Disorders of the Nervous System

Single Dose of Amphetamine Induces Delayed Subregional Attenuation of Cholinergic Interneuron Activity in the Striatum

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Abstract

Psychostimulants such as amphetamine (AMPH) target dopamine (DA) neuron synapses to engender drug-induced plasticity. While DA neurons modulate the activity of striatal (Str) cholinergic interneurons (ChIs) with regional heterogeneity, how AMPH affects ChI activity has not been elucidated. Here, we applied quantitative fluorescence imaging approaches to map the dose-dependent effects of a single dose of AMPH on ChI activity at 2.5 and 24 h after injection across the mouse Str using the activity-dependent marker phosphorylated ribosomal protein S6 (p-rpS6240/244). AMPH did not affect the distribution or morphology of ChIs in any Str subregion. While AMPH at either dose had no effect on ChI activity after 2.5 h, ChI activity was dose dependently reduced after 24 h specifically in the ventral Str/nucleus accumbens (NAc), a critical site of psychostimulant action. AMPH at either dose did not affect the spontaneous firing of ChIs. Altogether this work demonstrates that a single dose of AMPH has delayed regionally heterogeneous effects on ChI activity, which most likely involves extra-Str synaptic input.

Key words: acetylcholine; dopamine; fluorescence imaging; phosphorylated ribosomal protein S6; psychostimulant

Significance Statement

Using the activity dependent marker phosphorylated ribosomal protein S6 (p-rpS6240/244), we mapped amphetamine (AMPH) effects on the activity of cholinergic interneurons (ChIs) across the striatum (Str). AMPH reduced ChI activity in dose-dependent manner in the ventral Str/nucleus accumbens (NAc), a critical site of psychostimulant action.

Introduction

Psychostimulants such as amphetamine (AMPH) target dopamine (DA) neuron terminals (Sulzer, 2011) and engender dose-dependent behavioral effects. DA release in the ventral striatum/nucleus accumbens (Str/NAc) is associated with hyperlocomotion, whereas DA release in the dorsal Str is associated with stereotypies (Robinson and Becker, 1986; Kalivas and Stewart, 1991; Gaytan et al., 1998; Yates et al., 2007). DA neurons modulate the activity of cholinergic interneurons (ChIs), which comprise <2% of striatal (Str) neurons, and yet strongly control the Str circuitry (Goldberg and Wilson, 2010; Gonzales and Smith, 2015; Abdukeyoumu et al., 2019). Modulation of ChI activity is critical for the processing and reinforcement of reward-related behaviors (Atallah et al., 2014; Gonzales and Smith, 2015). ChIs in the
ventral Str are crucial for psychostimulant-dependent behaviors (Sofuoglu and Mooney, 2009; Witten et al., 2010; Lee et al., 2020; Lewis and Borrelli, 2020). However, whether AMPH has subregional effects on ChI activity has not been elucidated.

Previous studies have shown that the phosphorylated form of the ribosomal protein S6 at serine 240 and 244 residues (p-rpS6^{240/244}) reports activity of Chls under different pharmacological and/or behavioral conditions (Bertran-Gonzalez et al., 2012; Kharkwal et al., 2016; Matamales et al., 2016a,b). The phosphorylation of rpS6 can be induced by multiple signaling cascades; mTORC1 pathway and/or mTORC1-independent pathways such as the PKC, the MAPK or the cAMP/PKA pathways (Valjent et al., 2011; Bertran-Gonzalez et al., 2012; Gangarossa and Valjent, 2012). The phosphorylation of rpS6 appears to occur sequentially at five serine residues: in the order 236, 235, 240, 244, and 247 (Knight et al., 2012; Biever et al., 2015a). Bertran-Gonzalez and colleagues showed a clear p-rpS6^{240/244} signal preferentially expressed in Chls, in contrast to a much weaker signal of p-rpS6^{235/236} (Bertran-Gonzalez et al., 2012). Pharmacological modification of ChI firing leads to changes of p-rpS6^{240/244} intensity in Chls (Bertran-Gonzalez et al., 2012; Matamales et al., 2016b). To address regionality in AMPH modulation of ChI activity, we mapped p-rpS6^{240/244} intensity in Chls throughout the entire rostrocaudal axis of the Str after a single low-dose or high-dose of AMPH at two time points: 2.5 h postinjection (2.5hpi) and 24 h postinjection (24hpi). This revealed that AMPH induces a delayed regionally heterogeneous dose-dependent attenuation of ChI activity in the ventral Str/NAc.

Materials and Methods

Ethics

This research was performed in accordance with the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health, under a protocol approved by the Institutional Animal Care and Use Committee of New York State Psychiatric Institute (#NYSPI-1494).

