. Because AMPH simultaneously activates multiple monoamine neurotransmitter systems, here we examined the ability of dopamine (DA), serotonin (5-HT), and norepinephrine (NE) to induce pMeCP2. Selective blockade of the DA transporter (DAT) or the 5-HT transporter (SERT), but not the NE transporter (NET), was sufficient to induce pMeCP2 in the CNS. DAT blockade induced pMeCP2 in the prelimbic cortex (PLC) and nucleus accumbens (NAc), whereas SERT blockade induced pMeCP2 in the prelimbic cortex (PLC) and nucleus accumbens (NAc), whereas SERT blockade induced pMeCP2 induced pMeCP2 in the prelimbic cortex (PLC) and nucleus accumbens (NAc), whereas SERT blockade induced pMeCP2 only in the NAc. Administration of selective DA and 5-HT receptor agonists was also sufficient to induce pMeCP2; however, the specific combination of DA and 5-HT receptors activated determined the regional- and cell-type specificity of pMeCP2 induction. The D₁-class DA receptor agonist SKF81297 induced pMeCP2 widely; however, coadministration of the adenylate cyclase activator forskolin was sufficient to induce pMeCP2 in medium-spiny neurons, suggesting that the combinatorial regulation of cAMP by different classes of DA and 5-HT receptors may contribute to the cell-type specificity of pMeCP2 induction. Consistent with the regulation of pMeCP2 by multiple monoamine neurotransmitters, genetic disruption of any single monoamine transporter in DAT-, SERT-, and NET-knockout mice failed to eliminate AMPH-induced pMeCP2 in the NAc. Together, these studies indicate that combinatorial signaling through DA and 5-HT receptors can regulate the brain region- and cell-type specific pMeCP2 in the CNS. *Neuropsychopharmacology* (2012) **37**, 321–337; doi:10.1038/npp.2011.190; published online 28 September 2011

Keywords: MeCP2; monoamine transporters; signal transduction; dopamine; serotonin; nucleus accumbens

INTRODUCTION

Psychostimulant-induced activation of monoamine receptors is essential to the mechanism by which these drugs induce changes in behavior (Gainetdinov and Caron, 2003; Missale *et al*, 1998; Xu *et al*, 1997). Cocaine and amphetamine (AMPH) serve as indirect monoamine receptor agonists by inhibiting and/or reversing the function of monoamine transporters. However, despite the fact that these drugs rapidly increase extracellular levels of dopamine (DA), serotonin (5-HT), and norepinephrine (NE), repeated exposure is necessary to produce longlasting alterations in behavior (Hyman *et al*, 2006). This observation suggests that molecular mechanisms downstream of monoamine receptor activation are required to mediate the observed behavioral adaptations.

*Correspondence: Dr AE West, Department of Neurobiology, Duke University Medical Center, Box 3209, Durham, NC 27710, USA, Tel: + I 919 681 1909, Fax: + 919 684 4431,

E-mail: west@neuro.duke.edu

A substantial body of evidence indicates that psychostimulant-induced changes in gene expression contribute to the long-lasting effects of these drugs on behavior (Nestler, 2008; Kumar et al, 2005). Examination of the molecular processes underlying these changes in gene expression has revealed a significant role for epigenetic mechanisms of transcription, which are processes that mediate changes in gene expression by regulating chromatin structure (Maze et al, 2010; Renthal et al, 2009). Therefore, chromatin regulatory proteins that are targets of modulation by monoaminergic signaling pathways are strong candidates to link psychostimulant exposure to persistent changes in gene expression and behavior. Recently, we and others have demonstrated that the methyl-CpG-binding protein-2 (MeCP2) contributes to both neural and behavioral adaptations induced by repeated psychostimulant exposure (Deng et al, 2010; Im et al, 2010). We have shown that lentiviral shRNA-mediated knockdown of MeCP2 in the nucleus accumbens (NAc) increases sensitivity to AMPH-induced conditioned place-preference (CPP) whereas overexpression of MeCP2 decreases CPP (Deng et al, 2010). These data suggest that MeCP2 acts in the NAc to limit the rewarding properties of psychostimulants.

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Received 29 March 2011; revised 29 July 2011; accepted 29 July 2011

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Psychostimulants induce rapid and robust phosphorylation of MeCP2 at Ser421 (pMeCP2), a site that is thought to modulate MeCP2-dependent chromatin regulation (Deng et al, 2010; Zhou et al, 2006; Chen et al, 2003). MeCP2 regulates gene transcription by binding to methylated DNA and recruiting enzymes that modify histones (Nan et al, 1998). Although the specific mechanisms by which Ser421 phosphorylation affects MeCP2 function remain poorly understood, overexpression of a Ser421Ala mutant MeCP2 has been shown to inhibit neuronal activity-induced Bdnf expression, suggesting that Ser421 phosphorylation could contribute to gene derepression (Zhou et al, 2006). Interestingly, systemic administration of AMPH induces a restricted pattern of pMeCP2 induction in select neurons of the NAc and prelimbic cortex (PLC) (Deng et al, 2010). D₁-class DA receptor activation is required for AMPHdependent induction of pMeCP2 in the NAc, and administration of a D₁-class DA receptor agonist is sufficient to induce pMeCP2 (Deng et al, 2010). However, AMPH induces pMeCP2 selectively in GAD67- and parvalbuminpositive GABAergic interneurons of the NAc whereas it fails to induce pMeCP2 in medium-spiny neurons (MSNs), despite the fact that these cells express high levels of DA receptors. In addition to its effects on DA neurotransmission, AMPH also impacts 5-HT and NE neurotransmission. Thus, we considered the possibility that the selective pattern of pMeCP2 induced by AMPH might arise as a result of combinatorial signaling through one or more of these monoaminergic systems.

Here, we use both pharmacological and genetic techniques to characterize the differential roles of DA, 5-HT, and NE signaling in the regulation of the regional distribution and cell-type specificity of pMeCP2 induction. We find that selective inhibition of either the DA transporter (DAT) or 5-HT transporter (SERT) but not the NE transporter (NET) is sufficient to induce pMeCP2 in the NAc. However, only DAT inhibition induces pMeCP2 in the PLC. We also find that differential activation of specific classes of DA and 5-HT receptors induces pMeCP2 in distinct cell types and brain regions, suggesting that combinatorial signaling through these receptors sculpts the specific pattern of pMeCP2 induction in vivo. Finally, we discuss potential functional roles for pMeCP2 based on our analyses of AMPH-induced pMeCP2 in DAT-, SERT-, and NET-knockout (KO) mouse strains, each of which has a distinct profile of behavioral responses to AMPH administration. Taken together, these data significantly enhance understanding of the mechanisms and consequences of monoaminergic regulation of MeCP2 in the CNS.

MATERIALS AND METHODS

Animals

Adult (8–10 week old) male C57BL/6 mice (Jackson Laboratories, Bar Harbor, ME), DAT, NET, vesicular monoamine transporter 2 (VMAT2) mice (provided by Dr Marc Caron at Duke University Medical Center, Durham, NC), and SERT mice (provided by Dr Scott Hall, NIDA, Bethesda, MD) were used in these studies. The wild-type (WT), heterozygous (HET), and KO littermates were generated from HET breedings. Animals were weaned at

21–30 days of age, housed in groups of 3–5, and segregated by sex and genotype. All animals were given free access to standard laboratory chow and water and were housed in a humidity- and temperature-controlled room on a 14-h/10-h light/dark cycle (lights on at 0700 hours). All experiments were conducted with an approved protocol from the Duke University Institutional Animal Care and Use Committee in accordance with the guidelines from the National Institutes of Health for the Care and Use of Laboratory Animals.

