**NEUROSCIENCE**

**Dopaminylation of histone H3 in ventral tegmental area regulates cocaine seeking**

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Vulnerability to relapse during periods of attempted abstinence from cocaine use is hypothesized to result from the rewiring of brain reward circuitries, particularly ventral tegmental area (VTA) dopamine neurons. How cocaine exposure affects midbrain dopamine neurons to precipitate addiction-relevant changes in gene expression is unclear. We found that histone H3 glutamine 5 dopaminylation (H3Q5dop) plays a critical role in cocaine-induced transcriptional plasticity in the midbrain. Rats undergoing withdrawal from cocaine showed an accumulation of H3Q5dop in the VTA. By reducing H3Q5dop in the VTA during withdrawal, we reversed cocaine-mediated gene expression changes, attenuated dopamine release in the nucleus accumbens, and reduced cocaine-seeking behavior. These findings establish a neurotransmission-independent role for nuclear dopamine in relapse-related transcriptional plasticity in the VTA.

Cocaine increases dopamine neurotransmission from the ventral tegmental area (VTA) to reward-relevant brain regions. This action is central to its addictive properties. Non-neurotransmission roles for dopamine in cocaine dependency have not been considered. Our laboratory has recently described a role for serotonin (5-HT) in developing 5-HTergic neurons, whereby 5-HT located in the nucleus of these neurons was shown to covalently attach to histone proteins—specifically on H3 glutamine 5 (H3Q5)—to regulate gene expression through a process called serotonylation (1, 2). We had hypothesized that this mechanism may generalize to other monoamines in brain, such as dopamine. If true, this process could potentially play a role in the addiction-relevant actions of drugs that stimulate dopaminergic transmission.

Drug addictions are defined by pathological drug-seeking behavior that persists despite adverse consequences. Prolonged vulnerability to relapse is hypothesized to reflect the functional rewiring of brain reward circuitries (3, 4). This is precipitated, at least in part, by drug-induced transcriptional plasticity in midbrain dopamine neurons (5, 6). Histone mechanisms that control chromatin structures, and consequently gene expression, regulate addiction-relevant behaviors (7, 8). Given that histone H3 can be modified by monoamines in response to fluctuations in intracellular availability, we assessed whether dopamine, like 5-HT, can be transferred to the H3 N-terminal tail. We performed targeted, peptide-based liquid chromatography–tandem mass spectrometry (LC-MS/MS) after in vitro transglutaminase 2 (TGM2) (1, 2, 9) enzymatic assays with dopamine. Peptide LC-MS/MS analyses (fig. S1, A to D) revealed Q5 as a reactive substrate for the dopaminyl mark [H3 glutamine 5 dopaminylation (H3Q5dop)]. Given that the serotonyl modification can exist both in isolation (H3Q5ser) and tandem with a dopamine modification (H3Q5dop), we investigated the role of the dopamine modification in regulating gene expression.

**Fig. 1.** Histone H3 dopaminylation in the VTA is dysregulated by cocaine. (A) H3 dopaminylation in human postmortem VTAs from cocaine-dependent subjects versus controls. No changes were observed in H3K4me3Q5dop, H3K4me3, H3, or Tgm2 expression (fig. S5A). A.U., arbitrary units. (B) Experimental timeline of cocaine self-administration (SA) followed by tissue collection time points during withdrawal (WD). FR1-5, fixed-ratio 1 to 5. (C) Number of infusions earned in daily 6-hour test sessions in rats self-administering cocaine or saline. (D) Analysis of H3 dopaminylation in the VTAs (0 versus 1 versus 30 days of WD) from rats with extended access to cocaine versus saline (see fig. S5B for full scatter plots). No changes were observed in H3K4me3Q5dop, H3K4me3, H3, or Tgm2 expression (fig. S5, C to E). Data presented as averages ± SEM. See supplementary materials for full figure legends with statistical comparisons.