Experimental animals

Mice were 129 Sv/C57BL6J mixed background, backcrossed to C57BL6J at least five times and kept inbred. Mice were group housed and maintained on a 12/12 h light/dark cycle with lights on at 7 A.M. in a temperature-controlled room with food and water provided ad libitum. The DAT-IRES-Cre/++;ROSA26-flox-STOP-CAG-ChR2-YFP double mutant strain (The Jackson Laboratory, RRID: IMSR_JAX:006660, RRID:IMSR_JAX:024109) were used, with the same genotype as previous studies (Chuhma et al., 2014, 2018; Mingote et al., 2015, 2017). The presence of Cre is not essential for the present study; the IRES-cre transgene insertion in the DA transporter (DAT) locus modestly reduces DAT expression and AMPH responsiveness (Bäckman et al., 2006; Chohan et al., 2020).

For the immunocytochemistry experiments, 30 mice were used at postnatal day (P)56–P82, divided in two cohorts of 15 for 2.5hpi and 15 for 24hpi. Cohorts were balanced for sex: 16 male (2.5hpi cohort: saline, n = 3; low-dose AMPH, n = 3; high-dose AMPH, n = 3 and 24hpi cohort: saline, n = 2; low-dose AMPH, n = 2; high-dose AMPH, n = 3) and 14 female (2.5hpi cohort: saline, n = 2; low-dose AMPH, n = 2; high-dose AMPH, n = 2 and 24hpi cohort: saline, n = 3; low-dose AMPH, n = 3; high-dose AMPH, n = 2) mice. For the electrophysiological experiments, 40 male (saline, n = 17; low-dose AMPH, n = 11; high-dose AMPH, n = 12) and 40 female (saline, n = 13; low-dose AMPH, n = 11; high-dose AMPH, n = 16) mice at P52–P72 were used. No sex differences were observed, so data from male and female mice in each group were combined.

Drug treatment

D-AMPH hemisulfate (Sigma-Aldrich, A5880) either low-dose (2 mg/kg) or high-dose (16 mg/kg) was dissolved in 0.9% NaCl immediately before use. Injections were done intraperitoneally at a volume of 10 ml/kg body weight.

Behavioral monitoring

Mice were habituated to handling for 2 d before the drug administration. Monitoring took place under bright ambient light conditions during the light phase. On the injection day, mice were placed in the open field, equipped with infrared motion detectors (Plexiglas activity chambers, 40.6 cm long x 40.6 cm wide x 38.1 cm high; SmartFrame Open Field System, Kinder Scientific) for 1 h for habituation. Baseline activity was monitored for 30 min preinjection, then mice were injected intraperitoneally either with saline, 2 or 16 mg/kg AMPH, and observed for a 2-h postinjection period. Locomotor activity was recorded automatically in 10-min bins. Stereotyped behaviors, orofacial stereotypy (mouth movements, lick, bite, self-gnaw, taffy pull, jaw tremor, yawn) and grooming, were scored for 1 min every 5 min as previously described (Kelley, 2001).

Immunocytochemistry

For immunocytochemistry, mice were deeply anesthetized with ketamine (90 mg/kg)/xylazaine (7 mg/kg) and then perfused intracardially with cold PBS (100 mm; pH 7.4) followed by 4% paraformaldehyde (PFA). Brains were removed and postfixed overnight in 4% PFA. Coronal

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sections were cut at 50 μm with a vibrating microtome (Leica VT1200S), and stored in a cryoprotectant solution (30% glycerol, 30% ethylene glycol in 0.1 M Tris HCl, pH 7.4) at −20°C. Free-floating sections were washed in PBS and incubated in glycine (100 mM) for 30 min to quench aldehydes. Non-specific binding was blocked with 10% normal donkey serum (NDS) in 0.1 PBS Triton X-100 for 2 h. Sections were incubated in PBS containing 0.02% Triton X-100 and 2% NDS overnight at 4°C with primary antibodies: anti-ChAT (1:500, goat polyclonal, Millipore catalog #AB144P, RRID:AB_2079751) and anti-phosphorylated ribosomal protein S6 (p-rpS6240/244, 1:1500, rabbit polyclonal, Cell Signaling Technology catalog #2215, RRID:AB_331682). Sections were then washed with PBS, and secondary antibodies applied for 45 min in PBS containing 0.02% Triton X-100 at room temperature: anti-goat Alexa Fluor 594 (1:200, Thermo Fisher Scientific catalog #A-11 058, RRID:AB_2534105) and anti-rabbit Alexa Fluor 488 (1:200, Thermo Fisher Scientific catalog #A-21206, RRID:AB_2535792). Sections were mounted on gelatin subbed slides (SouthernBiotech) and coverslipped with ProLong Gold aqueous medium with DAPI (Thermo Fisher Scientific) and stored at 4°C until imaging.