Immunofluorescent Staining of the Brain Sections

Two hours after vehicle or drug injection, mice were perfused transcardially with 4% paraformaldehyde in 0.1 M PBS. The brains were post-fixed in 4% paraformaldehyde/PBS overnight, then sunk into 20% (wt/vol) sucrose/PBS overnight. Coronal sections (40 µm) were cut on a freezing microtome, and the brain regions were identified by anatomical landmarks. One section from each brain region of interest was selected for each mouse, based upon anatomical structures to represent the closest approximation of identical sections between individual mice. To minimize technical variations in immunostaining across genotypes, sections from different individual mice were first photographed for visual identification. The sections were then pooled and incubated with antibodies in a single chamber, and finally they were separated after processing for image analysis (see Deng et al, 2010). For immunostaining, tissue sections were permeabilized with either 1% (vol/vol) (for the pMeCP2 antibody) or 0.3% (vol/vol) (all other antibodies) Triton X-100 for 1 h and then sections were blocked with 3% (wt/vol) BSA (for the DARPP-32 antibody) or 16% (vol/vol) goat serum (all other antibodies) in PBS. Sections were incubated with the following primary antibodies overnight at 4 °C: rabbit anti-phospho-Ser421 MeCP2 1:15000 (Deng et al, 2010), mouse anti-GAD67 1:500 (MAB5406; Chemicon/Millipore, Billerica, MA), rabbit anti-c-Fos 1:15 000 (PC38; Calbiochem, San Diego, CA), goat anti-DARPP-32 1:50 (sc31519; Santa Cruz Biotechnology, Santa Cruz, CA), and mouse anti-GFAP 1:100 (Clone G-A-5; Sigma, St Louis, MO). Whenever possible, double immunostaining with primary antibodies raised in two different species was used for colocalization on single sections in order to determine the cell type in which pMeCP2 was induced. After three washes in PBS, sections were incubated with the following species-specific fluorescentconjugated secondary antibodies for 1 h at room temperature: goat anti-mouse antibodies conjugated to Cy3 or Cy2 1:500 (Jackson Immunoresearch, West Grove, PA), Alexa Fluor 488 goat anti-rabbit antibodies at 1:500 (A11034; Molecular Probes/Invitrogen, Carlsbad, CA), Cy3 donkey anti-rabbit antibodies at 1:500 (Jackson Immunoresearch), or Alexa Fluor 488 donkey anti-goat antibodies (A11055; Molecular Probes/ Invitrogen). Sections were washed in PBS, nuclei were labeled with Hoechst dye (Sigma) to facilitate anatomical localization of the brain structures, sections were mounted and coverslipped, and analyzed as described below.

Image Analyses

For quantitative immunofluorescence, images were captured on a Leica DMI4000 inverted fluorescence microscope using a Cascade 512B camera. Digital images were quantified using **Differential regulation of MeCP2 phosphorylation** AN Hutchinson *et al*

MetaMorph 7 Image Analysis software (Molecular Devices, Sunnyvale, CA). To minimize variation between samples, images were captured with a uniform exposure time within a single experiment, and immunofluorescence was quantified across a constant-sized region from a single field of each section (as shown in Figure 1a). We used the Count Nuclei module in MetaMorph 7.0 to first count the total number of cells per section that were positive for pMeCP2 or c-Fos immunoreactivity and then we determined the integrated immunofluorescence intensity of all the pMeCP2 or c-Fospositive nuclei in each image. We defined nuclei as objects of $3-8\,\mu\text{m}$ in diameter. For each experiment, an investigator blind to genotype and treatment condition chose a random section from which to set a single threshold value of fluorescence intensity above background to score objects of this size as positive for pMeCP2 or c-Fos expression. The threshold value was adjusted until the program's output of detected nuclei most closely matched the distribution of positive nuclei seen by eye. This threshold value was then held constant for every image within the experimental set. The first result of this analysis is a count of the number of cells that are pMeCP2 or c-Fos positive in each image. The Count Nuclei module then creates a mask that covers the area of all of the nuclei identified as positive on each section and quantifies the total immunofluorescence intensity under this area (the 'integrated intensity') for each image. Thus, our analysis allows us to tell how many cells have induced expression of pMeCP2 or c-Fos in each image as well as to evaluate the magnitude of pMeCP2 or c-Fos induction in these nuclei.



Figure 1 Selective DAT and SERT blockade differentially induces pMeCP2 in the NAc. (a) pMeCP2 and total MeCP2 induction following vehicle or 3 mg/kg AMPH injection. Solid gray box indicates representative section of NAc used to quantify pMeCP2 immunofluorescence. Dashed white box was used as the inset image to examine total MeCP2 and Hoechst (nuclear marker) overlay. ac, anterior commissure. (b) pMeCP2 immunostaining in the NAc (left column) and c-Fos immunostaining (right column) 2 h after treatment with Veh, 10 mg/kg GBR12909, 10 mg/kg citalopram, or 3 mg/kg reboxetine. (c, e, g) number of cells in which pMeCP2 was induced following treatment of GBR12909, citalopram, or reboxetine. (d, f, h) pMeCP2 integrated intensity in the NAc following treatment with GBR12909, citalopram, or reboxetine. All values of quantitated immunofluorescence intensity were normalized to Veh control for each drug. n = 5-8 mice/group. *p < 0.05 compared with Veh. Error bars indicate SEM. Scale bars = 10 µm.

Bilateral Guide Cannula Surgery

Mice were anesthetized with chloral hydrate (400 mg/kg i.p.). A 22-gauge bilateral stainless steel guide cannula assembly with center-to-center distance of 2.0 mm (PlasticsOne, Raanoke, VA) was implanted over the striatum in both hemispheres. Stereotaxic coordinates relative to bregma were A/P + 1.42 mm, $M/L \pm 1.0$ mm, D/V - 3.6 mm. Guide cannulae were anchored to the skull using carboxylate dental cement (Durelon, CMA Micro-dialysis). A dust cap was placed over the entire cannula assembly. The mice were housed individually after surgery and allowed to recover for 8–9 days. Mice were treated with the antibiotic cotrimoxazole, 3% w/v solution in water, for the first 3 days following surgery.

Intra-Striatal Infusions

All infusions were performed in an empty cage that was the same dimensions as the home cage and were administered in freely moving mice. A 28-gauge stainless steel injector cannula with 3.6 mm length was used for all drug infusions. The injection cannula was attached to a 25-µl Hamilton syringe (Hamilton, Reno, NV). In all, 1 µl of drug and saline was simultaneously infused into the left and right hemispheres over 6 min using a Razel infusion pump (Razel Scientific Instruments, St Albans, VT). The mice were infused with 2.5 µg/µl forskolin, 7-deacetyl-7-[0-(N-methylpiperazino)-g-butryl]-dihydrochloride (Calbiochem, La Jolla, CA) in one hemisphere and vehicle (saline) in the other so that each mouse served as its own control. Side selection was randomized. Injection cannulae remained in place for an additional 2 min to allow drug diffusion and then a dummy cannula was inserted into the guide to prevent backflow. The mice were perfused transcardially with 4% paraformaldehyde 90 min after infusion.

Striatal Cultures

Whole striatum was dissected from embryonic day 17 CD-1 rats (Charles River Laboratories, Wilmington, MA), dissociated mechanically, and plated on PDL-coated dishes containing DMEM/F12 medium (Invitrogen) with B27 supplements (Invitrogen). After 5 days in culture, neurons were stimulated with the following drugs: 55 mM KCl in an isotonic solution (see Tao et al, 1998), 50 µM NMDA (Sigma), 10 µM forskolin (EMD, Gibbstown, NJ), or 50 µM SKF38393 (Sigma). In some experiments, 5 µM Nimodipine (Sigma) or 100 µM APV (Tocris) was added 2 min before stimulation with KCl or NMDA. Sixty minutes after stimulation, cells were lysed in boiling SDS-PAGE sample buffer for analysis of protein phosphorylation by western blotting. In addition to the MeCP2 antibodies described above, the following antibodies were used: rabbit anti-CREB 1:1000 (06-863; Upstate Biotechnology/Millipore), mouse anti-phospho-Ser133 CREB 1:1000 (05-667; Upstate Biotechnology/Millipore), goat anti-rabbit-HRP 1:10 000 (Jackson Immunoresearch), and goat anti-mouse-HRP 1:10 000 (Jackson Immunoresearch).

