

Drug Predictive Cues Activate Aversion-Sensitive Striatal Neurons That Encode Drug Seeking

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Drug-associated cues have profound effects on an addict's emotional state and drug-seeking behavior. Although this influence must involve the motivational neural system that initiates and encodes the drug-seeking act, surprisingly little is known about the nature of such physiological events and their motivational consequences. Three experiments investigated the effect of a cocaine-predictive stimulus on dopamine signaling, neuronal activity, and reinstatement of cocaine seeking. In all experiments, rats were divided into two groups (paired and unpaired), and trained to self-administer cocaine in the presence of a tone that signaled the immediate availability of the drug. For rats in the paired group, self-administration sessions were preceded by a taste cue that signaled delayed drug availability. Assessments of hedonic responses indicated that this delay cue became aversive during training. Both the self-administration behavior and the immediate cue were subsequently extinguished in the absence of cocaine. After extinction of self-administration behavior, the presentation of the aversive delay cue reinstated drug seeking. *In vivo* electrophysiology and voltammetry recordings in the nucleus accumbens measured the neural responses to both the delay and immediate drug cues after extinction. Interestingly, the presentation of the delay cue simultaneously decreased dopamine signaling and increased excitatory encoding of the immediate cue. Most importantly, the delay cue selectively enhanced the baseline activity of neurons that would later encode drug seeking. Together these observations reveal how cocaine cues can modulate not only affective state, but also the neurochemical and downstream neurophysiological environment of striatal circuits in a manner that promotes drug seeking.

Key words: addiction; cocaine; dopamine; electrophysiology; relapse; voltammetry

Introduction

Informed by drug addicts' emotional and physiological reactions to drug-associated cues (Childress et al., 1999; Garavan et al., 2000), theories of drug addiction and relapse often posit an influence of cues on drug-seeking behavior (Di Chiara, 1999; Robinson and Berridge, 2003; Everitt et al., 2008). These theories often note the desirability of these incentivized cues, which have a well known neural representation in both mesolimbic dopamine (DA) signaling and cell firing in the nucleus accumbens (NAc). Such signals of immediate drug reward reliably increase striatal DA (Phillips et al., 2003), activate NAc neurons (Hollander and Carelli, 2007), and promote drug seeking (Shaham et al., 2003).

However, human addicts report that drug-associated cues induce negative feelings related to drug craving and anxiety that predict relapse (Sinha et al., 2000, 2003). The ability of drug cues

to cause negative affect can be studied in animal models, and such studies have demonstrated that predictors of delayed drug access can cause avoidance (Tzschentke, 1998; Grigson and Twining, 2002), negative affect (Wheeler et al., 2008), a reduction of DA (Wheeler et al., 2011), and enhanced excitatory activity in reward circuitry (Wheeler et al., 2008). Additionally, the degree of aversion correlates with enhanced drug-seeking motivation (Wheeler et al., 2011; Nyland and Grigson, 2013; Colechio et al., 2014). The fact that appetitive and aversive stimuli can differentially affect striatal circuitry has led us to hypothesize that aversive stimuli disinhibit a distinct aversion-sensitive striatal circuit that also drives drug seeking (Twining et al., 2014), possibly as an avoidance behavior (Baker et al., 2004; Koob and Le Moal, 2008). Although recent studies have shown that correcting a reduction in DA signaling can attenuate drug taking and seeking (Twining et al., 2014; Willuhn et al., 2014), it is not clear how an aversive stimulus that decreases DA signaling simultaneously impacts striatal cell firing to promote drug seeking.

Here we directly tested whether an aversive cue can cause drug seeking, if it does so in a state of low DA tone, and if it activates neurons that encode the drug-seeking act. Additionally, we investigated the impact of the aversive drug cue on the processing of another drug-associated stimulus and drug-seeking behavior. The potential change in the physiological response to a proximate cocaine cue (i.e., a cue that signals immediate drug infusion) was of particular interest because aversive stimuli can interact synergistically with proximate drug cues to drive drug seeking (Liu and

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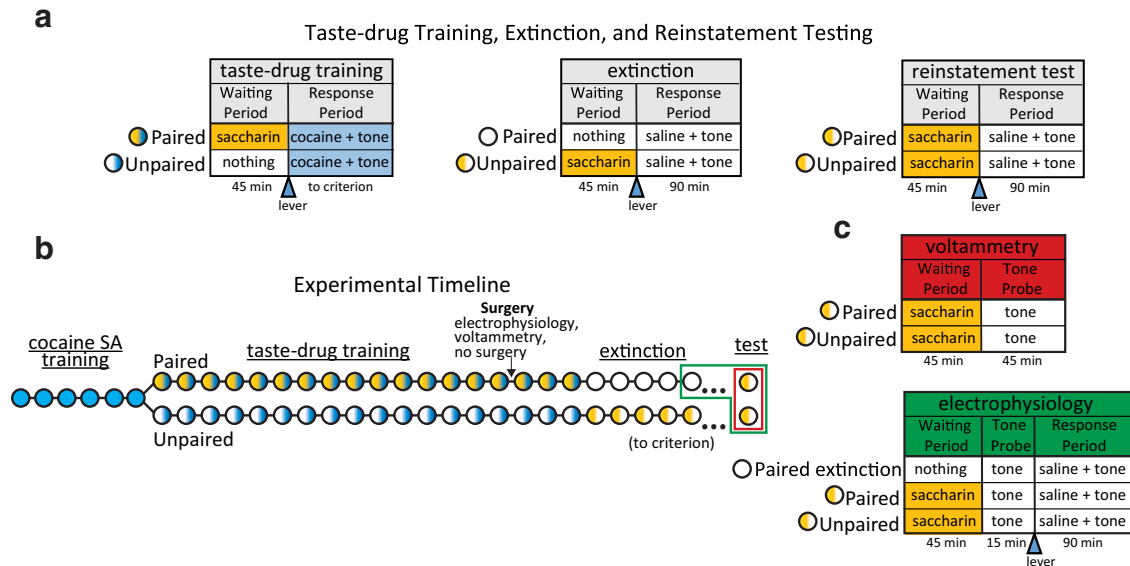


Figure 1. Overview of experimental design. **a**, Details of the three experimental phases. During taste-drug training, rats experienced a waiting period during which they received 45 saccharin infusions (yellow) or nothing (white), followed by lever extension (triangle), and a response period during which cocaine and a co-occurring tone stimulus could be self-administered (blue). During the response period of the extinction and reinstatement phases, cocaine was replaced by saline (white). Paired rats received saccharin exposure during the taste-drug training and reinstatement phases, and unpaired rats received saccharin during the extinction and reinstatement phases. **b**, Timeline of training and testing. Before being divided into groups, all rats self-administered cocaine with no waiting period for six daily sessions (circles). During the three experimental phases, rats received saccharin (yellow) or nothing (white) during the waiting period, and cocaine (blue) or saline (white) during the response period. Green and red boxes denote electrophysiology and voltammetry recording sessions, respectively. **c**, Details of voltammetry and electrophysiology testing. Following extinction, both groups received saccharin infusions followed by noncontingent presentations of the tone while rapid dopamine release (voltammetry) or neuronal firing rate (electrophysiology) was monitored. In the electrophysiology study, this was followed by a response period during which rats could self-administer saline.

Weiss, 2002; Shelton and Beardsley, 2005; Buffalari and See, 2009). The pattern of observed results supports the hypothesis that an aversive drug cue causes a sustained reduction in DA signaling, induces negative affect, and activates a subpopulation of striatal neurons that encode drug seeking and other drug-associated cues.