Imaging and analysis Images were acquired using an Axio Imager M2 fluorescence microscope (Zeiss) with a high-resolution digital camera (AxioCam 506 mono, 2752 × 2208 pixels, Zeiss), a 20×/0.8 objective and Zen 2.3 Digital Imaging software (Zeiss; RRID:SCR_013672). Ten coronal sections, spanning the rostrocaudal extent of the right Str (bregma 1.54, 1.18, 0.98, 0.62, 0.26, −0.10, −0.46, −0.82, −1.22, and −1.58 mm), were imaged. An image stack consisting of 5 planes at 5 μm intervals was obtained. Exposure time for each excitation was held constant throughout acquisition. Raw 16-bit images were analyzed using Fiji/ImageJ (version 2.0.0., NIH, RRID:SCR_002285). Z-projected images were obtained by taking pixels with the maximum intensity in a stack. The outer boundary of the Str and its intracellular solution was 135 mM K+ (World Precision Instruments). Pipette resistance was 4–5 MΩ and series resistance was 7–10 MΩ. Composition of intracellular solution was 135 mM K+ -methylene sulphonate (MeSO4), 5 mM KCl, 2 mM MgCl2, 0.1 mM CaCl2, 10 mM HEPES, 1 mM EGTA, 2 mM ATP, and 0.1 mM GTP; pH 7.25. Recording was done with an Axopatch 200B amplifier (Molecular Devices) in fast current clamp mode. All recordings were done at 32–34°C (TC 344B Temperature Controller, Warner Instruments). No more than four cells were recorded per animal. Data were filtered at 5 kHz using a four-pole Bessel filter, digitized at 5 kHz (Digidata 1550A, Molecular Devices) and recorded using pClamp 10 (Molecular Devices; RRID:SCR_011323). Electrophysiological data were analyzed with Axograph X (Axograph Science; RRID:SCR_014284). Firing frequencies were calculated as average frequency in a 2 s window obtained from 10 consecutive traces.

Statistical analysis Sample sizes were determined using G’Power 3.1 with effect sizes based on similar experiments (G’Power, 2016).

The p-rpS6240/244 fluorescence intensity were standardized to the corresponding saline group for each time point and subgroup by calculating z-scores: z = (x − μ)/σ, where x is the p-rpS6240/244 signal in individual ChIs, μ and σ are the mean and the SD, respectively, of p-rpS6240/244 signal in the corresponding saline group.

Slice electrophysiology and analysis For electrophysiology recording, mice were anesthetized with ketamine (90 mg/kg)/xylazine (7 mg/kg). After confirmation of full anesthesia, mice were decapitated and brains quickly removed in ice-cold high-glucose artificial CSF (ACSF; 75 mM NaCl, 2.5 mM KCl, 25 mM NaHCO3, 1.25 mM NaH2PO4, 0.7 mM CaCl2, 2 mM MgCl2, and 100 mM glucose; pH 7.4) saturated with carbogen (95% O2 + 5% CO2). Coronal sections of the Str (bregma from 1.70 to 0.26 mm) were cut, 300 μm thick, with a vibrating microtome (VT1200S, Leica), incubated in high-glucose ACSF at room temperature for at least 1 h for recovery, then transferred to the recording chamber (submerged, 500 µl volume) on the stage of an upright microscope (BX61WI, Olympus), continuously perfused with standard ACSF (125 mM NaCl, 2.5 mM KCl, 25 mM NaHCO3, 1.25 mM NaH2PO4, 2 mM CaCl2, 1 mM MgCl2, and 25 mM glucose; pH 7.4) saturated with carbogen. Recorded neurons were visualized using enhanced visible light differential interference contrast (DIC) optics with a scientific c-MOS camera (ORCA-Flash4.0LT, Hamamatsu Photonics).

Chls were identified visually by large soma size, confirmed by spontaneous firing, shallow resting membrane potentials (around 60 mV) and voltage sag by −400 pA current injection (700 ms in duration; Chuhma et al., 2014, 2018). Recording patch pipettes were fabricated from standard-wall borosilicate glass capillary with filament (World Precision Instruments). Pipette resistance was 4–9 MΩ and series resistance was 7–32 MΩ. Composition of intracellular solution was 135 mM K+ -methylene sulphonate (MeSO4), 5 mM KCl, 2 mM MgCl2, 0.1 mM CaCl2, 10 mM HEPES, 1 mM EGTA, 2 mM ATP, and 0.1 mM GTP; pH 7.25.Recording was done with an Axopatch 200B amplifier (Molecular Devices) in fast current clamp mode. All recordings were done at 32–34°C (TC 344B Temperature Controller, Warner Instruments). No more than four cells were recorded per animal.
or high-dose (16 mg/kg, per animal.

Figures 1A and 1B illustrate the timeline of AMPH experiments and the total distance traveled (left), orofacial stereotypy (middle), and grooming (right) scores shown after saline (0 mg/kg, n = 5 animals), low-dose (2 mg/kg, n = 5 animals), or high-dose (16 mg/kg, n = 5 animals) AMPH, for 2.5h cohort (red) and 24h cohort (blue). Dots in bar graphs show measurements per animal. *p < 0.05, **p < 0.01, ***p < 0.001 for comparison among doses; ○ p < 0.05 for comparison between 2.5h and 24h.

Statistical analyses were performed using Prism 8 (GraphPad Prism, RRID:SCR_002798) or SPSS 26 (SPSS; RRID:SCR_002865). p < 0.05 was considered as significant for all analyses. Data are presented as mean ± SEM. Parametric tests were used here because datasets followed a normal distribution (D’Agostino–Pearson normality test, p > 0.05). ANOVA was used for comparison among conditions. Where significance was detected, multiple pairwise comparisons with Bonferroni correction were performed as post hoc tests.