Locomotor Activity in the Open Field

Mice were habituated to the open field (Accuscan Instruments, Columbus, OH) for 1 h to establish baseline activity and then injected with vehicle (Veh) or 3 mg/kg AMPH (Sigma), 5 mg/kg SFK81297 (Tocris Bioscience, Ellisville, MO), 0.25 mg/kg quinpirole (Tocris), 5 or 10 mg/kg citalopram (Tocris), 10 or 15 mg/kg GBR12909 (Tocris), 3 or 5 mg/kg reboxetine (Tocris), or 3 or 4.5 mg/kg quipazine (Tocris). All injections were i.p., and all drugs were dissolved in sterile water, except SKF81297, which was dissolved in DMSO (final concentration 0.5% vol/vol). Following administration of Veh or drug, animals were immediately returned to the open field for 1 h. Horizontal (distance traveled in cm) was monitored under 340 lux illumination.

Behavioral Sensitization

Two different protocols were used as referenced in the text. In the first protocol, mice were injected with Veh or 3 mg/kg AMPH once a day for 5 consecutive days in the home cage, they were withdrawn from drug for 7 days, then they were challenged with 3 mg/kg AMPH in the open field while horizontal activity (distance traveled in cm) was monitored as described in the 'Locomotor Activity in the Open Field' section above. In the second protocol, the first injection of vehicle or AMPH was given in the open field so that locomotor activity could be recorded. The mice were then returned to the home cage where they received daily injections of Veh or 3 mg/kg AMPH for 4 additional consecutive days. The mice were then withdrawn from drug for 7 days, then they were challenged with 3 mg/kg AMPH in the open field while horizontal activity (distance traveled in cm) was monitored.

Statistical Analyses

Statistical analyses were performed using SPSS v11.0 statistical software (SPSS, Chicago, IL). The data are depicted as mean values and standard errors of the mean (SEM). The pMeCP2 immunoreactivities and locomotor activities of the transporter KO mice given Veh or AMPH were analyzed using two-way ANOVA for genotype and treatment. Activities of the mice were aggregated over the 1-h post-injection period. For C57BL/6J mice injected with Veh, GBR12909, citalopram, reboxetine, SKF81297, quinpirole, SKF81297 + quinpirole, quinpirole alone, or quipazine, pMeCP2 immunoreactivities in the NAc were analyzed using univariate ANOVA for treatment condition. Bonferroni corrected pairwise comparisons were used as the post hoc tests. Induction of Fos and pMeCP2 by intra-striatal forskolin was assessed by a Student's unpaired t-test. We assessed the correlation between pMeCP2 immunoreactivity and locomotor activity with Pearson correlation coefficients. In all cases, p < 0.05 was considered statistically significant.

RESULTS

Selective Blockade of the DAT or SERT Is Sufficient to Induce pMeCP2

Pharmacological targets of AMPH in the CNS include the DAT, SERT, and NET plasma membrane transporters, as well as the VMAT2 vesicular transporter (Fleckenstein *et al*,

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Drug	Distance (cm) Veh	Distance (cm) dose I	Distance (cm) dose 2
GBR12909	3390.3 ± 66.2	3645.6± 627.6*	2482.4±2 53.6*
Citalopram	3946.6 ± 690. I	2514.2 ± 407.9	729.0 ± 241.9*
Reboxetine	727.5 ± 96.	1036.2 ± 277.8	747.8±118.8*

The effects of 10 or 15 mg/kg GBR12909, 5 or 10 mg/kg citalopram, and 3 or 5 mg/kg reboxetine on locomotor activity in C57BL/6 mice 1 h post-injection. Dose I = lower dose; dose 2 = higher dose. All values of locomotor activity were normalized to Veh control for each drug. n = 6-8 mice/group. *p < 0.05 compared with Veh. Error bars indicate SEM.

2007; Han and Gu, 2006; Sulzer et al, 2005). Through its interactions with these target proteins, AMPH simultaneously raises extracellular levels of multiple monoamine neurotransmitters. To address the contributions of single monoaminergic neurotransmitter systems to the regulation of MeCP2 phosphorylation, we evaluated the ability of selective monoamine transporter inhibitors to induce pMeCP2 in the NAc (Figure 1). C57BL/6 mice were administered either Veh, a DAT inhibitor (GBR12909, 10 mg/kg or 15 mg/kg), a SERT inhibitor (citalopram, 10 mg/kg or 15 mg/kg), or a NET inhibitor (reboxetine, 3 mg/kg or 5 mg/kg). Open field locomotor activity was monitored for 1h then the mice were perfused 2h postinjection to assess pMeCP2 immunoreactivity. For GBR12909, an ANOVA for locomotor activity revealed a significant effect of treatment ($F_{2,15} = 13.313$, p < 0.001). Both doses of GBR12909 significantly increased locomotor activity compared with mice treated with Veh (p-values < 0.004) (Table 1). For pMeCP2 immunoreactivity in mice treated with GBR12909, an ANOVA indicated a significant effect of treatment ($F_{2,14} = 12.083$, p < 0.001), and Bonferroni comparisons showed that pMeCP2 intensities in both the 10- and 15-mg/kg groups were significantly higher than the Veh group (p-values < 0.05) (Figure 1b-d). These data demonstrate that selective blockade of the DAT is sufficient to induce pMeCP2 in the NAc.

ANOVA for locomotor activity in citalopram-treated mice revealed a significant effect of treatment ($F_{2,16} = 13.02$, p < 0.001). Unlike GBR12909, citalopram treatment led to a reduction in locomotor activity, although 10 mg/kg was the only dose to significantly lower activity (p < 0.001) (Table 1). When changes in pMeCP2 induction were considered for citalopram-treated mice, the ANOVA detected a significant effect of treatment ($F_{2,16} = 4.236$, p < 0.04), post hoc analyses found that the pMeCP2 intensity was significantly higher than the Veh group only for the 10-mg/kg dose (p < 0.05) (Figure 1b, e, and f). These data demonstrate that selective blockade of the SERT is also sufficient to induce pMeCP2 in the NAc. Intriguingly, we note that while GBR12909 and citalopram have opposite effects on locomotor activity, both drugs lead to the induction of pMeCP2 in the NAc. We discuss the relationship between pMeCP2 phosphorylation and locomotor activity further in the Discussion section.

Finally, inspection of the ANOVA for locomotor activity of mice treated with reboxetine revealed a significant effect of treatment ($F_{2,16} = 4.317$, p < 0.041). Like citalopram, reboxetine treatment led to a dose-dependent reduction in locomotor activity; only 5 mg/kg reboxetine significantly reduced locomotor activity compared with Veh-treated controls (p < 0.048) (Table 1). In contrast to the effects of GBR12909 and citalopram, pMeCP2 immunoreactivity in reboxetine-treated mice did not show an overall effect of treatment ($F_{2,16} = 0.806$, p < 0.463), suggesting that activation of NE signaling is not sufficient to induce pMeCP2 in this context (Figure 1b, g, and h). In order to ensure that the failure to induce pMeCP2 was not due to improper selection of doses, we also examined the ability of each of these drugs to induce c-Fos expression. In all, 5 mg/kg reboxetine induced expression of c-Fos in the NAc to a similar extent as that seen following administration of 3 mg/kg AMPH, 10 mg/kg GBR12909, and 10 mg/kg citalopram. Hence, the doses of reboxetine used in this study are able to induce a transcriptional response in neurons of the NAc (Figure 1b) (Miyata et al, 2005). Together, these data show that selective activation of either DA or 5-HT but not NE signaling is sufficient to drive induction of pMeCP2 in the NAc.