Materials and Methods

General experimental design. The novel procedure presented here was inspired by traditional studies of context-induced renewal of drug seeking in which the self-administration context is shifted during extinction, and returned during test (Crombag and Shaham, 2002; Bouton, 2004; i.e., ABA renewal). This treatment is typically compared with a control group that receives a context shift between self-administration and extinction, but not between extinction and test (the ABB control). In the present design, the paired group of rats received passive exposure to a normally appetitive saccharin solution before the opportunity to self-administer cocaine (Fig. 1) and the unpaired control group received cocaine access following a waiting period. During extinction, paired subjects experienced a waiting period followed by an extinction session, whereas unpaired subjects received saccharin infusions before extinction. Finally, paired rats experienced a reinstatement test session in which saccharin preceded extinction, and unpaired rats received another normal extinction session that was used as a test session for behavioral and physiological comparisons. One potential pitfall in this novel design concerns the fact that saccharin is not a motivationally neutral stimulus, even in the unpaired group. However, by comparing the final extinction session in the paired and unpaired groups with the test session in the paired group, we can determine the potential effect of the aversive cocaine cue, an inherently appetitive unconditioned stimulus, and no stimulus, on affect and drug-seeking behavior.

Subjects. Male Sprague-Dawley rats (Harlan Laboratories) weighing 275–300 g were individually housed in a temperature and humidity controlled AAALAC-accredited vivarium. Rats were maintained on a 12 h reversed dark/light cycle and all experimental procedures were conducted during the dark phase (starting at 07:00 h). Animals all had *ad*

libitum access to food and water except where otherwise noted. All experimental protocols were approved by the Institutional Animal Care and Use Committee at Marquette University in accordance with the National Institutes of Health *Guide for the Care and Use of Laboratory Animals*. A total of 48 rats were used in these studies, and trained as described above. Subsets of this group were assayed for electrochemical and electrophysiological encoding of various aspects of the task. Fifteen (8 paired and 7 unpaired) were used only for behavior. Twenty-five (16 paired and 9 unpaired) and eight (4 paired and 4 unpaired) were used for electrophysiology and voltammetry assays.

Surgical procedures. All surgical procedures were conducted under ketamine/xylazine (100 mg/kg; 20 mg/kg, i.p.) anesthesia. Intraoral and intrajugular catheter implantations were conducted as previously described (Wheeler et al., 2008). To prepare for voltammetric recordings, electrode guide cannula (BASi) were implanted above the NAc core (NAcC) unilaterally (AP: +1.3; ML: ±1.3), a bipolar stimulating electrode (Plastics One) was placed above the ipsilateral VTA (AP: −5.2; ML: ±1.0; DV: −8.0), and an Ag/AgCl reference electrode was placed contralateral to the guide cannula. For electrophysiological recordings, eight-wire microelectrode arrays (NB Labs) were implanted bilaterally in the NAc at AP: +1.7 mm, ML: ±0.8 NAc shell (NAcS) to ±1.3 mm (NAcC), DV: −6.3 mm. For each array, another wire was wrapped around a skull screw to serve as a ground. For all surgical procedures, rats were treated with the anti-inflammatory meloxicam (1% oral suspension) the day of surgery and for 2 d following surgery to reduce inflammation and postoperative pain. To maintain patency, the catheters were flushed daily with dH₂O (i.o.) or heparinized saline and the antibiotic cephazolin (i.v.).

Apparatus. Subjects were tested in standard operant chambers (Med Associates), interfaced with a computer, and housed in sound attenuating Faraday cages. Each operant chamber had two dedicated syringe pumps for intravenous and intraoral infusions that were delivered to each rat via a dual-channel fluid swivel (Instech Laboratories). Two retractable levers entered the chamber on the right side wall with a cue light directly above each. A food-pellet dispenser delivered 45 mg grain pellets (BioServe) to a recessed food cup positioned between the two levers. A

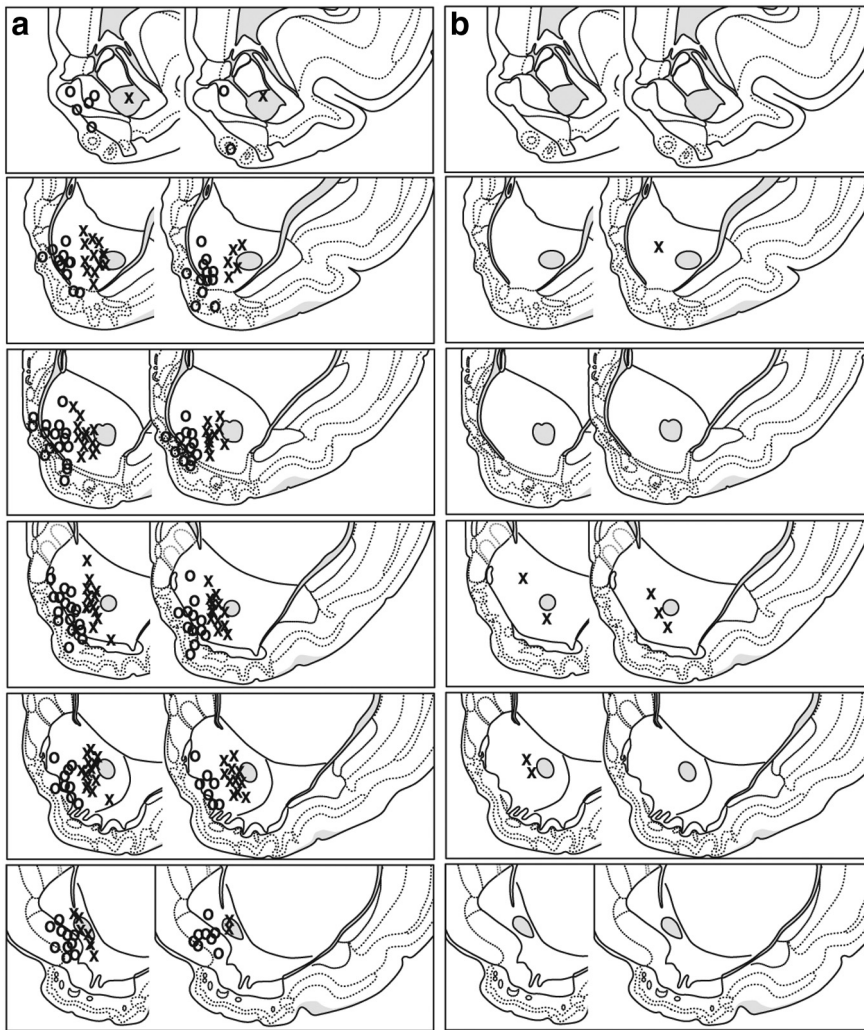


Figure 2. Electrode placements. *a*, Electrophysiology electrode placements in the NAcC (Xs) and NAcS (Os) in paired (left; $n = 16$) and unpaired (right; $n = 9$) rats. *b*, Voltammetry electrode placements in paired (left; $n = 4$) and unpaired (right; $n = 4$) rats.

speaker capable of producing a 65 dB 1 kHz tone was located on the left sidewall. Under each chamber, a camera was positioned to allow for recording behavioral responses (taste reactivity). The chamber floor was clear acrylic glass and a house light was positioned on the door of the sound attenuating chamber, outside of the operant chamber, to ensure recording quality.

Self-administration training. Following catheter implantation, subjects were allowed to recover until they reached presurgery bodyweight. Subjects were then food restricted to 90% bodyweight, and trained to press one of the two levers for food pellets. Upon acquisition of lever-press behavior, subjects were trained to self-administer cocaine on a fixed-ratio one schedule of reinforcement. The beginning of each cocaine self-administration session was signaled by the entry of both levers into the box and the illumination of two cue lights. Responses on the active lever resulted in a 3 s cocaine infusion (0.25 mg/0.1 ml) accompanied by cue-light offset, onset of a 5 s tone, and a 20 s timeout period. Responses on the active lever during the timeout resulted in no programmed consequences and are not included in the data analyses. The conclusion of the time out period was signaled by cue light onset. Each session was completed only when each subject reached a maximum infusion cap (detailed below).