Results

Dose-dependent effects of AMPH on locomotor activity and stereotypy

Behavioral observations were used to confirm the dose-dependent effects of AMPH. Mice received a single low-dose (2 mg/kg) or high-dose (16 mg/kg) of AMPH, and their brains were extracted for analysis either after 2.5h, when acute behavioral effects had subsided, or at 24h to assess enduring effects on ChI activity in the Str. One low-dose AMPH-injected mouse, in the 2.5h cohort, was excluded from the study as its locomotor activity decreased after injection.

To confirm differential behavioral effects of the two AMPH doses and similar behavioral effects in the two cohorts (2.5h and 24h), mice from each cohort received saline, low-dose or high-dose AMPH, and locomotion and stereotypy were monitored for 2 h in the open field (Fig. 1A). Total travel distance dose dependently increased in both the 2.5h cohort (saline 17.6 ± 3.4 m, low-dose 94.5 ± 13.8 m, high-dose 212.5 ± 29.3 m) and 24h cohort (saline 18.6 ± 3.4 m, low-dose 64.0 ± 5.0 m, high-dose 302.7 ± 35.8 m), while no significant difference was observed between the two cohorts (two-way ANOVA; treatment effect, F(2,24) = 78.15, p < 0.001; cohort effect, F(1,24) = 1.55, p = 0.23; Fig. 1A, left). Although there was a significant treatment × cohort interaction (F(2,24) = 4.95, p = 0.02), the two cohorts showed similar dose-dependent hyperlocomotion, a significant increase after low-dose and a further increase after high-dose.

Low-dose and high-dose AMPH increased orofacial stereotypy in both the 2.5h cohort (saline 2.0 ± 1.3, low-dose 2.2 ± 0.6, high-dose 17.6 ± 3.6) and 24h cohort (saline 0 ± 0, low-dose 8.0 ± 1.9, high-dose 20.2 ± 2.9), while no significant difference was observed between the two cohorts (two-way ANOVA; treatment effect, F(2,24) = 38.74, p < 0.001; cohort effect, F(1,24) = 1.50, p = 0.23; treatment × cohort interaction, F(2,24) = 1.69, p = 0.21; Fig. 1B, middle).

Low-dose AMPH did not affect grooming score in the two cohorts, while high-dose increased it in both 2.5h cohort (saline 9.6 ± 2.5, low-dose 6.6 ± 1.9, high-dose 16.2 ± 1.5) and 24h cohort (saline 5.8 ± 0.9, low-dose 7.8 ± 2.4, high-dose 18.8 ± 1.2; two-way ANOVA; treatment effect, F(2,24) = 19.86, p < 0.001). Neither a significant difference, nor a treatment × cohort interaction, was observed between the two cohorts (cohort effect, F(1,24) = 0, p > 0.99; treatment × cohort interaction, F(2,24) = 1.67, p = 0.21; Fig. 1B, right). These observations confirmed that AMPH elicited a comparable dose-dependent behavioral activation in the two cohorts, used for the 2.5h and 24h ChI studies.

RRID: SCR_013726, setting α = 0.05 and power = 0.8 (Cunningham and McCrum-Gardner, 2007; Faul et al., 2007). For the immunocytochemistry experiments, we used Cohen’s d = 0.97 as an effect size, resulting in 5 mice per group. For the electrophysiological experiments, we used Cohen’s d = 0.32 as an effect size, resulting in 12 mice per group.

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Distribution and morphology of CHIs is not affected by AMPH

To address potential neurotoxic effects of AMPH on CHIs, particularly high-dose (viz., Zhu et al., 2006), we examined CHI distribution and soma morphology. CHIs were identified by ChAT immunostaining and examined in 10 coronal sections spanning the rostro-caudal extent of the right Str in four subregions: NAc core and shell, DM and DL Str (Fig. 2A). The previously recognized rostro-caudal distribution of CHIs (Matamalos et al., 2016a) peaked at 0.98 mm from bregma and gradually declined caudally. The distribution was not affected by either AMPH dose or time after injection (three-way ANOVA; rostro-caudal effect, $F_{(6,240)}=204.13, p < 0.001$; treatment effect, $F_{(2,240)}=0.17, p = 0.85$; time effect, $F_{(1,240)}=0.66, p = 0.42$; rostro-caudal $\times$ treatment $\times$ time interaction, $F_{(18,240)}=1.43, p = 0.12$; Fig. 2B). Although numbers of CHIs varied significantly between Str subregions, AMPH dose or time after injection did not affect CHI count significantly in any Str subregion (three-way ANOVA; location effect, $F_{(6,96)}=954.82, p < 0.001$; treatment effect, $F_{(2,96)}=0.12, p = 0.89$; time effect, $F_{(1,96)}=0.42, p = 0.52$; location $\times$ treatment $\times$ time interaction, $F_{(6,96)}=0.52, p = 0.80$; Fig. 2C, top). Although CHI densities varied between Str subregions, highest in the NAc shell and lowest in the NAc core, AMPH dose or time after injection did not affect densities in any subregion (three-way ANOVA; location effect, $F_{(3,96)}=73.76, p < 0.001$; treatment effect, $F_{(2,96)}=0.15, p = 0.86$; time effect, $F_{(1,96)}=0.57, p = 0.45$; location $\times$ treatment $\times$ time interaction, $F_{(6,96)}=0.30, p = 0.94$; Fig. 2C, bottom).