Although most neurons within the NAc express MeCP2, we have shown that AMPH selectively induces pMeCP2 in a specific population of GABAergic interneurons that express the GABA synthesizing enzyme GAD67 and the calciumbinding protein parvalbumin (Deng *et al*, 2010). To determine whether selective monoamine transporter blockade can recapitulate this cell-type specificity of AMPH-induced pMeCP2, we colabeled NAc sections from mice treated with the selective transporter blockers with antibodies against both pMeCP2 and GAD67. Like AMPH, GBR12909 induced pMeCP2 only in GAD67-positive GA-BAergic interneurons of the NAc (Figure 2). However, examination of double-immunolabeled sections following citalopram administration revealed that the induced pMeCP2 is not restricted to this cell population.

AMPH-induced phosphorylation of MeCP2 is most robust in the NAc; however, AMPH drives a small, but statistically significant increase in pMeCP2 in the PLC (Deng et al, 2010). To determine which monoamine neurotransmitters are sufficient to induce pMeCP2 in this brain region, we asked whether GBR12909, citalopram, or reboxetine induced pMeCP2 in the PLC (Figure 3). ANOVA revealed a significant effect of treatment with GBR12909 (F_{2,14} = 5.667, p < 0.02); 10 mg/kg led to increased pMeCP2 induction compared with Veh controls (p < 0.02) (Figure 3a). Unlike GBR12909, citalopram or reboxetine failed to induce pMeCP2 in the PLC, despite the fact that citalopram induced pMeCP2 in the NAc of the same animals (Figures 3b and c and 1f, respectively). These data suggest that there is a selective role for DA in pMeCP2 induction in the PLC. Examination of PLC sections double immunolabeled for pMeCP2 and GAD67 revealed that the GBR12909-induced pMeCP2 in the PLC was not restricted to GAD67-positive GABAergic interneurons (Figure 3d). Taken together, these data indicate that DA and 5-HT play selective roles in the induction of pMeCP2 in different the brain regions and cell types.

DA and 5-HT Receptor Agonists Differentially Induce pMeCP2

Because selective inhibition of the DAT and SERT transporters is sufficient to drive pMeCP2, we next asked



Figure 2 DAT and SERT blockers differentially regulate the cell-type specificity of MeCP2 phosphorylation in the NAc. pMeCP2 (red), GAD67 (green), Hoechst nuclear dye (blue), and overlay immunostaining in the NAc 2 h after treatment with Veh, 3 mg/kg AMPH, 10 mg/kg GBR12909, or 10 mg/kg citalopram. White arrows indicate neurons coimmunolabeled with antibodies for pMeCP2 and GAD67, and blue arrows indicate pMeCP2 induction in neurons that are not GAD67-positive GABAergic interneurons. Scale bar = $20 \,\mu$ m.

which DA and 5-HT receptor subtypes mediate this response. DA acts through both D_1 - (D_1 and D_5) and D_2 -class (D_2 , D_3 , and D_4) receptors (Missale *et al*, 1998). Previously, we showed that SKF81297, a D₁-class receptor agonist, is sufficient to drive robust phosphorylation of MeCP2 in the NAc (Deng et al, 2010). To determine the magnitude and regional distribution of pMeCP2 following D₁-class receptor activation, C57BL/6J mice were treated with 5 or 7.5 mg/kg SKF81297. To examine pMeCP2 induction following activation of D2-class receptors, pMeCP2 levels were quantified following treatment with quinpirole (0.4 mg/kg), a D₂-class receptor agonist. Finally, because AMPH activates both D1- and D2-class receptors simultaneously, we examined how combinatorial DA receptor activation influenced the level and distribution of pMeCP2 induction by coinjecting SKF81297 (7.5 mg/kg) and quinpirole (0.4 mg/kg).

When an ANOVA was applied to pMeCP2 immunoreactivity in the NAc following administration of Veh, SKF81297 (7.5 mg/kg), quinpirole, or coinjection of SFK81297 + quinpirole, an overall effect of treatment was observed ($F_{3,16} = 22.3$, p < 0.001). SKF81297 led to higher levels of pMeCP2 induction compared with Veh controls (p < 0.001). Parenthetically, activation of D₁-class receptors alone led to more widespread induction of pMeCP2 than we observed following AMPH, both in the NAc, PLC, and most notably in the hippocampus, where we have not observed any AMPH-induced pMeCP2 (Figure 4a-c). Furthermore, upon examining the cell types in which pMeCP2 is induced within the NAc following injection of 7.5 mg/kg SKF81297, we find that unlike AMPH, the D₁-class receptor agonist induces pMeCP2 in both GAD67-positive interneurons and additional cell types (Figure 4d). These data suggest that additional AMPH-activated receptor systems may antagonize the effects of D₁-class signaling on the regional and cell-type specific induction of pMeCP2. One candidate for inhibition is D₂-class receptors, which can be activated by the agonist quinpirole. Treatment of mice with 0.4 mg/kg quinpirole alone leads to a small induction of pMeCP2 that is confined to the GABAergic interneurons of the NAc (Figure 4a, b, and d). Intriguingly, coinjection of this D₂-class receptor agonist significantly reduces the extent of SKF81297-induced pMeCP2 both in the NAc (p < 0.001) (Figure 4b), as well as in the hippocampus (Figure 4c). Furthermore, coinjection of SKF81297 and quinpirole restricts the induction of pMeCP2 in the NAc to the GABAergic interneurons (Figure 4d). These data are consistent with the known mechanism of action of AMPH on both D₁- and D₂-classes of receptors, and they suggest that combinatorial D₁- and D₂-class receptor signaling may restrict the pattern and extent of AMPH-induced pMeCP2 to a select population of neurons within the NAc.

In addition to examining induction of pMeCP2 following activation of DA receptors, we also wanted to investigate the pattern of pMeCP2 produced by stimulation of 5-HT receptors. There are 14 5-HT receptor subtypes, which comprise seven different classes of receptors (Barnes and Sharp, 1999). Since 5-HT₂ and 5-HT₃ receptors are highly expressed in the NAc, we asked whether administration of quipazine (3 or 4.5 mg/kg), a 5-HT receptor agonist that

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Figure 3 Selective DAT blockade induces pMeCP2 in the PLC. (a–c) pMeCP2 integrated intensity in the PLC following treatment with GBR12909, citalopram, or reboxetine. All values of quantified immunofluorescence intensity were normalized to Veh control for each drug. (d) pMeCP2 (red), GAD67 (green), and overlay immunostaining in the PLC 2 h after treatment with Veh, 10 mg/kg GBR12909, 10 mg/kg citalopram, or 3 mg/kg reboxetine. White arrows indicate neurons coimmunolabeled with antibodies for pMeCP2 and GAD67, and blue arrows indicate pMeCP2 induction in neurons that are not GAD67-positive GABAergic interneurons. n = 5-7 mice/group. *p < 0.05 compared with Veh. Error bars indicate SEM. Scale bars = 20 μ m.

shows preferential activation of 5-HT_2 and 5-HT_3 receptors, can induce pMeCP2. ANOVA revealed a significant effect of treatment with quipazine ($F_{2,9} = 7.898$, p < 0.01). Both 3 and 4.5 mg/kg quipazine drove significant induction of pMeCP2 in the NAc, and these effects were restricted to the GABAergic interneurons (*p*-values < 0.05) (Figure 5a and b). We note that the cell-type specific induction of pMeCP2 in the NAc following quipazine is in contrast to the pattern of phosphorylation following administration of the indirect agonist citalopram, which can activate signaling at all 14 5-HT receptor subtypes and which leads to induction of pMeCP2 that is more widespread (Figure 3). Together, these data suggest that different classes of DA and 5-HT receptors may have distinct effects on the pattern of pMeCP2 induction in the NAc.