Self-administration sessions occurred in a series of six experimenter-controlled 6 d cycles consisting of 3 d of cocaine self-administration and 3 d without cocaine access in the home cage. The maximum number of infusions taken per session increased each cycle from a 20 infusion cap in the first cycle, to a 50 infusion cap in the final cycle. Upon completion of

cycles 1 and 2, subjects were separated into groups, counterbalanced for weight and average latency to reach to the infusion cap in the second cycle. Rats were initially trained for six sessions without a waiting period to balance the groups and to ensure the acquisition of a tone–cocaine association independent from saccharin exposure. For paired rats, all subsequent cocaine self-administration sessions were preceded by a 45 min waiting period during which each subject received 45 intraoral saccharin infusions (0.15%; 0.2 ml/6 s; 1 inf/min). Unpaired subjects received an equivalent waiting period with no taste before cocaine access.

Extinction and reinstatement. Upon completion of six cycles of cocaine self-administration all subjects were transitioned into the extinction phase. Before every extinction session, unpaired rats received 45 intraoral infusions of saccharin, whereas paired rats experienced a waiting period. Following the saccharin period, a 90 min extinction session began in which presses on the formerly active lever resulted in cue light offset, tone onset, and an infusion of saline. After a minimum of five extinction sessions, subjects were tested for reinstatement when they registered an average of 10 presses or less on the formerly active lever (excluding presses made during the 20 s time out period) over two consecutive sessions. Extinction responding was analyzed with a mixed ANOVA comparing the paired and unpaired groups across matched extinction sessions.

Once each subject reached extinction criteria they were tested for saccharin-induced reinstatement of cocaine seeking. This session was procedurally identical to an extinction session with the exception that all subjects received saccharin infusions. Reinstatement was assessed by comparing responding in the paired group at test with the final extinction session in both the paired and unpaired groups using *t* tests. To assess the relationship between

reinstatement behavior and aversion, a regression analysis was performed (Pearson's *R*). Because of variability in baseline rates of lever pressing at the end of extinction, a reinstatement ratio was computed for the regression analysis by dividing the total number of lever presses during test by the sum of number of lever presses during test and extinction.

Taste reactivity scoring/analysis. Taste reactivity was analyzed in a frame-by-frame analysis using digital video recorded on the test session in paired and unpaired rats. Appetitive and aversive responses were counted in the saccharin period using the technique of Grill and Norgren (1978). Bouts of “wet dog shaking,” paw flailing, and mouth movements that matched the “triangle” shape for a duration exceeding 90 ms were counted as aversive. Instances in which the tongue protruded and crossed the midline were counted as appetitive. The rates of aversive and appetitive events (per trial) were computed for each animal. Taste reactivity data were analyzed with a mixed ANOVA and subsequent planned comparisons of appetitive and aversive taste reactivity (events/trial) in the paired and unpaired groups.

Voltammetry testing and data collection. Subjects underwent fast-scan cyclic voltammetry surgery after their 15th session of cocaine self-administration training. Following recovery, subjects received their final 3 d of self-administration and extinction as previously described. To familiarize the rats with the recording situation, the VTA stimulating electrode was harnessed to a rotating commutator (Crist Instruments) during at least one of the final extinction sessions. Once subjects reached extinction criteria DA responses to saccharin and tone were measured. A

carbon fiber electrode was lowered into the NAc. The fiber was held at -0.4 V against Ag/AgCl between scans and then driven to $+1.3$ V and back in a triangular fashion at 400 V/s for each voltammetric measurement. The application of this triangle waveform causes oxidation and reduction of chemical species that are electroactive within this potential range, producing a change in current at the carbon fiber. Specific analytes (including dopamine and pH) are identified by plotting these changes in current against the applied potential to produce a cyclic voltammogram (Heien et al., 2004, 2005). The current arising from electrode processes was removed by using background subtraction. For data collected during testing, the background period was taken as the minima during the 10 s before saccharin or cue presentation. This practice does not subtract the presence of phasic dopamine release events because the background was explicitly selected for the absence of dopamine signals. Measurements were made every 100 ms and, after driving the electrode into the NAc core, the electrode equilibrated for 40 min before any data were collected. The position of the microelectrode was then optimized by monitoring the presence of spontaneously occurring dopamine release events (Roitman et al., 2008; Wheeler et al., 2011). Once this was observed, the electrode was locked in place and data collection proceeded. The experiment consisted of a 20 min baseline dopamine monitoring period, then a 45 min saccharin delivery period, followed by a 45 min tone test period during which the proximate cocaine cues (tone onset and cue light offset) were presented every 90 s. Orofacial expressions during saccharin administration were recorded and scored. Following the conclusion of testing, several electrical stimulation trains that varied in number of pulses (1, 2, 5, 10, and 20) were administered for the generation of a training set for principal component analysis for the detection of dopamine and pH changes during the behavioral session. Due to technical limitations of our voltammetry testing apparatus, DA was not measured during lever-press behavior.

Analyte identification and quantification were achieved using principal component regression analysis described previously in detail (Heien et al., 2004). All data presented here fit the resulting model at the 95% confidence level. Briefly, training sets were generated from background-subtracted cyclic voltammograms collected during and after electrical stimulations. At least 10 voltammograms were obtained for DA and pH. The resulting current amplitude was converted to DA concentration based on calibration of the electrode in a flow injection analysis system after the *in vivo* experiment. To convert current due to oxidation of DA, 500 nM and 1 μ M dopamine in a buffer, pH 7.4, were used in the flow-injection analysis system. Data from each saccharin or cue probe presentation were background subtracted using a 1 s block at the local minima in the 10 s before presentation. The resultant current changes over time were converted into DA concentrations over time using a principle component regression.

Voltammetry data analysis. For each rat, trials were aligned to saccharin or tone presentation and averaged together. To analyze the acute effects of saccharin on DA, average concentration during a baseline epoch (10 s before infusion onset) was compared with average concentration during an effect epoch (10 s after infusion onset). Additionally, the diffuse effect of saccharin on non-time-locked DA was examined by blocking baseline epoch measurements into thirds (early, middle, and late). Unlike the acute dopaminergic response to saccharin, which lasted several seconds, the observed effect of the tone was rapid (~ 1 s). Thus, to analyze the effect of the tone on DA concentration, the peak DA concen-

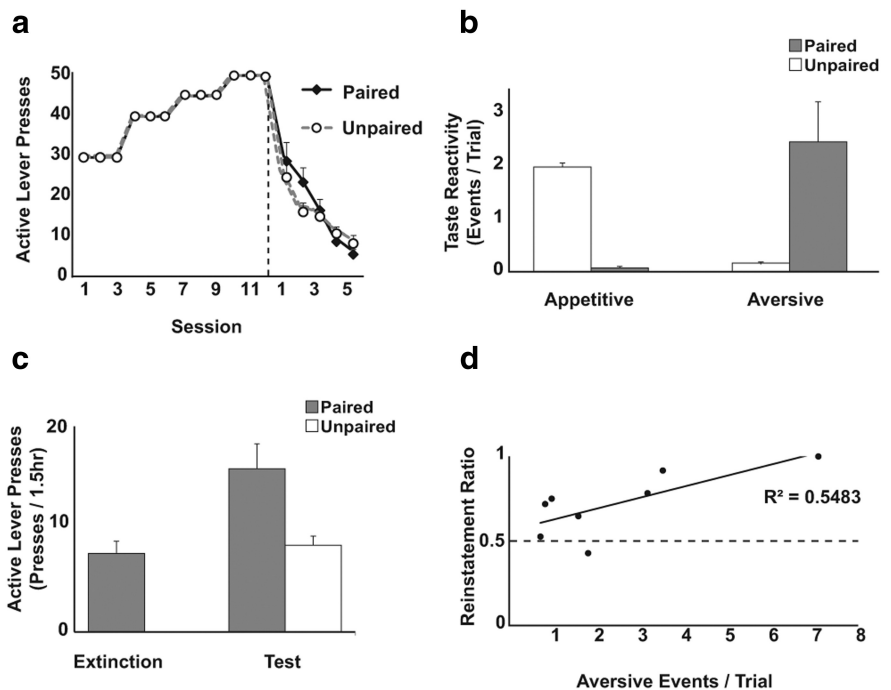


Figure 3. The aversive drug cue causes drug seeking. *a*, Lever pressing during self-administration and extinction. Session duration varied during self-administration because all rats responded until they reached a progressive cap. During extinction, session duration was 1.5 h. The vertical dashed line depicts day of switch to nonreinforced responding. Responding during extinction was similar in paired ($n = 8$) and unpaired ($n = 7$) rats, mean \pm SE. *b*, Paired rats showed more aversive taste responses to intraoral saccharin infusions per trial than unpaired rats ($p < 0.02$), whereas unpaired rats showed more appetitive responses than paired rats ($p < 0.01$). *c*, Paired rats increased responding on the previously active lever from extinction to test ($p < 0.01$) and responded more than unpaired rats ($p < 0.02$). *d*, The magnitude of the aversion caused by the drug cue predicted the reinstatement of drug seeking ($p < 0.04$). A ratio > 0.5 (the horizontal dashed line) indicates an increase in responses on the previously active lever during test compared with extinction.

tration observed during the 5 s tone was compared with the peak DA observed in the final 5 s of the baseline epoch. Changes in DA were analyzed using repeated-measures ANOVAs and planned contrasts were used for direct comparisons.