We examined the shape of CHIs based on their cytoplasmic ChAT immunoreactivity. AMPH did not affect CHI soma area (2.5hpi: saline 242.5 $\pm$ 4.4 $\mu$m$^2$, low-dose 243.6 $\pm$ 10.5 $\mu$m$^2$, high-dose 240.4 $\pm$ 9.6 $\mu$m$^2$; 24hpi: saline 242.6 $\pm$ 14.6 $\mu$m$^2$, low-dose 250.1 $\pm$ 3.0 $\mu$m$^2$, high-dose 242.5 $\pm$ 5.5 $\mu$m$^2$; two-way ANOVA; treatment effect, $F_{(2,24)}=0.21, p = 0.81$; time effect, $F_{(1,24)}=0.16, p = 0.69$; treatment $\times$ time interaction, $F_{(2,24)}=0.07, p = 0.94$), perimeter (2.5hpi: saline 72.3 $\pm$ 1.4 $\mu$m, low-dose 74.3 $\pm$ 3.4 $\mu$m, high-dose 75.3 $\pm$ 3.1 $\mu$m; 24hpi: saline 76.3 $\pm$ 1.1 $\mu$m, low-dose 76.6 $\pm$ 1.4 $\mu$m, high-dose 76.3 $\pm$ 1.4 $\mu$m; two-way ANOVA; treatment effect, $F_{(2,24)}=0.23, p = 0.80$; time effect, $F_{(1,24)}=1.14, p = 0.24$; treatment $\times$ time interaction, $F_{(2,24)}=0.29, p = 0.75$) or circularity (2.5hpi: saline 0.60 $\pm$ 0.02, low-dose 0.59 $\pm$ 0.04, high-dose 0.65 $\pm$ 0.03; 24hpi: saline 0.56 $\pm$ 0.04, low-dose 0.58 $\pm$ 0.04, high-dose 0.55 $\pm$ 0.04; two-way ANOVA; treatment effect, $F_{(2,24)}=0.28, p = 0.76$; time effect, $F_{(1,24)}=0.56, p = 0.46$; treatment $\times$ time interaction, $F_{(2,24)}=0.07, p = 0.93$) in the whole Str, at any time points (Fig. 3A).

Although CHI morphology differed between Str subregions, AMPH did not affect soma area, perimeter, or circularity in any Str subregion or at the two different time points (area: three-way ANOVA; location effect, $F_{(3,96)}=13.17, p < 0.001$; treatment effect, $F_{(2,96)}=0.37, p = 0.68$; time effect, $F_{(1,96)}=0.31, p = 0.58$; location $\times$ treatment $\times$ time interaction, $F_{(6,96)}=0.18, p = 0.98$; perimeter: three-way ANOVA; location effect, $F_{(3,96)}=22.45, p < 0.001$; treatment effect, $F_{(2,96)}=0.38, p = 0.68$; time effect, $F_{(1,96)}=2.42, p = 0.12$; location $\times$ treatment $\times$ time, $F_{(6,96)}=0.83, p = 0.55$; circularity: three-way ANOVA; location effect, $F_{(3,96)}=7.69, p < 0.001$; treatment effect, $F_{(2,96)}=0.58, p = 0.57$; time effect, $F_{(1,96)}=1.36, p = 0.25$; location $\times$ treatment $\times$ time interaction, $F_{(6,96)}=0.35, p = 0.91$; Fig. 3B). Thus, neither low- nor high-dose AMPH affected CHI distribution or morphology, arguing against neurotoxic effects of a single dose of AMPH.

AMPH attenuation of CHI activity in vivo

We mapped AMPH effects on CHI activity using p-rpS6240/244 as a reporter. Double immunostaining showed colocalization of ChAT and p-rpS6240/244 (Fig. 4). P-rpS6240/244 signal was also present in other Str cells, so ChAT staining was used to extract the signal specifically deriving from CHIs. We quantified p-rpS6240/244 intensity as the average pixel intensity in each ChAT-positive neuron, in sections from the saline, low-dose and high-dose AMPH-injected mice, at 2.5hpi or 24hpi (n = 5 animals/treatment, 10 hemisections/animal). Individual ChI locations were plotted in coronal hemisections of the Str and p-rpS6240/244 intensities were color-scaled (Fig. 5A, C).