Among the pharmacological reagents tested here, we saw the most widespread induction of pMeCP2 following administration of a D_1 -class DA receptor agonist. By contrast, a D_2 -class DA receptor agonist opposed the effects of D_1 -class receptor induction of pMeCP2. Because D_1 -class receptors are coupled through G_s to increases in intracellular cAMP whereas D_2 -class receptors oppose the increase in cAMP through G_i , we considered the possibility that differential regulation of pMeCP2 by distinct DA and 5-HT receptors could be integrated at the level of cAMP signaling. To test whether the elevation of cAMP is sufficient to induce pMeCP2 *in vivo*, we infused the adenylate cyclase activator forskolin directly into the striatum of C57BL/6J mice. All sections selected for staining had similar cannula placement (Figure 6a). Compared with saline infusion, forskolin robustly induced expression of the cAMPregulated immediate early gene product Fos in neurons below the tip of the cannula, demonstrating the efficacy of the drug treatment (p < 0.01) (Figure 6b and c). Similarly, we found that forskolin infusion was sufficient to induce pMeCP2 (p < 0.05) (Figure 6b and d). Unlike the sparse pattern of pMeCP2 induced by systemic AMPH administration (Figure 1), intra-striatal forskolin induced pMeCP2 in most of the cells near the cannula tip. This forskolininduced pMeCP2 colocalized in cells with DARPP-32, a marker of MSNs, and showed no overlap with GFAP, an abundantly expressed marker of glial cells (Figure 6e). Thus, these data show that intra-striatal forskolin delivery is sufficient to induce pMeCP2 in MSNs.

Forskolin infusion into the striatum could drive pMeCP2 by activating postsynaptic signaling cascades that directly induce MeCP2 phosphorylation in MSNs. Alternatively, because forskolin can act on presynaptic terminals to enhance the probability of neurotransmitter release (Chen and Regehr, 1997; Chavez-Noriega and Stevens, 1994) the induction of pMeCP2 in MSNs could be an indirect consequence of forskolin-induced release of neurotransmitter from the terminals of neurons that project to the striatum from other brain regions. To determine whether forskolin can act in MSNs to induce pMeCP2, we isolated the striatum from embryonic rat brains and treated



Figure 4 Stimulation of D₁- and D₂-class DA receptors induces different patterns of pMeCP2. (a) pMeCP2 immunostaining in the NAc 2 h after treatment with Veh, 7.5 mg/kg SKF81297, 0.4 mg/kg quinpirole, or coinjection of SKF81297 and quinpirole. (b) Quantitation of pMeCP2 immunostaining from (a). (c) pMeCP2 induction in the hippocampus 2 h after treatment with Veh, 5 or 7.5 mg/kg SKF81297, or coinjection of 7.5 mg/kg SKF81297 and quinpirole. Enlargement images show the dotted inset regions. (d) pMeCP2 (red), GAD67 (green), and overlay immunostaining in the NAc 2 h after treatment with 7.5 mg/kg SKF81297, 0.4 mg/kg quinpirole, or coinjection of SKF81297 and quinpirole. White arrows indicate neurons coimmunolabeled with antibodies for pMeCP2 and GAD67, and blue arrows indicate pMeCP2 induction in neurons that are not GAD67-positive GABAergic interneurons. n = 4-7 mice/group. *p < 0.05 compared with Veh, #p < 0.05 SKF compared with SKF + quinpirole. Error bars indicate SEM. Scale bar = 20 µm.

dissociated cultures of striatal neurons with either forskolin or the D₁-class DA receptor agonist SKF38393. Although these treatments induced phosphorylation of the protein kinase A (PKA) target and transcription factor CREB at Ser133, elevation of cAMP in cultured striatal neurons was not sufficient to induce pMeCP2 (Figure 6f). By contrast, both membrane depolarization and NMDA application induced pMeCP2 in these neurons in a manner that was dependent on the activation of L-type voltage-gated calcium channels and NMDA-type glutamate receptors, respectively (Figure 6g). We cannot rule out that the inability of forskolin to activate pMeCP2 in cultured MSNs is not a consequence of the immaturity of these cells. Nonetheless, taken all together, our data suggest that the differential ability of specific DA and 5-HT receptor agonists to drive pMeCP2 in MSNs (1) is integrated at the level of cAMP

signaling and (2) may depend, at least in part, on presynaptic cAMP-dependent effects.

AMPH-Induced pMeCP2 Is Reduced in SERT-KO Mice

The data presented thus far have addressed the sufficiency of activating specific monoamine neurotransmitter systems for induction of pMeCP2. However, to understand whether regulation of specific monoamine neurotransmitter systems is required for the induction of pMeCP2 following AMPH administration, we assessed the ability of AMPH to induce pMeCP2 in the NAc in strains of mice lacking expression of single monoamine transporters (Figure 7).

To determine whether AMPH can induce pMeCP2 in the NAc in the absence of the DAT, we administered either Veh or 3 mg/kg AMPH to DAT-KO mice and their DAT-WT

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Figure 5 Activation of 5-HT receptors by quipazine induces pMeCP2 in the NAc. (a) Quantitation of pMeCP2 integrated intensity in the NAc 2 h after treatment with Veh or 3 or 4.5 mg/kg quipazine. (b) pMeCP2 (red), GAD67 (green), and overlay immunostaining in the NAc 2 h after treatment with Veh or 3 mg/kg quipazine. White arrows indicate neurons coimmunolabeled with antibodies for pMeCP2 and GAD67. Values of quantitated immunofluorescence were normalized to Veh control. n = 4-6 mice/group. *p < 0.05 compared with Veh. Error bars indicated SEM. Scale bar = 20 µm.

littermate controls, monitored their activity in the open field for 1 h post-injection, and then quantified pMeCP2 in the NAc. A two-way ANOVA applied to locomotor activity 1 h post-injection revealed an overall main effect of genotype $(F_{3,22} = 74.516, p < 0.001),$ treatment $(F_{3,22} = 8.911, p < 0.008)$, and a significant genotype by treatment interaction ($F_{3,22} = 38.785$, p < 0.001). Consistent with previous reports (Gainetdinov et al, 1999; Giros et al, 1996), we found that DAT-KO mice had higher levels of baseline locomotor activity (p < 0.001) compared with the DAT-WT Veh controls and that their locomotor activity decreased following AMPH exposure (p < 0.001) compared with DAT-KO mice given Veh (Figure 7a). When an ANOVA was applied to pMeCP2 immunoreactivity, a main effect of treatment ($F_{3,22} = 17.709$, p < 0.001) was observed. However, there was no effect of genotype ($F_{3,22} = 2.082$, p < 0.166), and the genotype by treatment interaction was not significant ($F_{3,22} = 1,192, p < 0.289$). Bonferroni corrected pairwise comparisons noted that pMeCP2 levels were increased by AMPH in both genotypes compared with their Veh-treated controls (p < 0.033 and p < 0.002, respectively) (Figure 7b). However, the AMPH-induced levels of pMeCP2 did not differ between DAT-KO mice and their DAT-WT littermates (p < 0.807). Although the data did not reach

significance, DAT-KO mice showed a trend toward increased basal levels of pMeCP2 compared with DAT-WT mice (Figure 7b), which may arise as a consequence of the increased basal levels of extracellular DA in DAT-KOs (Gainetdinov *et al*, 1999). Microdialysis studies have shown that psychostimulants can increase extracellular DA levels in the NAc of DAT-KO mice through mechanisms that may involve blockade of the SERT or NET, although the functional relevance of a further increase in extracellular DA in face of the chronic hyperdopaminergia found in this strain remains unclear (Gainetdinov and Caron 2003; Carboni *et al*, 2001). Nonetheless, since AMPH can also bind the SERT and NET, we next asked whether these transporters are required for AMPH-induced pMeCP2.

To determine whether AMPH-induced pMeCP2 in the NAc is impaired in the absence of the SERT or NET, SERT-KO and NET-KO mice and their respective WT littermates were treated with Veh or AMPH, locomotor activity was monitored 1 h post-injection, and subsequently pMeCP2 was quantified in the NAc. For the SERT mice, a two-way ANOVA for locomotor activity revealed only a significant effect of treatment ($F_{3,23} = 38.262$, p < 0.001). Hence, locomotor activity was stimulated by AMPH to similar extent in both SERT-KO mice and their WT littermates (Figure 7c).