Electrophysiology recording procedures. Subjects underwent electrode implantation surgery after their 15th session of self-administration training. Recordings were conducted with microelectrode arrays featuring eight stainless steel wires (50 μ m diameter) arranged in a 2×4 configuration (NB Labs). Following recovery, subjects received their final 3 d of self-administration and extinction. To familiarize the rats with the recording situation, they were connected to a flexible recording cable (Plexon) attached to a commutator (Crist Instruments) during these sessions. To assess encoding of the tone independent of lever pressing, eight paired and eight unpaired subjects also experienced noncontingent tone presentations during the last extinction session, and during the test session. Fifteen tone trials were delivered at 60 s intervals after the saccharin-infusion period. After the tone testing, the levers were extended, and extinction/reinstatement testing commenced. The headstage contained 16 miniature unity-gain field effect transistors. NAc activity was recorded differentially between each active wire and an inactive wire chosen for an absence of neuronal activity. Online isolation and discrimination were accomplished using a commercially available neurophysiological system (OmniPlex system; Plexon). Multiple-window discrimination modules and high-speed analog-to-digital signal processing in conjunction with computer software enabled isolation of neuronal signals based on waveform analysis. The neurophysiological system incorporated an array of digital signal processors (DSPs) for continuous spike recognition. The DSPs provided a continuous parallel digital output of neuronal events to a computer. Another computer controlled behavioral events of the experiment (Med Associates) and sent digital outputs corresponding to each event to the OmniPlex to be time-stamped along with the neural data. Criteria for identifying different

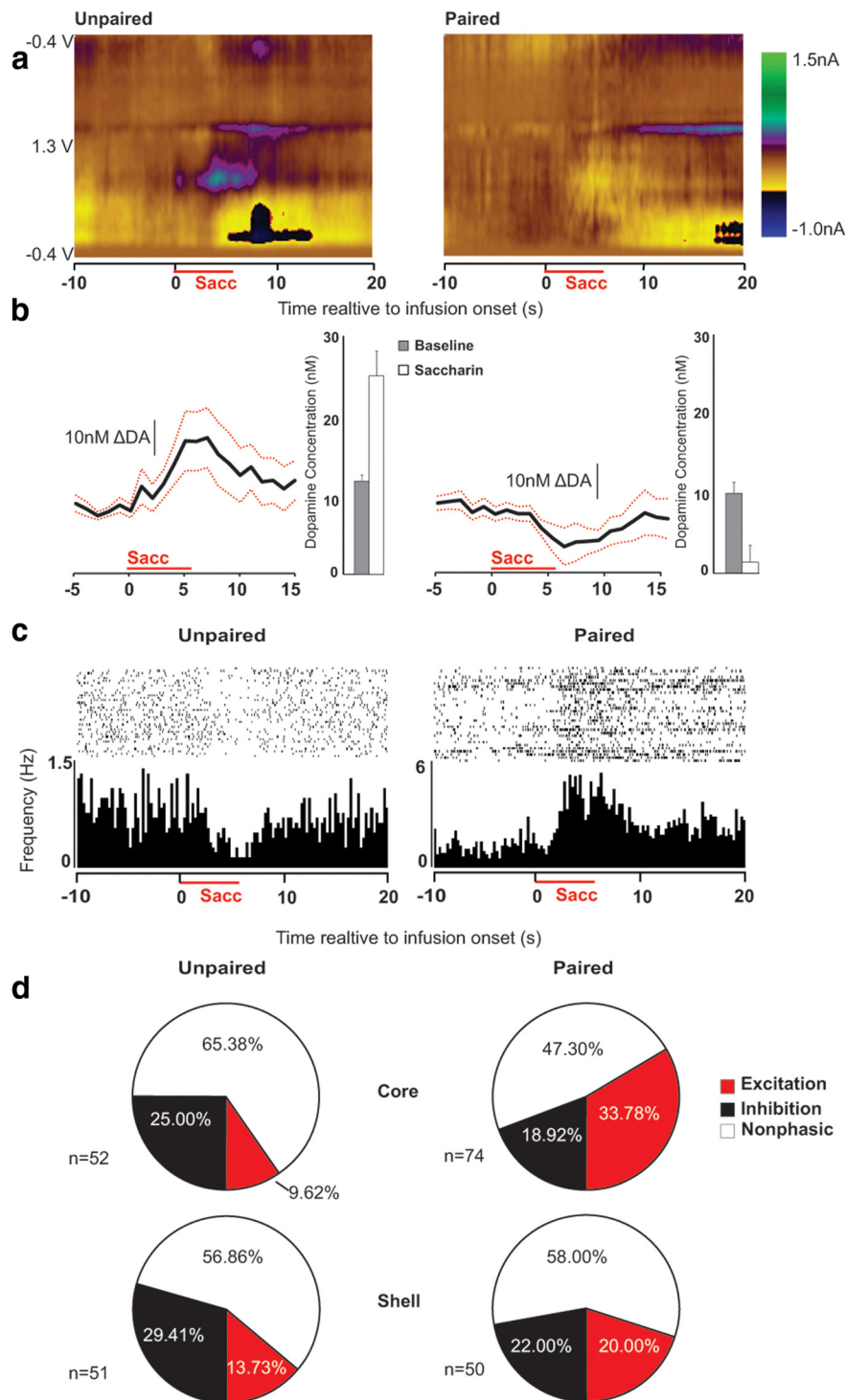


Figure 4. The aversive drug cue reduces DA and enhances neural excitation acutely. *a*, Average color plots from representative unpaired (left) and paired (right) rats showing dopaminergic responses to saccharin (horizontal red line). The figures show current changes (nA; z-axis) plotted against voltage (V; y-axis) and time (s; x-axis). *b*, Relative to the 10 s baseline epoch, saccharin acutely increased DA concentration (nM) in unpaired rats (left) and decreased DA concentration in paired rats (right; $p < 0.02$). Black lines show average concentration over time and red lines indicate SE for each group. Bars depict overall averages from the baseline and saccharin epochs. *c*, Rasters and histograms depict examples of inhibitory (left) and excitatory (right) changes in firing rate time-locked to the presentation of saccharin. Individual action potentials during each saccharin presentation are shown in the raster, and firing rate (Hz) is shown in the histogram below. *d*, Categorization of responses to saccharin in all recorded neurons. Unpaired neurons (left) show more excitatory than inhibitory responses to saccharin, whereas paired neurons (right) show more inhibitory than excitatory responses ($p < 0.01$).

neurons on a single wire were described previously in detail (Roitman et al., 2005). Briefly, discrimination of individual waveforms corresponding to a single neuron was accomplished using template and principle component analysis procedures provided by the PlexControl software sys-

tem. The template analysis procedure involves taking a sample of the waveform and building a template of that extracellular waveform. Subsequent neurons that match this waveform are included as the same neuron. Cell sorting was further accomplished after the experiment was over using additional principle components analysis in Offline Sorter v3.3.2 (Plexon).