At 2.5hpi, p-rpS6240/244 intensity varied among Str subregions, with higher p-rpS6240/244 intensity in the DM Str decreasing ventrally to the NAc shell (Fig. 5A). There was no apparent difference in the distribution of p-rpS6240/244 intensity between saline, low-dose, and high-dose AMPH-injected animals (Fig. 5A). Average p-rpS6240/244 intensity in the whole Str did not differ among saline, low-dose and high-dose AMPH-injected animals (one-way ANOVA, $F_{(2,12)}=0.003, p = 0.99$; Fig. 5B, left). Although average CHI p-rpS6240/244 intensities differed in Str subregions, neither low- nor high-dose AMPH affected CHI p-rpS6240/244 intensity in any Str subregion at 2.5hpi (two-way ANOVA; treatment effect, $F_{(2,48)}=0.62, p = 0.54$; location effect, $F_{(3,48)}=41.28, p < 0.001$; treatment $\times$ location interaction, $F_{(6,48)}=0.66, p = 0.68$; Fig. 5B, right).

At 24hpi, CHI p-rpS6240/244 intensity was reduced by low-dose AMPH particularly in the NAc (Fig. 5C). Low-dose AMPH reduced p-rpS6240/244 staining more than high-dose (Fig. 5C). Indeed, low-dose AMPH reduced average CHI p-rpS6240/244 staining in the whole Str, while high-dose AMPH did not show a significant effect (one-way ANOVA, $F_{(2,12)}=4.35, p = 0.03$; Fig. 5D, left). Low-dose AMPH significantly reduced CHI p-rpS6240/244 intensity in the NAc core ($p = 0.043$) and NAc shell ($p = 0.047$), but not in the dorsal Str (DM Str, $p = 0.46$; DL Str, $p = 0.40$; Fig. 5D, right; two-way ANOVA; treatment effect, $F_{(2,48)}=8.56, p < 0.001$; location effect, $F_{(3,48)}=0.20, p = 0.89$; treatment $\times$ location interaction, $F_{(6,48)}=0.35, p = 0.91$). High-dose AMPH does not affect CHI p-rpS6240/244 intensity in any Str subregion.

To compare AMPH effects on p-rpS6240/244 intensity between the two time points, p-rpS6240/244 intensities in CHIs were standardized to the respective saline groups and the differences expressed as z-scores for each Str subregion. Z-scores in AMPH-injected animals at 2.5hpi showed no difference from saline-injected animals in any Str subregion (two-way ANOVA; treatment effect $F_{(2,48)}=0.70, p = 0.50$; location effect $F_{(3,48)}=1.98, p = 0.13$; Fig. 5D, right).
Figure 2. Distribution of Chls in the Str is not affected by AMPH. A, Schematic representations of 10 coronal sections of the Str (from bregma +1.54 to −1.58 mm). Delineations of Str subregions are shown in the right Str (Paxinos and Franklin, 2008): NAc core
Figure 3. Morphology of Chls is not affected by AMPH. A–B, Morphologic characteristics of Chls in the whole Str (A) and each Str subregion (B) in the same hemisections as shown in the previous figure: area (µm²), perimeter (µm), and circularity after saline (0 mg/kg), low-dose (2 mg/kg), or high-dose (16 mg/kg) AMPH, at 2.5hpi (top) and 24hpi (bottom). Dots in bar graphs show the average measurements per animal; *p < 0.05, **p < 0.01, ***p < 0.001.

C, Total Chl count (top) and Chl density (neurons/mm²; bottom) in each Str subregion are shown, at 2.5hpi (left) and 24hpi (right). Group ns are given in Figure 1. Dots in bar graphs show the average per animal; **p < 0.01 and ***p < 0.001.
treatment \times location interaction, $F_{(6,48)} = 0.65, p = 0.69; \text{ Fig. 6A}$. At 24hpi, p-rpS6$^{240/244}$ intensity z-scores became negative after low-dose or high-dose AMPH in all Str subregions, indicating a reduction in ChI activity (two-way ANOVA; treatment effect $F_{(2,48)} = 8.97, p < 0.001$; location effect $F_{(3,48)} = 0.54, p = 0.66$; treatment \times location interaction, $F_{(6,48)} = 0.49, p = 0.81; \text{ Fig. 6B}$). Low-dose AMPH significantly attenuated ChI p-rpS6$^{240/244}$ intensity z-scores in the ventral subregions: NAc core ($p = 0.012$) and shell ($p = 0.048$), but not in the dorsal Str (DM Str, $p = 0.50$; DL