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and in combination with H3 lysine 4 trimethylation (H3K4me3) and K4me3, we examined the effect of K4me3 on TGM2-mediated dopaminylation in vitro. Unmodified versus methylated mononucleosomes were subjected to TGM2 dopaminylation assays. Using an antibody against H3K4me3, vide infra, we found that TGM2 equally dopaminylates unmodified and K4me3 substrates, which suggests that both modifications may occur in vivo.

To assess roles for H3 dopaminylation in the context of adult neuronal plasticity, we raised and fully validated single (H3K4me3) and dual (H3K4me3K4dop) modification-specific antibodies (Fig. S3, A to J). We examined whether dopaminyl modifications in the adult brain are modulated by clinically relevant levels of drug exposure. We assessed the expression of these modifications in postmortem human brain tissues obtained from cocaine-dependent individuals compared with matched controls. We focused our investigations on the VTA, the origin of many of the dopaminergic projection neurons that compose the mesocorticolimbic dopamine system (10, 11).

H3K4me3Q5dop, but not H3K4me3Q5dop, was significantly reduced in its expression in the VTAs of cocaine users; H3K4me3, total H3, and Tgm2 were unchanged in their relative levels of expression (Fig. 1A and fig. S5A). However, nearly all of the cocaine users examined in this study displayed pronounced peripheral concentrations of cocaine metabolites at time of death, which may confound the acute pharmacological actions of cocaine with long-term adaptive responses to the drug.

Fig. 2. H3Q5dop in the VTA contributes to cocaine-mediated gene expression. (A) VTA transduced with a lentivirus expressing H3.3Q5A-HA-EF1-RFP (red fluorescent protein) overlayed with a nuclear co-stain [4′,6-diamidino-2-phenylindole (DAPI)]. (B) H3 dopaminylation in VTAs infected with lentiviral vectors. (C) Experimental timeline of self-administration RNA-seq experiment after viral transduction. (D) RRHO map key describing the extent and directionality of overlap between differential gene expression. (E) RRHO comparing differential expression between cocaine-regulated genes. Each pixel represents the overlap between differential transcriptomes, with the significance of overlap of a hypergeometric test color-coded. Coc, cocaine; Sal, saline. (F and G) Overlap of differentially expressed (DEx) PCGs in VTA tissues comparing cocaine versus saline (empty) (F) or cocaine versus saline (H3.3 WT) (G) to H3.3Q5A versus empty or H3.3 WT (cocaine) animals, respectively. (H and I) Heat maps of overlapping genes obtained from RNA-seq data comparing cocaine versus saline (empty) and H3.3Q5A versus empty (cocaine) animals (H) or cocaine versus saline (H3.3 WT) and H3.3Q5A versus H3.3 WT (cocaine) animals (I) using normalized RNA expression values. (J) KEGG 2019 pathway enrichment analysis for the 211 overlapping PCGs displaying reversals in cocaine-mediated gene expression from (G) and (I). LTP, long-term potentiation. Data presented as averages ± SEM.
Therefore, we employed intravenous cocaine self-administration in rats, a well-established procedure to study drug abuse (12, 13), to further explore potential contributions of H3Q5dop to addiction-relevant behaviors. Animals were trained to self-administer cocaine (or saline) under a fixed-ratio 5 schedule of reinforcement (see materials and methods for details). After training, independent cohorts of animals were allocated to two drug treatment groups (as were the respective controls): extended access (6-hour sessions) or restricted access (1-hour sessions) (Fig. 1B and fig. S4A). Animals with extended access to self-administration, but not those with restricted access, demonstrate a gradual escalation of intake across sessions (Fig. 1C and fig. S4A) (12, 14, 15). After 10 days of self-administration, VTA tissues were collected at three different time points: 0, 1, or 30 days. Global levels of H3Q5dop were significantly down-regulated in the VTAs of animals with extended access to cocaine at day 0 (Fig. 1D and fig. S5B), a time point that most appropriately mimics our human subject conditions, in that cocaine is still present at time of death. This reduction was transient, as H3Q5dop levels steadily increased over the course of the next 30 days (Fig. 1D and fig. S5B). By contrast, no changes in VTA H3Q5dop were observed in rats with restricted access to cocaine after 30 days of withdrawal (fig. S4B). Alterations in H3K4me3Q5dop, H3K4me3, total H3, or Tgm2 expression were not observed in animals with either extended or restricted access to cocaine compared with their controls (Fig. 1D, fig. S4B, and fig. S5, C to F). Finally, we assessed levels of the marks in VTA tissues at 30 days of withdrawal in animals receiving cocaine passively—either by being yoked to extended-access rats (fig. S4C and fig. S5G) or through experimenter-administered (intrapерitoneal) injections (fig. S4D and fig. S5H)—and in animals trained to self-administer food under extended-access conditions (fig. S4E and fig. S5I). In all three cases, levels of H3Q5dop, along with expression of H3K4me3Q5dop, H3K4me3, total H3, and Tgm2, remained unaffected.