Phasic response categorization. Phasic encoding of the saccharin, tone, and lever-press response was characterized by generating peri-event histograms (500 ms bins) surrounding each event in Neuroexplorer 4.126 (Nex Technologies). To ensure a reliable histogram, lever-press responses were only considered whether the subject registered 10 or more responses spaced at least 20 s apart. Each histogram was divided into a baseline epoch and an effect epoch based on the type of event: (1) 10 s immediately following the onset of the saccharin infusion, (2) 5 s immediately following the onset of the tone, (3) 5 s immediately preceding a lever press, and (4) 5 s immediately following the lever press. Individual bins within the effect epochs were then standardized based on the mean and SD of the appropriate baseline epoch. Neurons that exhibited two or more bins with Z-scores that exceeded or fell < 3 SD were classified as excitatory or inhibitory. Because some inhibitions reached 0 Hz, but still did not fall below three Z-scores, inhibitory responses were also included if two or more bins had the minimum possible Z-score, provided that the effect epoch showed at least twice as many minimum bins as the baseline epoch. Neurons were also categorized based on a *t* test comparison of the frequency bins of neurons that exhibited $> 15\%$ shifts in frequency from baseline to the effect epoch ($\alpha = 0.05$). After neurons were characterized, Fisher's exact tests were used to detect any differences in phasic responses and the distribution of excitatory and inhibitory responses in the core and shell during paired extinction, unpaired extinction, and paired testing.

Baseline analysis. An average firing rate (Hz) for each trial was drawn from the baseline epoch for the saccharin infusion and transformed [$\log_{10}(x + 0.001)$] to normalize the distributions (Guillem et al., 2014). Trials were grouped in thirds to capture any changes in baseline firing that might occur during the saccharin period. Previously categorized tone and lever-press encoding neurons were selectively analyzed with repeated-measures ANOVAs to determine whether their baseline firing rates are altered during saccharin exposure. One inhibitory tone-encoding neuron in the unpaired group did not have sufficient data for a complete time course analysis.

Histology. After voltammetry and electrophysiology testing, subjects were killed with CO_2 . To verify placements of voltammetry recordings, small electrolytic lesions were created by running a current (250 μA) through a stainless steel electrode placed at the depth at which the recording took place. Brains were then removed and submerged in 10% formaldehyde for 14 d. For electrophysiology recordings, a current (20 μA) was run through the implanted microwires, and brains incubated in a 10% formaldehyde/4% potassium ferrocyanide solution. All brains were then sliced into 40 μm sections, mounted, stained with

0.25% thionin, and coverslipped. Figure 2 shows the electrode placements from the voltammetry and electrophysiology experiments.

Results

The aversive drug cue reinstates extinguished drug seeking

Although prior studies have demonstrated a predictive relationship between cue-induced negative affect and drug taking, here we tested the ability of an aversive drug cue to cause reinstatement. Because self-administration sessions had a quota/cap, all subjects took the same amount of cocaine. To examine the extinction data (Fig. 3*a*), responses on the formerly active lever (excluding those made during the time out period) were analyzed. Both groups decreased responding similarly on the active lever across sessions (session: $F_{(4,52)} = 22.41$, $p < 0.01$; all other p values > 0.19).

Taste reactivity to saccharin at test resembled previous reports (Fig. 3*b*), with paired rats showing more aversive ($p < 0.02$) and fewer appetitive ($p < 0.01$) responses relative to unpaired rats (group \times taste reactivity: $F_{(1,13)} = 25.75$, $p < 0.01$; all other p values > 0.45). Paired rats exhibited enhanced drug seeking relative to their final extinction session (Fig. 3*c*; $t_{(7)} = 3.85$, $p < 0.01$), and unpaired rats at test ($t_{(13)} = 2.90$, $p < 0.02$). In contrast, responses on the inactive lever did not differ between paired (mean = 2.88; SE = 1.08) and unpaired rats (mean = 2.88; SE = 1.08; $t_{(13)} = 1.68$, $p = 0.12$). Furthermore, the aversive reaction to cocaine-associated saccharin was correlated with reinstatement in paired rats (Fig. 3*d*; $r_{(6)} = 0.74$, $p < 0.04$).

The aversive drug cue reduces DA and enhances neural excitation acutely

Consistent with observations in the NAcS (Wheeler et al., 2011), the aversive and appetitive stimuli had opposite effects on NAcC DA (Fig. 4*a,b*). Cocaine-predictive saccharin caused an acute decrease in NAcC DA concentration in the paired group, whereas appetitive unpaired saccharin caused an acute increase in DA concentration (group \times epoch: $F_{(1,6)} = 10.45$, $p < 0.02$; group: $F_{(1,6)} = 6.42$, $p < 0.01$; epoch: $p > 0.50$).

Also consistent with prior reports, aversive and appetitive stimuli had differential effects on NAc neuronal firing rates. Figure 4*c* shows examples of inhibitory and excitatory neural encoding of the saccharin stimulus at test. Saccharin infusions were encoded by 48.39% ($n = 60/124$) of the paired neurons and 38.84% of the unpaired neurons ($n = 40/103$). Encoding of the aversive drug cue was predominantly excit-