Figure 4. Phosphorylation of ribosomal protein S6 (p-rpS6$^{240/244}$) in Chls. A, Low-magnification images of ChAT (purple) and p-rpS6$^{240/244}$ (green) in a Str hemisection (bregma 0.98 mm) with merged images in the middle. Colored rectangles are representative locations of Str subregions and magnified in B–E. Expanded images of the NAc core (B, orange), NAc shell (C, magenta), DM Str (D, green), and DL Str (E, blue) subregions.
Figure 5. P-rpS6\textsuperscript{240/244} intensity in Chls 2.5h\textsubscript{pi} and 24h\textsubscript{pi} AMPH. A, C, Spatial distribution of Chls with relative p-rpS6\textsuperscript{240/244} intensity in the same 10 hemisections as shown in the previous morphology figures along the rostrocaudal axis (from bregma –1.54 to –1.58 mm), at 2.5h\textsubscript{pi} (A) and 24h\textsubscript{pi} (C). Spatial distribution of Chls from five animals was superimposed for each injection group: saline (top), low-dose (middle), or high-dose (bottom) AMPH. Each dot represents one Chl and intensity of p-rpS6\textsuperscript{240/244} is shown on a blue (low level) to red (high level) color scale. B, D, left, Average p-rpS6\textsuperscript{240/244} intensity in Chls in the whole Str at 2.5h\textsubscript{pi} (B) and 24h\textsubscript{pi} (D) after saline (0 mg/kg), low-dose (2 mg/kg), or high-dose (16 mg/kg) AMPH. Right, Box and whiskers plots showing p-rpS6\textsuperscript{240/244} intensity in Chls in each Str subregion at 2.5h\textsubscript{pi} (B) and 24h\textsubscript{pi} (D). Dots in bar graphs show the average measurements per animal; *p < 0.05.
spontaneous firing of ChIs in slices in the four Str subregions after saline, low-dose or high-dose AMPH at 24h$_{pi}$ (Fig. 7A). ChIs were identified visually by large soma size, confirmed by spontaneous firing and voltage sag in response to hyperpolarizing-current injection (Fig. 7B), as described previously (Chuhma et al., 2014). Although firing frequencies of ChIs varied significantly among Str subregions, AMPH did not affect firing frequencies of ChIs in any Str subregion (two-way ANOVA; treatment effect, $F_{(2,134)} = 1.21$, $p = 0.30$; location effect, $F_{(3,134)} = 13.30$, $p < 0.001$; treatment × location interaction, $F_{(6,134)} = 1.12$, $p = 0.36$; Fig. 7C). Thus, neither low- nor high-dose AMPH affected the intrinsic firing of ChIs in the deafferented slice, at 2.5h$_{pi}$ or 24h$_{pi}$, suggesting AMPH effects on ChI activity are because of extra-Str synaptic input.

**Discussion**

ChIs are principal targets of DA neurons and subject to regionally heterogeneous modulation. Here, we mapped the downstream effects of a single AMPH dose on ChI activity using p-rpS6$^{240/244}$ as a ChI-preferential activity-dependent marker. The single dose of AMPH did not affect the distribution, overall morphology, or spontaneous firing of ChIs in any Str subregion, arguing against neurotoxic effects of AMPH. While AMPH had no effect on *in vivo* ChI activity at 2.5h$_{pi}$, it significantly attenuated ChI activity at 24h$_{pi}$ in the ventral Str/NAc. In the NAC, the attenuation in ChI activity after low-dose was greater than after high-dose. In the dorsal Str, no significant difference in ChI activity was observed after either low-dose or high-dose AMPH. Thus, a single dose of AMPH has delayed regionally heterogeneous effects on ChI activity, with a dose-dependency in the NAC.

**Distribution, morphology, and spontaneous firing of ChIs in the Str**

In rodents (Gonzales and Smith, 2015), non-human primates (Brauer et al., 2000) and humans (Holt et al., 1996), the average size of ChIs in the NAC is smaller than in the dorsal Str. Here, we found that ChIs in the NAC core were significantly smaller and more elongated compared with those in other Str subregions, and that the morphology of ChIs soma (area, perimeter and circularity) differed among Str subregions. We also confirmed the differential distribution of ChIs in Str subregions (Gonzales and Smith, 2015). ChIs are denser in the NAC medial shell, as previously described in mice (Matamales et al., 2016a), rats (Phelps and Vaughn, 1986), and primates (Brauer et al., 2000).

A single injection of AMPH, either low-dose or high-dose, did not affect ChI distribution or soma morphology in any Str subregion, at either time point, showing these doses were not neurotoxic. Although AMPH neurotoxicity on DA neurons has been known for some time (Wagner et al., 1980; Ricaurte et al., 1984; Ryan et al., 1990; Miller and O’Callaghan, 1996; Krasnova et al., 2001, 2005; Granado et al., 2018), no study has focused on downstream neurotoxic effect on Str ChIs. To cause a significant toxic effect on ChIs, a higher dose of a more potent psychostimulant appears to be required; a single high-
AMPH at 24hpi. Dots in bar graphs show measurements for individual animals; the numbers of ChIs recorded were 12–13 cells/Str subregion/treatment; **p < 0.001.

Figure 7. Spontaneous ChI firing 24hpi AMPH. A, Whole-cell recordings were made from ChIs in the four Str subregions. B, An example of ChI firing recorded in the DL Str shows the characteristic spontaneous firing (black trace), and the prominent sag in response to hyperpolarizing-current injection (gray trace). C, Spontaneous firing frequencies of ChIs in each Str subregion are shown after saline (0 mg/kg, n = 30 animals), low-dose (2 mg/kg, n = 22 animals), or high-dose (16 mg/kg, n = 28 animals) AMPH at 24hpi. Dots in bar graphs show measurements for individual animals; the numbers of Chls recorded were 12–13 cells/Str subregion/treatment.