Figure 6 Forskolin is sufficient to induce pMeCP2 in the MSNs *in vivo* but not *in vitro*. (a) Anatomical location of microinjection sites in the striatum. (b) pMeCP2 and Fos immunostaining in the striatum 90 min after treatment with Veh or forskolin. (c, d) Quantitation of Fos and pMeCP2 integrated intensity in the striatum 90 min after treatment with Veh or forskolin. Values of quantitated immunofluorescence were normalized to Veh control. n = 9 mice/group. *p < 0.05 compared with Veh. Error bars indicated SEM. (e) Colocalization of pMeCP2 immunoreactivity with cell-type markers in the region at the tip of the cannula in sections from forskolin-treated mice. pMeCP2 (red), the MSN marker DARPP-32 (green, top), the astrocyte marker GFAP (green, bottom). Scale bar = 20 μ m. (f, g) Cultured embryonic rat striatal neurons (5 days *in vitro*) were treated for 60 min with various pharmacological reagents, the cells were lysed directly in boiling SDS and separated by SDS-PAGE, western blotting was performed using the antibodies indicated, and the blots were visualized with chemiluminescence on film. (f) Cultures were left either untreated (none), or treated with 55 mM KCl in an isotonic solution, 10 μ M forskolin (Fsk.) to elevate intracellular cAMP, or 50 μ M of the D₁-class DA receptor agonist SKF38393 (SKF). Western blots were probed for pMeCP2 and total MeCP2, or pCREB (Ser133) and total CREB as a control for the activation of cAMP signaling pathways. No protein was loaded in lane 2. (g) Cultures were left either untreated, or treated with 55 mM KCl or 50 μ M NMDA as indicated. In the lanes indicated, the L-type voltage-gated calcium channel inhibitor Nimodipine (Nim.) or the NMDA-receptor antagonist APV was added 2 min before stimulation with NMDA or KCl.

When changes in pMeCP2 were considered, the ANOVA detected significant main effects of genotype ($F_{3,23} = 5.542$, p < 0.029) and treatment ($F_{3,23} = 31.20$, p < 0.001), and a significant genotype by treatment interaction ($F_{3,23} = 5.467$, p < 0.03). Interestingly, Bonferroni comparisons demonstrated that AMPH-induced pMeCP2 was significantly reduced in SERT-KO mice compared with SERT-WTs

(p < 0.003), even though AMPH induced locomotor activity to the same extent in both genotypes (Figure 7d). Taken together with our pharmacological studies, the observation that AMPH-induced pMeCP2 in the NAc is significantly reduced in SERT-KO mice provides independent evidence of a role for 5-HT signaling in the regulation of pMeCP2 in the CNS.

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Figure 7 AMPH-induced pMeCP2 is reduced in the NAc of SERT-KO mice. (a, c, e) The effects of Veh or AMPH treatment on locomotor activity in DAT, SERT, and NET mice I h post-injection. (b, d, f) pMeCP2 induction in the NAc following Veh or AMPH treatment in DAT, SERT, and NET mice. All values of locomotor activity and quantitated immunofluorescence were normalized to the WT Veh control for each strain of mice. Veh treatment is indicated with a white bar, and AMPH treatment (3 mg/kg) is indicated with a dark gray bar. n = 6-8 mice/group. *p < 0.05 AMPH compared with Veh within the same genotype, *p < 0.05 WT Veh compared with mutant Veh, #p < 0.05 WT AMPH compared with mutant AMPH. (g) pMeCP2 (red) and GAD67 (green overlay immunostaining in the NAc 2 h after treatment with 3 mg/kg AMPH in the DAT-, SERT-, and NET-KO mice. White arrows indicate neurons coimmunolabeled with antibodies for pMeCP2 and GAD67. Error bars indicate SEM. Scale bar = 20 μ m.

With regards to NET-KO mice, a two-way ANOVA for locomotion revealed a significant effect of genotype ($F_{3,21} = 22.008$, p < 0.001), treatment ($F_{3,21} = 641.144$, p < 0.001), and significant treatment by genotype interaction ($F_{3,21} = 20.578$, p < 0.001). As previously reported (Xu *et al*, 2000), NET-KO mice were hyperresponsive to the locomotor-stimulating effects of acute AMPH administration compared with their WT littermates (p < 0.001) (Figure 7e). By contrast, an ANOVA for pMeCP2 immunoreactivity in the NAc found only a significant effect of treatment ($F_{3,21} = 104.690$, p < 0.001) (Figure 7f). Therefore, pMeCP2 was normally induced by AMPH in the absence of the NET.

Analysis of AMPH-stimulated pMeCP2 induction within the NAc revealed that in all genotypes tested, pMeCP2 remained selective for GAD67-positive GABAergic interneurons (Figure 7g). These data suggest that no single monoamine neurotransmitter system dictates the cell-type specificity of this pattern of pMeCP2 induction in the NAc. Furthermore, the evidence that genetic disruption of any single transporter failed to eliminate AMPH-induced pMeCP2 supports our pharmacological evidence that combinatorial signaling through multiple monoamine receptors drives the overall extent and pattern of pMeCP2 induced in the CNS.

pMeCP2 in the NAc Following Repeated AMPH Does Not Predict Behavioral Sensitization

Beyond providing a means to study the mechanistic actions of pharmacological agents, monoamine transporter KO mice have proven to be useful for investigating the AN Hutchinson et al

physiological functions of monoamine-regulated signaling pathways (Gainetdinov and Caron, 2003). For example, these mice have been used to gain insight into the mechanisms that underlie behavioral sensitization (Rocha, 2003; Xu et al, 2000), which is a progressive and persistent increase in behavioral responses to psychomotor stimulants that develops following their repeated administration (Pierce and Kalivas, 1997). Our previous data revealed a strong correlation between the magnitude of pMeCP2 induced in the NAc and the level of open field locomotor activity following repeated AMPH administration, which is one measure of behavioral sensitization (Deng et al, 2010). To test the hypothesis that pMeCP2 levels in the NAc predict behavioral sensitization to AMPH, we next asked whether pMeCP2 would correlate with locomotor activity following repeated AMPH administration in the NET-KO mice, which fail to sensitize (Xu et al, 2000).

NET-KO and their NET-WT littermates were injected with either Veh or AMPH once daily for 5 consecutive days in their home cages, then following a 7-day withdrawal they were administered an AMPH challenge in the open field to assess locomotor sensitization. The NET-KO mice were significantly more responsive to the locomotorstimulating effects of a single injection of AMPH in this paradigm than similarly treated WT littermates, as we have shown previously (Figure 7c). However, unlike their NET-WT littermates, the NET-KO mice failed to exhibit significantly enhanced locomotor activity following repeated AMPH administration (p < 0.0033 for WT and p < 0.12 for KO, VEH:AMPH compared with AMPH:AMPH) (Figure 8a and b). When changes in pMeCP2 were considered, ANOVA failed to detect significant effects of genotype ($F_{3,52} = 0.001$, p < 0.98) or treatment ($F_{3,52} = 0.063$, p < 0.803). pMeCP2 in the NAc was induced to similar



Figure 8 pMeCP2 in the NAc of NET-KO mice correlates with locomotor activity but not behavioral sensitization after repeated AMPH treatment. (a, b) The effects of acute (VEH:AMPH) or repeated (AMPH:AMPH) AMPH administration on locomotor activity in NET-WT and NET-KO mice I h post-injection on the challenge day. (c) pMeCP2 integrated intensity in the NAc following AMPH treatment on the challenge day in NET-WT and NET-KO mice. (d–g) Pearson product correlations of pMeCP2 immunofluorescence intensity in the NAc and cumulative locomotor activity (I h following the challenge injection) from individual mice. (h, i) Comparison of locomotor activity I h post-injection on day I and the challenge day (day 12) in NET-WT and NET-KO mice under the AMPH:AMPH condition. (j, k) Pearson product correlations of pMeCP2 immunofluorescence intensity in the NAc and the change in locomotor activity from day I to day I2 from individual mice in the NET-WT and NET-KO AMPH:AMPH groups. *p < 0.05 compared with VEH:AMPH. Error bars indicate SEM.