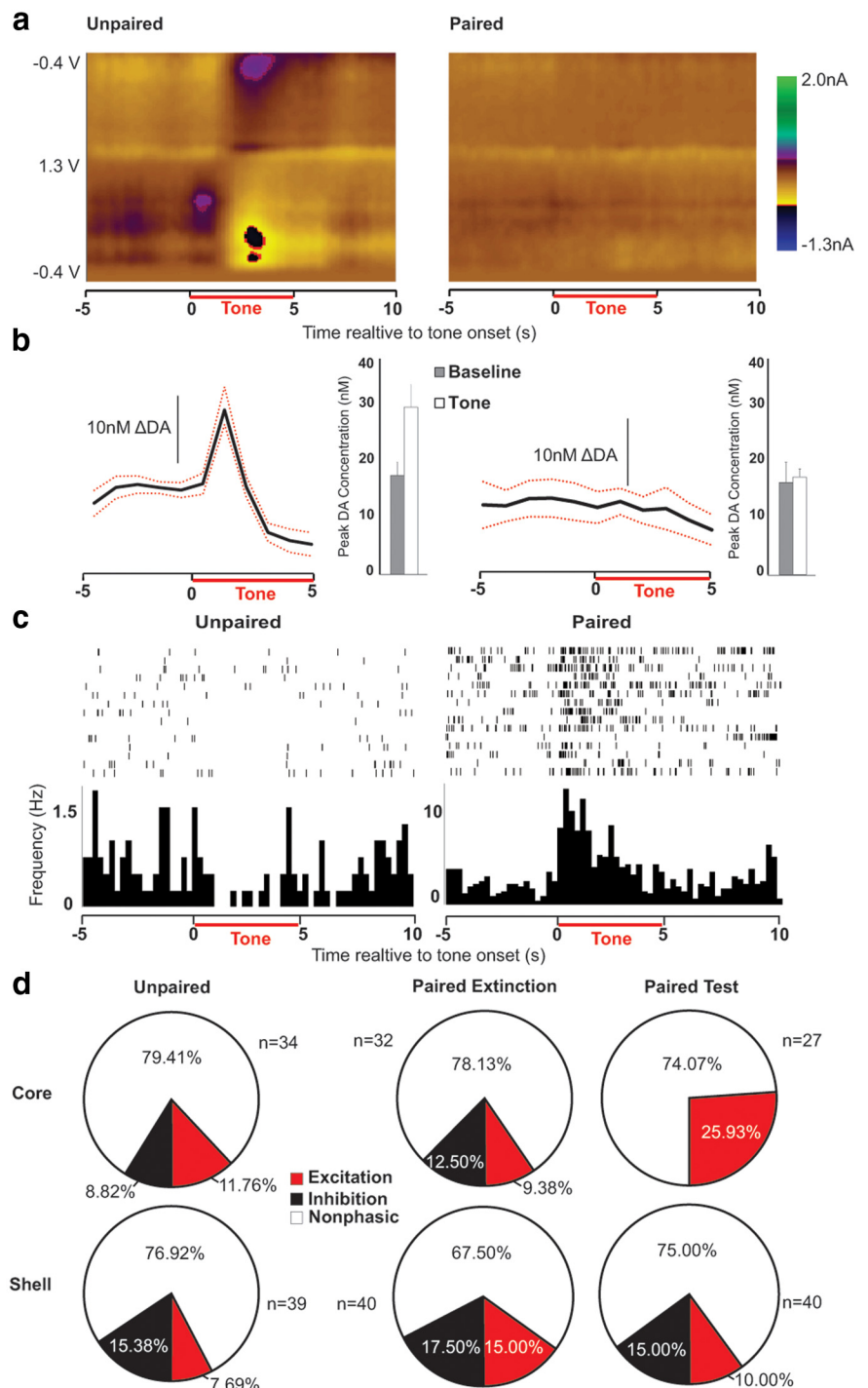


Figure 5. The aversive drug cue eliminates DA encoding and augments excitatory neural encoding of the proximate drug cue. *a*, Average color plots from a representative unpaired (left) and paired (right) rat showing the dopaminergic response to the proximate drug cue (tone; horizontal red line). The figures show current changes (nA; z-axis) plotted against voltage (V; y-axis) and time (s; x-axis). *b*, Relative to the 5 s baseline epoch, the tone caused a brief increase in DA concentration (nM) in unpaired rats ($p < 0.01$), and this response was abolished by the preceding experience of the aversive drug cue in paired rats ($p = 0.70$). Black lines show average concentration over time and red lines indicate SE for each group. Bars depict average peak DA from the baseline and tone epochs. *c*, Rasters and histograms depict examples of inhibitory (left) and excitatory (right) neural encoding of the tone. Individual action potentials during each tone presentation are shown in the raster, and firing rate (Hz) is shown in the histogram below. *d*, Categorization of responses to the tone in all recorded neurons. Paired neurons (right) show more excitatory encoding in the core relative to the shell at test ($p < 0.04$), but this difference in excitatory encoding was not apparent in paired neurons during extinction (center; $p = 0.63$) or in unpaired neurons (left; $p = 0.62$).

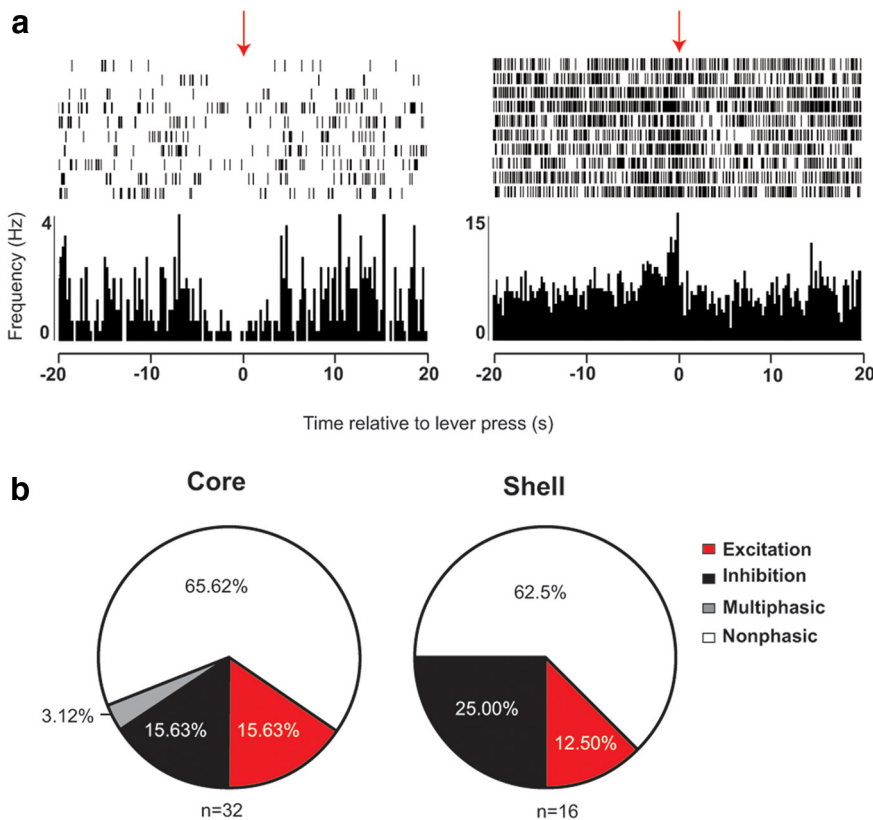


Figure 6. NAc neurons encode responses during reinstatement of cocaine seeking. **a**, Representative inhibitory (left) and excitatory (right) encoding of lever press at test. Individual action potentials aligned to each lever press (red arrows) are shown in the raster, and firing rate (Hz) is shown in the histogram below. **b**, Categorization of lever-press encoding in paired neurons at test.

atory in nature ($n = 35/60$) rather than inhibitory ($n = 25/60$). This ratio was reversed ($p < 0.01$) in the unpaired group, in which encoding was predominantly inhibitory ($n = 28/40$) rather than excitatory ($n = 12/40$). Figure 4*d* depicts the proportion of responsive neurons in each subregion.

The aversive drug cue eliminates DA encoding of the extinguished proximate drug cue

The tone was presented after saccharin exposure to determine whether reinstatement behavior co-occurs with renewed DA encoding of the extinguished proximate drug cue. The results revealed the opposite (Fig. 5*a,b*; group \times epoch: $F_{(1,6)} = 6.30$, $p < 0.05$). The proximate drug cue caused a rapid, transient increase in DA in the unpaired group ($p < 0.01$), but this DA response was not observed in the paired group ($p > 0.70$). Surprisingly, these data suggest that dampened dopamine signaling caused by aversive cocaine-predictive saccharin persists during the time in which an animal transitions to drug seeking and diminishes the dopaminergic encoding of other drug predictive cues.

The aversive drug cue augments excitatory neural encoding of the extinguished proximate drug cue

Considering the suppression of DA encoding of the tone in the paired group, the neural encoding of this stimulus was of particular interest. The tone was encoded by 27.78% ($n = 20/72$) of paired neurons on the last day of extinction and 25.37% ($n = 17/67$) of paired neurons at test. In the unpaired group, 22.0% ($n = 16/73$) of neurons encoded the tone at test, indicating no effect of group or treatment on the overall proportion of tone-encoding neurons in the NAc ($p > 0.40$). However, the nature of these phasically active neurons (excitations or inhibi-

tions) varied based on the treatment and brain region (Fig. 5*c,d*). Tone-encoding neuronal responses were roughly evenly distributed between excitations and inhibitions in the unpaired condition (excitation/inhibition ratio: core, $n = 4/3$; shell, $n = 3/6$; $p = 0.62$), as well as the paired condition during extinction (core, $n = 3/4$; shell, $n = 6/7$; $p = 0.63$). However, the presence of the aversive, drug predictive cue induced a significant shift toward excitatory responses in the NAcC in the paired group at test ($n = 7/0$) relative to the NAcS ($n = 4/6$; $p < 0.04$). Thus, a shift toward excitatory encoding co-occurred with the suppression of DA encoding.

The examination of neural encoding of drug-seeking was restricted to rats that registered enough lever presses to produce a reliable histogram, which limited the analysis to rats in the paired group at test ($n = 5$). Overall, 34.38% ($n = 11/32$) NAcC and 37.5% ($n = 6/16$) NAcS neurons encoded one or more aspects (pre- and/or postresponse) of the lever-press behavior. Of the phasically active NAcC neurons, 54.55% ($n = 6/11$) exhibited some form of excitatory encoding, and 54.55% ($n = 6/11$) exhibited inhibitory encoding (1 neuron exhibited a phasic decrease before and a phasic increase after the response), whereas NAcS neuronal responses were 33.33% ($n = 2/6$) excitatory and 66.67% ($n = 4/6$) inhibitory (Fig. 6*a,b*).