To explore the functional consequences of H3Q5dop during withdrawal, we delivered a virus vector into the VTA that expresses the H3 variant, H3.3—containing a glutamine-to-alanine substitution at position 5 on its N-terminal tail (Fig. 2A), leading to significant down-regulation of H3Q5dop (Fig. 2B). This reduced the expression of the mark in a dominant-negative fashion without our need for manipulating the activity of Tgm2, an enzyme with diverse functions in the brain that are independent from its histone transamidase activity. H3K4me3Q5dop expression was unaffected by H3.3Q5A delivery. We explored the
impact of attenuating H3Q5dop accumulation in the VTA on drug-induced transcriptional programs that may be responsible for cocaine seeking after prolonged withdrawal. After chronic cocaine versus saline self-administration, rats were infected, intra-VTA, with one of the three viruses, followed by a 30-day period of enforced abstinence (Fig. 2C). After withdrawal, infected VTA tissues were microdissected and processed for RNA sequencing (RNA-seq) (fig. S6, A and B). We first used a threshold-free rank-rank hypergeometric overlap test (RRHO) to assess the patterns and significance of the overlap (Fig. 2D) between differential (cocaine versus saline) gene expression profiles observed after infection with our viral control vectors (H3.3 WT versus empty). The effect of cocaine self-administration on gene expression significantly overlapped between the two groups, which indicates that viral H3.3 WT expression does not affect cocaine-mediated gene expression profiles (Fig. 2E, left panel). We then compared differential gene expression profiles after infection—with either empty (Fig. 2E, middle panel) or H3.3 WT viruses (Fig. 2E, right panel)—with H3.3Q5A-infected VTA after cocaine self-administration. In both cases, attenuating H3Q5dop expression resulted in significant reversals of cocaine-mediated gene expression. Pairwise comparisons were used to identify specific genes displaying dysregulated expression between cocaine versus saline animals (empty or H3.3 WT). Then, overlay assessments were performed to align these sets of genes with those that are also regulated in cocaine self-administering animals expressing H3.3Q5A versus empty (Fig. 2F and tables S1 and S2; 720 differentially expressed protein coding genes (PCGs) were found to overlap) or H3.3 WT (Fig. 2G and tables S3 and S4; 211 differentially expressed PCGs were found to overlap) viruses. Further evaluation of these overlapping PCGs found that ~95 to 87% were significantly rescued in their expression with H3.3Q5A delivery; H3.3Q5A expression in saline self-administering animals had little to no impact on the expression of overlapping genes (Fig. 2, H and I). Gene enrichment analysis [Human Kyoto Encyclopedia of Genes and Genomes (KEGG) 2019] indicated strong associations with pathways involved in the regulation of synaptic function and drug addiction (Fig. 2J and table S5). These findings suggest that H3Q5 dopamine plays an important role in cocaine-induced transcriptional plasticity in the VTA.