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extents in the two treatment groups (Veh:AMPH or AMPH:AMPH) for each genotype (Figure 8c). These data confirm that the lack of NET expression has no effect on the magnitude of pMeCP2 induction following either acute or repeated AMPH exposure, thus allowing us to analyze the correlation between induced pMeCP2 and locomotor activity in these mice.

Surprisingly, when a correlational analysis was applied to locomotor activity and NAc pMeCP2 induction in individual mice, a marked difference between the acute AMPH (VEH:AMPH) and repeated AMPH (AMPH:AMPH) treatment groups emerged for both NET genotypes (Figure 8d–g). No significant relationship between locomotor activity and pMeCP2 induction was observed in the VEH:AMPH group of either NET-WT (r=0.16, p=0.578) or NET-KO mice (r=0.25, p=0.379) (Figure 8d and f). By contrast, we observed a strong correlation of pMeCP2 immunoreactivity with locomotor activity in the AM-PH:AMPH treatment group of both genotypes (r=0.67, p=0.012; NET-WT AMPH:AMPH and r=0.71, p=0.007NET-KO AMPH:AMPH) (Figure 8e and g).

Although the level of locomotor activity after repeated AMPH can be used as an indication of behavioral sensitization, this metric does not reveal how the locomotor behavior of any single animal changed over the course of the repeated AMPH treatment. To obtain a more direct measure of locomotor sensitization in individual mice, we measured locomotor responses to AMPH in a different sensitization paradigm in which we monitored open field locomotor activity of each mouse on both the first day of AMPH treatment as well as on the challenge day. This experimental design allowed us to calculate a sensitization score for each mouse by subtracting the locomotor activity on the first day of AMPH treatment from the locomotor activity following AMPH exposure on the challenge day (Figure 8h and i). When a correlational analysis was applied to sensitization scores and NAc pMeCP2 levels in individual mice, no correlation was seen for either the NET-WT (r=0.373, p=0.467 AMPH:AMPH) or the NET-KO mice (r = 0.314, p = 0.492 AMPH:AMPH) (Figure 8j and k).

Taken together, these data replicate our previous observation that pMeCP2 levels correlate with open field locomotor activity only following repeated and not after acute AMPH. However, the levels of pMeCP2 in the NAc after repeated AMPH administration do not predict the degree of behavioral sensitization in individual mice, at least as measured by open field locomotor activity. An alternate hypothesis suggested by our data is that levels of locomotor activity following repeated AMPH exposure and pMeCP2 levels in the NAc may instead reflect other functional consequences of repeated AMPH treatment that persist in the absence of NET expression.

DISCUSSION

The net effect of AMPH interactions with monoamine transporters in the CNS is the activation of multiple monoaminergic receptor-coupled signaling pathways, culminating in the induction of intracellular processes including gene expression (Moratalla *et al*, 1996). Previously, we showed that AMPH administration induces Ser421

phosphorylation of MeCP2 in select populations of neurons within the NAc and PLC. Here, we have characterized the specific monoaminergic neurotransmitter pathways that regulate this phosphorylation event. Our data demonstrate that activation of both DA and 5-HT signaling is sufficient to induce pMeCP2. By contrast, we find no evidence for NE-dependent regulation of pMeCP2 even though three classes of NE receptors are expressed in the brain, several of which have been implicated in mediating certain aspects of behavioral responses to psychostimulants (Weinshenker and Schroeder, 2007; Xu *et al*, 2000; Laakso and Hietala, 2000). Although these data do not exclude roles for specific NE receptor-dependent regulation of pMeCP2 in other contexts, they do suggest that induction of pMeCP2 is selective for specific stimuli.

Interestingly, we find that the regional- and cell-type specific pattern of pMeCP2 induction depends on the particular combination of DA and 5-HT receptors activated. Consistent with the hypothesis that combinatorial signaling through multiple monoamine receptors sculpts the pattern of pMeCP2 in vivo, we find that genetic disruption of any single plasma membrane monoamine transporter in DAT-KO, SERT-KO, and NET-KO mice fails to completely block the ability of AMPH to induce pMeCP2 in the NAc and does not change the cell-type selectivity of the induced pattern. Parenthetically, VMAT2 is the other major target of AMPH in the CNS, and although VMAT2-KO mice are not viable after birth, we find that AMPH induces a similar magnitude and pattern of pMeCP2 in the NAc of VMAT2-HETs compared with their VMAT2-WT littermates (data not shown). Analysis of the distribution of pMeCP2 in the CNS following the various pharmacological stimuli used in this study also suggests that no single transporter or receptor expression pattern is sufficient to fully explain the selectivity of pMeCP2 induction. For example, the DAT is highly concentrated in DA-rich brain regions including the neostriatum, NAc, and olfactory tubercle (Ciliax et al, 1995). By contrast, the SERT and NET have more widespread distributions in the brain; high levels of SERT are found in the dorsal striatum, cortex, hippocampus, amygdala, and brainstem (Sur et al, 1996), whereas NET expression is concentrated in the PLC, bed nucleus of stria terminalis, and ventral regions of the striatum, including the NAc (Schroeter et al, 2000). However, only selective inhibition of the DAT induces pMeCP2 in the PLC, even though the SERT and NET are also highly expressed in this brain region (Miner et al, 2000, 2003). Taken together with the failure of either AMPH or the selective DAT inhibitor GBR12909 to induce pMeCP2 in MSNs of the NAc, which express high levels of DA receptors, these data suggest that either combinatorial patterns of transporter and receptor expression or additional mechanisms of specificity are important for the regulation of pMeCP2.

Our data suggest that one way different combinations of DA and 5-HT receptors influence the cell-type specific phosphorylation of MeCP2 is via the integration of receptor-induced signaling cascades at the level of cAMP. Both D_1 - and D_2 -class DA receptors as well as most types of 5-HT receptors are coupled by G-proteins to the regulation of adenylate cyclase and cAMP signaling (Barnes and Sharp, 1999; Missale *et al*, 1998). D_1 -class DA receptors elevate cAMP levels and we find that they drive widespread

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induction of pMeCP2 both in MSNs within the NAc and in other brain regions including the hippocampus. D2-class DA receptors inhibit cAMP production and we see that they oppose the D₁-class DA receptor-dependent induction of pMeCP2 in MSNs and hippocampal neurons. Elevation of cAMP levels leads to the activation of intracellular signaling cascades that can directly regulate nuclear signaling events, such as the induction of PKA-dependent Ser133 phosphorvlation on the transcription factor CREB (Yamamoto et al, 1988). However, MeCP2 Ser421 is not a direct target of phosphorylation by PKA (Zhou et al, 2006). Instead, calcium-calmodulin-activated kinases (CaMKII or CaM-KIV) have been identified as the likely MeCP2 Ser421 kinases following the activation of NMDA-type glutamate receptors and the opening of L-type voltage-gated calcium channels (Tao et al, 2009; Zhou et al, 2006). It is possible that DA and 5-HT receptors could induce pMeCP2 via the regulation of intracellular calcium levels. G_{q/11}-coupled DA and 5-HT receptors can release calcium from intracellular stores (Missale et al, 1998), and the 5-HT₃ receptor is unique in that it is a ligand-gated ion channel directly permeable to calcium (Derkach et al, 1989). Activation of D_1 -class DA receptors in cultured striatal neurons can elevate intracellular calcium levels indirectly through a mechanism that may involve DA- and PKA-dependent phosphorylation of the NR1 subunit of the NMDA-type glutamate receptor (Dudman et al, 2003). Stimulation of D₁class DA receptors also leads to PKA-dependent potentiation of L-type voltage-gated calcium channels in striatal neurons (Surmeier et al, 1995), and the subsequent activation of CaMKII by this pathway has been shown to be required for the regulation of AMPA-type glutamate receptor trafficking in the NAc in vivo in a cocaine-seeking paradigm (Anderson et al, 2008).

