Persistent coding of outcome-predictive cue features in the rat nucleus accum-

bens.

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1 Abstract

outcome associations to inform future behavior.

The nucleus accumbens (NAc) is important for learning from feedback, and for biasing and invigorating behavior in response to cues that predict motivationally relevant outcomes. NAc encodes outcome-related cue features such as the magnitude and identity of reward. However, little is known about how features of cues themselves are encoded. We designed a decision making task where rats learned multiple sets of outcome-predictive cues, and recorded single-unit activity in the NAc during performance. We found that coding of cue identity and location occurred alongside coding of expected outcome. Furthermore, this coding persisted both during a delay period, after the rat made a decision and was waiting for an outcome, and after the outcome was revealed. Encoding of cue features in the NAc may enable contextual modulation of ongoing behavior, and provide an eligibility trace of outcome-predictive stimuli for updating stimulus-

12 Introduction

- Theories of nucleus accumbens (NAc) function generally agree that this brain structure contributes to motivated behavior, with some emphasizing a role in learning from reward prediction errors (RPEs) (Averbeck & Costa 2017; Joel et al. 2002; Khamassi & Humphries 2012; Lee et al. 2012; Maia 2009; Schultz 2016; see also the addiction literature on effects of drug rewards; Carelli 2010; Hyman et al. 2006; Kalivas & Volkow 2005), and others a role in the modulation of ongoing behavior through stimuli associated with motivationally relevant outcomes (invigorating, directing; Floresco, 2015; Nicola, 2010; Salamone & Correa, 2012). These proposals echo similar ideas on the functions of the neuromodulator dopamine (Berridge, 2012; Maia, 2009; Salamone & Correa, 2012; Schultz, 2016), with which the NAc is tightly linked functionally as well as anatomically (Cheer et al., 2007; du Hoffmann & Nicola, 2014; Ikemoto, 2007; Takahashi et al., 2016).
- Much of our understanding of NAc function comes from studies of how cues that predict motivationally relevant outcomes (e.g. reward) influence behavior and neural activity in the NAc. Task designs that associate such cues with rewarding outcomes provide a convenient access point, eliciting conditioned responses such as sign-tracking and goal-tracking (Hearst & Jenkins, 1974; Robinson & Flagel, 2009), Pavlovian-instrumental transfer (Estes, 1943; Rescorla & Solomon, 1967) and enhanced response vigor (Nicola, 2010; Niv et al., 2007), which tend to be affected by NAc manipulations (Chang et al. 2012; Corbit & Balleine 2011; Flagel et al. 2011; although not always straightforwardly; Chang & Holland 2013; Giertler et al. 2004). Similarly, analysis of RPEs typically proceeds by establishing an association between a cue and sub-sequent reward, with NAc responses transferring from outcome to the cue with learning (Day et al., 2007; Roitman et al., 2005; Schultz et al., 1997; Setlow et al., 2003).
- Surprisingly, although substantial work has been done on the coding of outcomes predicted by such cues
 (Atallah et al., 2014; Bissonette et al., 2013; Cooch et al., 2015; Cromwell & Schultz, 2003; Day et al., 2006;
 Goldstein et al., 2012; Hassani et al., 2001; Hollerman et al., 1998; Lansink et al., 2012; McGinty et al., 2013;
 Nicola, 2004; Roesch et al., 2009; Roitman et al., 2005; Saddoris et al., 2011; Schultz et al., 1992; Setlow et

al., 2003; Sugam et al., 2014; West & Carelli, 2016), much less is known about how outcome-predictive cues themselves are encoded in the NAc (but see; Sleezer et al., 2016). This is an important issue for at least two 37 reasons. First, in reinforcement learning, motivationally relevant outcomes are typically temporally delayed 38 relative to the cues that predict them. In order to solve the problem of assigning credit (or blame) across such temporal gaps, some trace of preceding activity needs to be maintained (Lee et al., 2012; Sutton & Barto, 40 1998). For example, if you become ill after eating food X in restaurant A, depending on if you remember the 41 identity of the restaurant or the food at the time of illness, you may learn to avoid all restaurants, restaurant 42 A only, food X only, or the specific pairing of X-in-A. Therefore, a complete understanding of what is learned following feedback requires understanding what trace is maintained. Since NAc is a primary target 44 of dopamine signals interpretable as RPEs, and NAc lesions impair RPEs related to timing, its activity trace will help determine what can be learned when RPEs arrive (Hamid et al., 2015; Hart et al., 2014; Ikemoto, 46 2007; McDannald et al., 2011; Takahashi et al., 2016). Similarly, in a neuroeconomic framework, NAc is thought to represent a domain-general subjective value signal for different offers (Bartra et al., 2013; Levy 48 & Glimcher, 2012; Peters & Büchel, 2009; Sescousse et al., 2015); having a representation of the offer itself alongside this value signal would provide a potential neural substrate for updating offer value.