Next, we examined the effect of H3Q5dop on dopaminergic neuronal activity. Cocaine-naïve rats were injected intra-VTA with viruses expressing empty vector or H3.3 WT controls versus H3.3Q5A. We recorded spontaneous action potentials (sAPs) from infected dopaminergic neurons expressing hyperpolarization-activated currents (Fig. 3A). Attenuation of H3Q5dop significantly reduced the frequency of sAPs arising from dopaminergic neurons.

To more directly assess the impact that blocking H3Q5dop during cocaine withdrawal has on dopamine release into the nucleus accumbens (NAc), rats were allowed to self-administer cocaine under extended-access conditions and were infected intra-VTA with one of the three viruses. After 30 days of withdrawal and cue-induced cocaine seeking, ex vivo assessments of dopamine release using fast-scanning cyclic voltammetry (FSCV) were performed (Fig. 3B). Stimulation-induced dopamine release into the NAc was significantly attenuated (Fig. 3, C to F) in response to reductions in H3Q5dop accumulation.

To investigate what consequences modifying H3Q5dop levels has on relapse-relevant behaviors, we allowed an independent cohort of animals to self-administer cocaine under extended-access conditions followed by intra-VTA viral manipulations. After withdrawal, rats were returned to a drug-paired context, and cocaine seeking was assessed (Fig. 4A). Preventing H3Q5dop accumulation in animals with extended access significantly reduced their drug-seeking behavior (Fig. 4, B and C). Reducing H3Q5dop levels in the VTA of a separate cohort of animals with restricted access did not affect their drug-seeking behavior (fig. S7, A to C). To ensure that such manipulations do not impair motivation to seek natural rewards, additional rats were trained to respond for food rewards under the same extended-access schedule of reinforcement, followed by viral manipulations, 30 days of ad libitum feeding, and subsequent reward seeking (fig. S8A). Food-seeking responses remained unaffected by manipulations of H3Q5dop (fig. S8, B and C). Given the fact that chronic cocaine use can elicit numerous additional behavioral abnormalities, we performed intra-VTA viral manipulations in cocaine-naïve rats, followed by assessments of psychomotor sensitization in response to repeated experimenter-administered cocaine. Both control (H3.3 WT) and knockdown (H3.3Q5A) animals displayed similar levels of cocaine-induced locomotor sensitization (fig. S8, D and E).

We demonstrated that a previously uncharacterized chromatin modification, histone dopaminylation, is critically involved in regulating aberrant neuronal gene expression patterns in the VTA in response to cocaine consumption. The increased expression of H3Q5dop that follows prolonged withdrawal from extended, but not restricted, access to cocaine self-administration regulates relapse-like behaviors. It does so, in part, through the dysregulated transcription of addiction- and synaptic plasticity–related genes in the VTA, as well as through aberrant dopamine release dynamics in the NAc after cocaine withdrawal (fig. S9, model). Gaining a better understanding of the mechanistic roles for H3Q5dop in
mediating permissive compared with repressive transcription, as well as the genes regulated through this mechanism, will greatly improve our knowledge of the molecular underpinnings of drug addiction.

REFERENCES AND NOTES


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SUPPLEMENTARY MATERIALS

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Materials and Methods
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More than a normal neurotransmitter
The molecular mechanisms underlying the persistence of addiction remain largely unclear. Lepack et al. found that, with cocaine exposure, there is an intracellular accumulation of dopamine in neurons of a brain region called the ventral tegmental area (see the Perspective by Girault). Dopamine associates with chromatin to initiate a previously unknown form of epigenetic regulation called dopaminylation. This modification has an impact on ventral tegmental area function and, consequently, on dopaminergic action potentials. The result is aberrant dopamine signaling in the ventral striatum during periods of drug seeking.

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