In addition to these postsynaptic sites of DA and 5-HT receptor action, our data raise the possibility that the induction of pMeCP2 in MSNs may be modulated by cAMPregulated processes in presynaptic terminals of afferent neurons that project to the striatum. Forskolin has been widely shown to enhance presynaptic vesicle release probability (Robbe et al, 2001; Chen and Regehr, 1997; Chavez-Noriega and Stevens, 1994). In striatal slice preparations, DA-dependent modulation of glutamate release from corticostriatal terminals allows for filtering of less active inputs and reinforcement of specific sets of corticostriatal synaptic connections (Bamford et al, 2004). Application of AMPH to this preparation inhibits presynaptic vesicle release from corticostriatal glutamatergic terminals in a manner that depends on D₂-class DA receptors (Bamford et al, 2004). Since NMDA-type glutamate receptors are robust activators of pMeCP2 in MSNs (Figure 6g), we propose that the inhibition of glutamate release by D₂-class receptors may underlie the ability of D₂-class receptor agonists to inhibit the D₁-class agonist induction of pMeCP2 in this population of neurons.

One commonality we observed among all of the DA and 5-HT reagents tested in this study is their ability to induce robust pMeCP2 in GAD67-positive GABAergic interneurons of the NAc. Although the signaling mechanisms that mediate this induction remain to be fully established, several lines of evidence suggest that this may reflect increases in firing of this cell population. In vivo recordings from presumed GAD67-positive striatal GABAergic interneurons in freely moving rats demonstrate that AMPH administration induces a rapid increase in the firing rate of most of these neurons (Wiltschko et al, 2010). D₂-class receptors appear to play an important facilitative role in this excitability because administration of the D₂-class receptor agonist eticlopride uniformly decreases the firing rate of these cells (Wiltschko et al, 2010). DA and 5-HT could act directly on GAD67-positive GABAergic interneurons to modulate their activity. Striatal GABAergic interneurons have been shown to express and be excited by D₁-class DA receptors; in slice preparations they are also activated indirectly through actions of presynaptic D₂-class receptors (Centonze et al, 2003; Bracci et al, 2002). Application of 5-HT to striatal slices also increases firing of GABAergic interneurons via activation of 5-HT_{2C} receptors (Blomeley and Bracci, 2009). Alternatively, DA and 5-HT could modulate properties of the afferent inputs to the striatum in a manner that changes the drive to GAD67-positive GABAergic interneurons. Excitatory input to GAD67positive GABAergic interneurons in the NAc comes from the prefrontal cortex (Gruber et al, 2009), while the major source of GABAergic inhibitory feedback onto these neurons comes from the globus pallidus (Bevan et al, 1998). D₂-receptor antagonists enhance the expression of Fos in the globus pallidus (Billings and Marshall, 2003), suggesting that increased firing of globus pallidus neurons induced by D₂ blockade could contribute to the suppression in striatal GABAergic interneuron firing described above (Wiltschko et al, 2010). Increased firing of GABAergic interneurons would lead to the opening of voltage-gated calcium channels, thus activating the CaM kinases that phosphorylate MeCP2. Although GABAergic interneurons are not thought to express the α isoform of CaMKII (Liu and Jones, 1996), the β , γ , and δ isoforms are widely expressed throughout the brain although at much lower levels than CaMKIIα (Sakagami and Kondo, 1993; Tighilet et al, 1998). Interestingly, a subset of hippocampal GABAergic interneurons have been shown to have an NMDA-receptordependent form of long-term potentiation that is blocked by CaM kinase inhibitors, suggesting the functional importance of non- α isoform-mediated CaMKII activity in these neurons (Lamsa et al, 2007).

Our behavioral data yield some insights into the functional implications of pMeCP2 induction. First, although the induction of pMeCP2 in the NAc is concurrent with the increase in locomotor activity following acute AMPH administration, pMeCP2 coincides with decreased locomotor activity following acute citalopram and with the paradoxical decrease in locomotor activity following AMPH administration in DAT-KO mice. Furthermore, following a single dose of AMPH, there is no correlation between the level of pMeCP2 induced in the NAc and the locomotor activity of an individual animal (Deng et al, 2010). Thus, pMeCP2 induction in the NAc reflects the effects of these drugs on the brain that are not directly involved in determining or sensing the level of locomotor activity. Interestingly, pMeCP2 in the NAc does correlate with locomotor activity following repeated AMPH treatment. Locomotor activity after repeated AMPH exposure is one measure of behavioral sensitization, which presumably occurs as a result of psychostimulant-induced neural circuit adaptations, although these remain poorly defined (Kauer and Malenka, 2007). However, pMeCP2 still correlates with locomotor activity following repeated AMPH treatment in NET-KO mice, which do not sensitize. Importantly, although the NET-KO mice do not sensitize to repeated AMPH administration, they do show other behavioral adaptations to repeated psychostimulant administration. For example, the NET-KO mice show preference for the cocaine-paired chamber in a CPP assay (Xu et al, 2000), and they self-administer cocaine in a dose-dependent manner (Rocha, 2003). Thus, we hypothesize that pMeCP2 induction may reflect the neural circuit activities that underlie these or other related behavioral adaptations to repeated AMPH exposure. Because the phosphorylation of pMeCP2 is selective for GABAergic interneurons of the NAc following AMPH and many of the other drugs tested here, it is tempting to speculate that pMeCP2 may play a role in plasticity of this neuronal population. It should be emphasized that the functions of GAD67-positive GABAergic striatal interneurons in behavioral responses to AMPH are entirely unknown. However, these cells exert robust feedforward inhibition over striatal output (Koos and Tepper, 1999), and changes in their firing patterns have been suggested to underlie the initiation of chosen actions while suppressing unwanted alternatives (Gage et al, 2010).

Finally, a novel finding of our study is the observation that the selective SERT inhibitor citalopram is sufficient to induce pMeCP2 in the CNS. 5-HT depletion has been shown to significantly reduce the inflammation-induced phosphorylation of MeCP2 in sensory neurons of the dorsal horn of the spinal cord (Geranton et al, 2008). Together with our findings, these data raise the possibility that serotonergic regulation of pMeCP2 could be important in a range of experimental paradigms and cell types. Drugs that target the SERT and 5-HT receptors are widely used in treating neuropsychiatric disorders such as depression (Krishnan and Nestler, 2008; Morilak and Frazer, 2004; Charney, 1998). In rodents, serotonergic regulation of affective behaviors has been shown in social-defeat stress and extended social isolation (Wallace et al, 2009; Tsankova et al, 2006), and drugs that alter serotonergic transmission modulate depressive-like behavior in these paradigms through mechanisms that are thought to involve changes in gene transcription (Vialou et al, 2010; Wallace et al, 2009; Tsankova et al, 2006; Nestler and Carlezon, 2006). Our evidence that a selective SERT inhibitor induces pMeCP2 in neural circuits that are functionally relevant to affective behaviors raises the possibility that this pathway could be involved in the mechanism of antidepressant action. From this perspective, investigating the regulation of pMeCP2 in the response to chronic antidepressant treatment may yield new insights into additional functional roles for MeCP2 in monoamine-regulated neural plasticity.

ACKNOWLEDGEMENTS

We thank MG Caron for generously sharing the DAT, NET, and VMAT2 mice and FS Hall for kindly supplying the SERT mice. We also thank RM Rodriguiz for technical assistance. Support for this work was provided by National Institute of Drug Abuse grants R01-DA022202 (AEW) and F32-DA025447 (JVD).

DISCLOSURE

The authors declare that this work was funded by NIH and that over the past three years WCW has received research funds from Abbott Laboratories, whose research objective was unrelated to the work contained within the present manuscript.

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