Second, for ongoing behavior, the relevance of cues typically depends on context. In experimental settings, context may include the identity of a preceding cue, spatial or configural arrangements (Bouton, 1993;
Holland, 1992; Honey et al., 2014), and unsignaled rule changes, as occurs in set shifting and other cognitive control tasks (Cohen & Servan-Schreiber, 1992; Floresco et al., 2006; Grant & Berg, 1948; Sleezer et
al., 2016). In such situations, the question arises how selective, context-dependent processing of outcomepredictive cues is implemented. For instance, is there a gate prior to NAc such that only currently relevant
cues are encoded in NAc, or are all cues represented in NAc but their current values dynamically updated
(FitzGerald et al., 2014; Goto & Grace, 2008; Sleezer et al., 2016)? Representation of cue identity would
allow for context-dependent mapping of outcomes predicted by specific cues.

60 Thus, both from a learning and a flexible performance perspective, it is of interest to determine how cue iden-

tity is represented in the brain, with NAc of particular interest given its anatomical and functional position at the center of motivational systems. We sought to determine whether cue identity is represented in the NAc, if cue identity is represented alongside other motivationally relevant variables, such as cue outcome, and if these representations are maintained after a behavioral decision has been made (see Figure 1 for a schematic representation of the specific hypotheses tested). To address these questions, we recorded the activity of NAc units as rats performed a task in which multiple, distinct sets of cues predicted the same outcome.

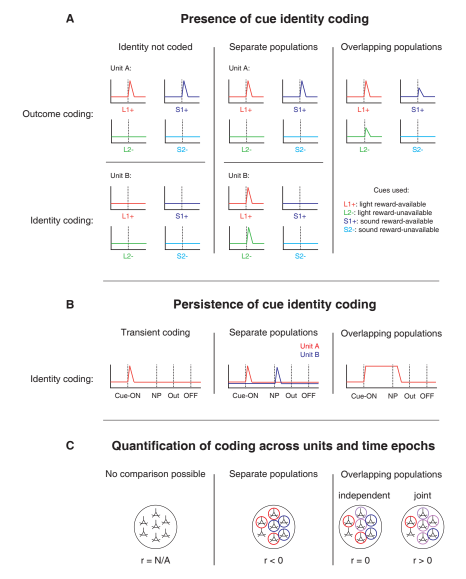


Figure 1: Schematic of hypothetical coding scenarios for cue feature coding employed by single units in the NAc across different cue features (A) and phases of a trial (B). A: Schematic peri-event time histograms (PETHs) illustrating putative responses to different cues under different hypotheses of how cue identity (light, sound; L, S) and outcome (reward-available, reward-unavailable; +, -) are coded. Left panel: Coding of identity is absent in the NAc. Top: Unit A encodes a motivationally relevant variable, such as expected outcome, similarly across other cue features, such as identity or physical location. Hypothetical plot is firing rate across time. L1+ (red) signifies a reward-available light cue, S1+ (navy blue) a reward-available sound cue, L2- (green) a reward-unavailable light cue, S2- (light blue) a reward-unavailable sound cue. Dashed line indicates onset of cue. Bottom: No units within the NAc discriminate their firing according to cue identity. Middle panel: Coding of identity occurs in a separate population of units from coding of other cue features such as expected outcome or physical location. Top: Same as left panel, with unit A discriminating between reward-available and reward-unavailable cues. Bottom: Unit B discriminates firing across stimulus modalities, depicted here as firing to light cues but not sound cues. Note lack of coding overlap in both units. Right panel: Coding of identity occurs in an overlapping population of cells with coding of other motivationally relevant variables. Hypothetical example demonstrating a unit that responds to reward-available cues, but firing rate is also modulated by the stimulus modality of the cue, firing most for the reward-available light cue. B: Schematic PETHs illustrating potential ways in which identity coding may persist over time. Left panel: Cue-onset triggers a transient response to a unit that codes for cue identity. Dashed lines indicate time of a behavioral or environmental event. 'Cue-ON' signifies cue-onset, 'NP' signifies nosepoke at a reward receptacle, 'Out' signifies when the outcome is revealed, 'OFF' signifies cue-offset. Middle and right panel: Identity coding persists at other time points, shown here during a nosepoke hold period until outcome is revealed. Coding can either be maintained by a sequence of units (middle panel) or by the same unit as during cue-onset (right panel). C: Schematic pool of NAc units, illustrating different analysis outcomes that discriminate between hypotheses. R values represent the correlation between sets of recoded regression coefficients (see text for analysis details). Left panel: Cue identity is not coded (A: left panel), or is only transiently represented in response to the cue (B: left panel). Middle panel: Negative correlation (r < 0) suggests that identity and outcome coding are represented by separate populations of units (A: middle panel), or identity coding is represented by distinct units across different points in a trial (B: middle panel). Red circles represents coding for one cue feature or point in time, blue circles for the other cue feature or point in time. Right panel: Identity and outcome coding (A: right panel), or identity coding at cue-onset and nosepoke (B: right panel) are represented by overlapping populations of units, shown here by the purple circles. The absence of a correlation (r = 0) suggests that the overlap of identity and outcome coding, or identity coding at cue-onset and nosepoke, is expected by chance and that the two cue features, or points in time, are coded by overlapping but independent populations from one another. A positive correlation (r > 0) implies a higher overlap than expected by chance, suggesting coding by a joint population. Note: The same logic applies to other aspects of the environment when the cue is presented, such as the physical location of the cue, as well as other time epochs within the task, such as when the animal receives feedback about an approach.