The aversive drug cue reduces baseline DA and enhances the baseline excitability of neurons that will encode future drug-seeking behavior

In the NAcS, an aversive drug cue causes a long term, diffuse depression of DA concentration, as well as an acute time-locked DA decrease (Wheeler et al., 2011). The presence of the diffuse depression might be especially important for reinstatement of drug seeking (Twining et al., 2014). We tested for the presence of this phenomenon here by performing an analysis of baseline (10 s preinfusion) DA concentration in the NAcC across trials during the saccharin exposure period (Fig. 7*a*). The baseline period was blocked into early and late session DA concentration. One subject in the paired group lacked sufficient data to construct a complete time course and could not be included in this analysis, but it was clear that baseline DA concentration changed in a non-time-locked manner across trials differentially depending on group (Fig. 7*c*; group \times trial block: $F_{(1,5)} = 22.39$, $p < 0.01$; all other p values > 0.20). Planned comparisons revealed that appetitive, unpaired saccharin increased basal DA concentration across trials ($p < 0.02$), and aversive, cocaine-predictive saccharin decreased basal DA concentration across trials ($p < 0.03$).

Because fluctuations in DA can modulate neural excitability (Surmeier et al., 2007), we were interested in how diffuse changes in DA might relate to the baseline excitability of NAc neurons. Furthermore, we isolated neurons that would later go on to encode the proximate drug cue or drug-seeking behavior in the next phase of testing to determine whether paired saccharin differentially impacted the behavior of these neurons (Fig. 7*b*). Because of

the limited number of previously categorized neurons, both categories of responses (inhibitory and excitatory) were combined for this analysis. Changes in baseline firing rates were assessed by blocking the saccharin period into thirds and generating three average baseline firing rate measurements for each neuron. An ANOVA revealed a significant interaction between all factors (trial block \times group \times subregion: $F_{(2,56)} = 4.20$, $p = 0.02$) and a marginal interaction between group and trial block ($F_{(1,25)} = 3.87$, $p = 0.05$) but no other effects ($p > 0.17$). A *post hoc* analysis showed that the aversive cocaine cue caused an increase in the baseline firing rates from the beginning to the end of the saccharin period specifically in NAcC neurons that would later encode the tone cue (Fig. 7*d*; $p < 0.05$; all other p values > 0.80). Notably, all of the paired core neurons that increased their baseline firing rates were recorded in subjects that exhibited reinstatement (reinstatement ratios > 0.50), whereas the single neuron that did not increase firing rate across the saccharin session was recorded in a subject that did not show appreciable aversive taste reactivity (< 0.50 aversive events/trial) or reinstatement.

Similar to tone-encoding neurons in the core, neurons that encoded drug seeking exhibited a change in baseline activity during exposure to the aversive saccharin stimulus (Fig. 7*e*). In this case, baseline firing rates in both the core and shell neurons increased during the saccharin period ($F_{(2,18)} = 4.51$, $p < 0.03$; all other p values > 0.09). Importantly, an analysis of all neurons (including nonencoding neurons) showed that most baseline firing rates decreased during the saccharin period regardless of group (Fig. 7*f*; $F_{(2,420)} = 4.88$, $p < 0.01$), and baseline activity in the paired group was generally higher than the unpaired group ($F_{(1,210)} = 7.37$, $p < 0.01$; all other p values > 0.07). Considering the general downward trend in baseline activity, the increase in firing rate in neurons that encode drug seeking or the proximate drug cue is exceptional.

Discussion

The present findings show that a natural reward that signals delayed access to cocaine becomes aversive and reinstates drug seeking after extinction. The presence of this aversive drug cue decreases acute and diffuse DA release in the NAcC and increases the baseline firing rates of neurons that will subsequently encode the proximate cocaine cue and cocaine-seeking behavior. Furthermore, the aversive drug cue also eliminates the dopaminergic encoding of the proximate drug cue. This demonstrates a pronounced depression of DA signaling that persists long after the aversive drug cue is removed. At the same time, excitatory encoding of the proximate cue is enhanced specifically in the NAcC. This relationship between

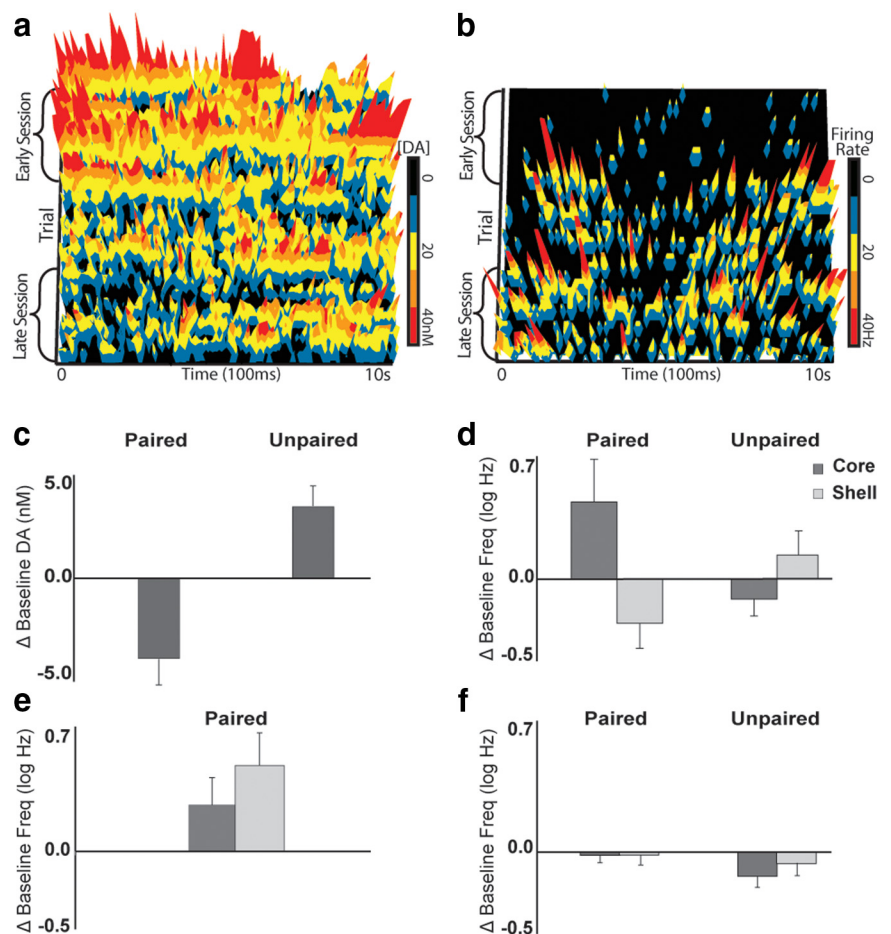


Figure 7. The aversive drug cue reduces DA and enhances excitability of neurons that will encode future drug-seeking behavior. *a*, Baseline DA (nM; z-axis) across trials (y-axis) throughout the saccharin period in a representative paired rat. *b*, Baseline firing rate (Hz; z-axis) across trials (y-axis) throughout the saccharin period in a representative tone-encoding neuron from a paired rat. *c*, Difference scores (late–early) of average baseline DA concentrations (nM) in paired and unpaired rats. Saccharin reduced baseline [DA] over time in paired rats ($p < 0.03$), and increased baseline [DA] in unpaired rats ($p < 0.02$). *d*, The change in baseline firing rates (log Hz) in units that subsequently encoded the tone. In the paired group, saccharin increased firing rates in tone responsive units in the NAcC ($p < 0.05$), but had no effect in the NAcS or in unpaired rats (p values > 0.80). *e*, The change in baseline firing rates in lever press response units in paired rats. Saccharin increased baseline firing rates in both the NAcC and NAcS ($p < 0.03$). *f*, The change in baseline firing rates in all recorded units before saccharin infusion in paired and unpaired rats. Saccharin reduced baseline firing rates in the full population of recorded neurons ($p < 0.01$).

dampened DA signaling and enhanced neural excitability provides insight into how negative affect can drive compensatory drug-seeking behavior.