Single dose of AMPH affects ChI activity

P-rpS6\textsuperscript{240/244} signal reports the integrated activity and p-rpS6\textsuperscript{240/244} intensity changes appear to be detected 60 min after pharmacological or behavioral manipulations (Bertran-Gonzalez et al., 2012), suggesting that p-rpS6\textsuperscript{240/244} is suitable to study ChI activity at 2.5hpi or later (Knight et al., 2012). Therefore, the lack of AMPH effect at 2.5hpi is not because of temporal limits of p-rpS6\textsuperscript{240/244} measurement. Stress increases p-rpS6\textsuperscript{240/244} intensity (Knight et al., 2012; Biever et al., 2015a), this may be reflected in the greater p-rpS6\textsuperscript{240/244} intensity in the 2.5hpi compared with the 24hpi saline controls.

In the present study, p-rpS6\textsuperscript{240/244} intensity in Chls was not affected at 2.5hpi after low-dose or high-dose AMPH, while Chl activity modulation via DA neuron glutamatergic cotransmission is dose dependently attenuated after a single dose of AMPH 2.5hpi (Chuhma et al., 2014). This discrepancy could be because of differences in the measurements; p-rpS6\textsuperscript{240/244} reflects the tonic in vivo activity of Chls, which also receive cortical and thalamic glutamatergic inputs in addition to DA neuron inputs (Lim et al., 2014), in contrast to the short phasic firing control of Chls by DA neuron synaptic inputs.

Psychostimulants, including cocaine, methamphetamine and AMPH, are associated with an overall downregulation of DA transmission, both DA release and D2 receptor levels (Ashok et al., 2017). So, we should have expected an increase in ChI activity because of the loss of D2 receptor inhibition. In contrast, the attenuation of ChI activity at 24hpi argues for polysynaptic effects extending beyond direct effects on DA neuron presynaptic terminals. Indeed, AMPH-induced DA release has an onset of minutes and lasts for about 1 h in rodents (Sulzer, 2011), in parallel with behavioral activation. Tonic attenuation of cortical or thalamic glutamatergic inputs may be caused by polysynaptic modulation, resulting in delayed attenuation of ChI activity. Since AMPH does not affect p-rpS6\textsuperscript{240/244} levels or protein synthesis in the Str within 2 h following injection (Rapanelli et al., 2014; Biever et al., 2015b), 2.5 h does not appear to be sufficient to cause long-term circuit changes.

Polysynaptic mechanisms that could contribute to observed decreases in ChI activity in the ventral Str/NAc may involve AMPH effects on other neurotransmitters besides DA. Glutamate efflux in the ventral tegmental area (VTA) is affected by AMPH administration, although both an increase (Xue et al., 1996) and a decrease (Wolf and Xue, 1998) of glutamate efflux have been observed. Acute AMPH exposure induces attenuation of excitatory glutamatergic synaptic transmission in the VTA by activation of serotonin receptors (Jones and Kauer, 1999). AMPH also indirectly affects DA release by stimulating the trace amine-associated receptors (TAAR1) expressed in DA neuron presynaptic terminals (Underhill et al., 2021).

Chls in psychostimulant-induced changes

In the present study, low-dose AMPH significantly attenuated Chl activity in the ventral Str/NAc, a crucial site of psychostimulant action (Russo et al., 2010; Sulzer, 2011). DA neurons projecting to the ventral Str/NAc that corelease glutamate (Hnasko et al., 2010; Stuber et al., 2010) can drive burst firing in Chls (Chuhma et al., 2014; Mingote et al., 2019). A single dose of AMPH attenuates glutamate cotransmission (Chuhma et al., 2014), and mice with conditional reduction in glutamate cotransmission show an attenuated sensitization to repeated AMPH (Mingote et al., 2017). Similarly, we found here that AMPH attenuated Chl activity at 24hpi only in the ventral Str/NAc, suggesting that DA neuron glutamate cotransmission may be one of the factors responsible for NAc-selective attenuation of Chls by low-dose AMPH, in addition to attenuation of phasic firing control through direct synaptic connections of DA neurons.

Although psychostimulant addiction involves repeated use, a single dose of AMPH can induce enduring Str...
circuit changes, drug-dependent behavior and negative affective states, such as anhedonia, depression and anxiety (Vanderschuren et al., 1999; Koob and Le Moal, 2001; Xia et al., 2008; Kameda et al., 2011; Li et al., 2017; Jing et al., 2018; Rincón-Cortés et al., 2018; Jayanthi et al., 2020). Interestingly, even a single dose of AMPH has been found to induce behavioral and neurochemical sensitization, which appears to increase over weeks (Robinson, 1984; Vanderschuren et al., 1999). Our results, in line with these previous findings, point to the relevance of a single dose of AMPH for elucidating drug-induced plasticity. Enduring alterations in ChI activity following acute AMPH exposure point to ChIs as a key component of drug-induced plasticity in the Str circuitry. Further studies using mice with restricted expression of opsins in ChAT neurons will be required to explore whether this reduction in NACh ChI activity is important in subsequent drug-dependent behavior.

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