Behavioral design

The present design differs from studies of noncontingent drug cue- and context-induced reinstatement (Crombag and Shaham, 2002; Saunders et al., 2013) because it allows for direct assessment of the affective state of the animal during cue exposure. It differs from cue-induced reinstatement designs in which subjects receive response-contingent presentations of an unextinguished drug cue (Davis and Smith, 1976) because it is not directly driven by the conditioned incentive value of the saccharin cue (which is presumably punishing). As such, it is capable of providing unique insight to the physiological mechanisms underlying the induction of negative affect by drug cues, and is useful for modeling cue-induced changes in affective state that may promote relapse in human addicts.

The present design resembles other context-induced reinstatement models in that the extinction context (unpaired saccharin in our design) could be an omission cue. Thus, the critical

behavioral and physiological differences between the groups could be due to the presence of an omission signal suppressing behavior in the unpaired group, rather than the presence of an aversive drug cue in the paired group inducing behavior. Several observations make this unlikely. First, both groups extinguished at the same rate, suggesting that the unpaired cue was not being used as an omission signal to guide behavior. Also, electrophysiological responses to the tone did not differ during extinction in the paired and unpaired groups, indicating that the presence of a potential omission signal did not influence encoding of the proximate cue. Finally, recent findings indicate that the presence of an innately aversive taste stimulus is sufficient to cause drug seeking. We observed a similar level of reinstatement following exposure to quinine, an inherently aversive stimulus that had no association with drug access (Twining et al., 2014). Similarly, in the present study the magnitude of the reinstatement effect was positively correlated with the expression of aversion. Together these observations indicate that it is the aversive status of the cocaine cue, and the negative affect and neurochemical environment that it engenders, that induce drug seeking.

The impact of an aversive drug cue on DA encoding of a proximate drug cue

The observation of persistent dopaminergic encoding of an extinguished drug cue in the unpaired group may be surprising, but it is consistent with contemporary theory regarding the product of extinction. Extinction produces suppression of behavior, but without direct physiological intervention (Lee et al., 2006; Otis et al., 2013) or perhaps precise timing (Monfils et al., 2009), it does not result in a loss of learning (Bouton, 2004; Miller and Matzel, 2006). Neural processing of an extinguished drug cue is an interaction between memories of reinforcement and nonreinforcement, and the DA response likely reflects this ambiguity. As such, it is not surprising that brain chemistry is more reactive to an extinguished drug cue than a nonassociated behaviorally inert stimulus, which does not typically elevate NAc DA (Phillips et al., 2003).

The fact that reinstatement behavior was accompanied by a suppression of the dopaminergic encoding of a proximate drug-cue was surprising. Even so, the present result is more intuitive when informed by observations of DA signaling during cocaine self-administration. For example, rats that exhibit escalated drug use actually show attenuated DA encoding of drug-taking behavior (Willuhn et al., 2014). Furthermore, drug taking is reduced when positive DA encoding is restored through pharmacological intervention. That is, rats seek more cocaine because DA encoding is depressed, and reduce responding when it is restored (Willuhn et al., 2014). In this light, it is provocative to consider the possibility that if the acute DA elevation mediates the suppression of drug taking, then it is possible that positive DA signaling also works to suppress drug seeking after extinction. The aversive drug cue may spur behavior, in part by eliminating this neurochemical event. This explanation is consistent with the observation that vigorous cocaine seeking/taking behavior tends to be preceded by reductions in NAc DA (Wise et al., 1995; Tsibulsky and Norman, 1999; Stuber et al., 2005).

In contrast to the present results, it is important to recognize that some aversive stimuli can increase DA and reinstate drug seeking. For example, electric footshock increases NAc DA as measured by microdialysis, and this increase in DA is sustained as drug seeking is reinstated (McFarland et al., 2004). There are many factors that may contribute to this discrepancy (Wenzel et al., 2015). The nociceptive stimulation produced by electrical

footshock is typically very brief, discrete, and naturally engages robust locomotor behaviors that facilitate escape from the experimental context. In contrast, the taste infusions used here are relatively long, occur over an extended period of time, and induce behaviors of gustatory rejection that involve little locomotion (culminating with passive rejection). It is noteworthy that aversive gustatory stimuli can elevate DA in the NAc (Bassareo et al., 2002), indicating that modality is not the only critical factor. Regardless of the root cause(s) of this discrepancy, it is important to examine these different situations because both can lead to enhanced drug seeking, and may do so through dissociable mechanisms.

The role of NAc neural and neurochemical signaling in drug seeking

Broadly, activity in the NAc mediates reinstatement phenomena (Bossert et al., 2013). Disrupting activity in the NAc attenuates stress-, cue-, cocaine-, and context-induced reinstatement (Cornish and Kalivas, 2000; Fuchs et al., 2004, 2008; McFarland et al., 2004; Xie et al., 2012). Specifically, corticostriatal glutamatergic projections drive relapse phenomena (McFarland et al., 2003, 2004; Stefanik et al., 2013). In the present experiments, reinstatement was associated with enhanced excitability in two ways. First, exposure to the aversive drug cue elevated the baseline excitatory activity of neurons that would eventually encode the proximate drug cue and/or drug-seeking behavior. This observation reveals a physiological mechanism that directly connects cue-induced negative affect to the subsequent drug-seeking act. Second, the aversive drug cue caused drug seeking and enhanced excitatory encoding of the proximate drug cue. This change in cue reactivity could contribute to the reinstated drug seeking observed here and in studies that show behavioral synergy between aversive stimuli and proximate drug cues (Liu and Weiss, 2002; Buffalari and See, 2009), even if the proximate drug cue has been extinguished (Shelton and Beardsley, 2005).

The relationship between decreased DA signaling and a shift toward excitatory activity

The effect of DA on striatal neural activity is complex and depends on a number of factors, such as D1-like and D2-like receptor expression, as well as the collateral influence of DA-sensitive interneurons (Surmeier et al., 2007). In general, D1-like receptor activation enhances excitability (Surmeier et al., 1995), appetitive stimuli preferentially activate D1-expressing neurons (Xiu et al., 2014), and excitatory encoding of motivated behavior coincides with and is sometimes contingent upon elevated DA signaling in the NAc (Owesson-White et al., 2009; Cacciapaglia et al., 2011). In contrast, D2-like receptor activation inhibits neuron excitability (Hernandez-Lopez et al., 2000), D2-expressing neurons are preferentially activated by aversive stimuli (Xiu et al., 2014), and acute inhibition of DA signaling induces aversion by activating D2-expressing neurons in the NAc (Danjo et al., 2014). Given our coincident observations of decreased dopamine signaling, increased NAc activity, and behavioral signs of negative affect, we hypothesize that the current results reflect a disinhibited D2-mediated aversion-sensitive striatal circuit.

The product of aversion-induced neural activation has been linked to behavioral inhibition (i.e., no-go; Nakanishi et al., 2014), but activation of D2-like-expressing neurons is also important for directing new behaviors that avoid aversive outcomes (Mehta et al., 2001; Yawata et al., 2012; Porter-Stransky et al., 2013). This convergent evidence strongly suggests that cue- and aversion-induced reductions in DA signaling play a critical role in enhancing the excitability of neurons that encode proximate drug

cues and drug-seeking behavior. The interaction between negative affect, dampened DA signaling, and enhanced neural excitability highlights a physiological substrate by which negative reinforcement mechanisms can augment cue reactivity and drive behavior. This aversion-driven behavioral modulation could be directed toward escape in adaptive behavioral contexts, but in a drug-seeking context, it may promote drug seeking in an effort to correct the negative affective state. Future experiments will be needed to investigate the precise causal role of physiologically relevant DA reductions in initiating this maladaptive cascade of events.

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