Results

8 Behavior

Rats were trained to discriminate between cues signaling the availability and absence of reward on a square track with four identical arms for two distinct set of cues (Figure 2A). During each session, rats were pre-70 sented sequentially with two behavioral blocks containing cues from different sensory modalities, a light and 71 a sound block, with each block containing a cue that signaled the availability of reward (reward-available), 72 and a cue that signaled the absence of reward (reward-unavailable). To maximize reward receipt, rats should 73 approach reward sites on reward-available trials, and skip reward sites on reward-unavailable trials (see Fig-74 ure 2B for an example learning curve). All four rats learned to discriminate between the reward-available and 75 reward-unavailable cues for both the light and sound blocks as determined by reaching significance (p < .05) 76 on a daily chi-square test comparing approach behavior for reward-available and reward-unavailable cues for 77 each block, for at least three consecutive days (range for time to criterion: 22 - 57 days). Maintenance of behavioral performance during recording sessions was assessed using linear mixed effects models for pro-79 portion of trials where the rat approached the receptacle. Analyses revealed that the likelihood of a rat to 80 make an approach was influenced by whether a reward-available or reward-unavailable cue was presented, 81 but was not significantly modulated by whether the rat was presented with a light or sound cue (Percent-82 age approached: light reward-available = 97%; light reward-unavailable = 34%; sound reward-available = 83 91%; sound reward-unavailable 35%; cue identity p = .115; cue outcome p < .001; Figure 2C). Additional analyses separated each block into two halves to assess possible within session learning. Adding block half 85 into the model did not improve prediction of behavioral performance (p = .86), arguing against within session learning. Thus, rats successfully discriminated the cues according to whether or not they signaled the 87 availability of reward at the reward receptacle.

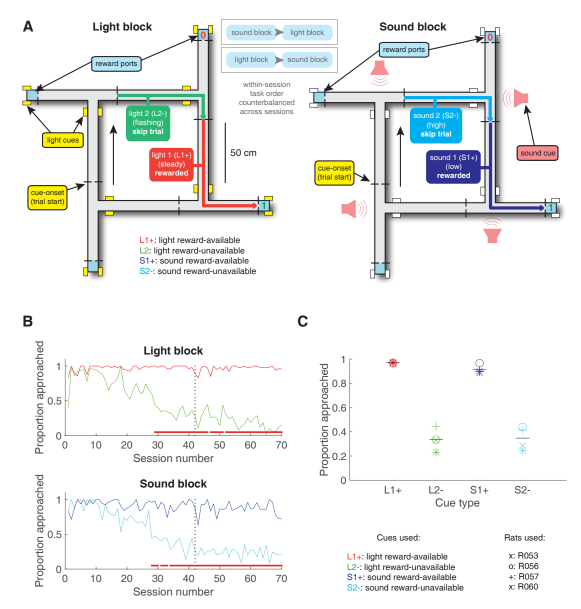


Figure 2: Schematic and performance of the behavioral task. A: Apparatus was a square track consisting of multiple identical T-choice points. At each choice point, the availability of 12% sucrose reward at the nearest reward receptacle (light blue fill) was signaled by one of four possible cues, presented when the rat initiated a trial by crossing a photobeam on the track (dashed lines). Photobeams at the ends of the arms by the receptacles registered nosepokes. Arrows outside of track indicate correct running direction. Left: light block showing an example trajectory for a correct reward-available (approach trial; red) and reward-unavailable (skip trial; green) trial. Rectangular boxes with yellow fill indicate location of LEDs used for light cues. Right: sound block with a correct reward-available (approach trial; navy blue) and reward-unavailable (skip trial; light blue) trial. Speakers for sound cues were placed underneath the choice points, indicated by magenta speaker icons. Ordering of the light and sound blocks was counterbalanced across sessions. Reward-available and reward-unavailable cues were presented pseudo-randomly, such that not more than two of the same type of cue could be presented in a row. Location of the cue on the track was irrelevant for behavior, all cue locations contained an equal amount of reward-available and reward-unavailable trials. B-C: Performance on the behavioral task. B. Example learning curves across sessions from a single subject (R060) showing the proportion of trials approached for reward-available (red line for light block, navy blue line for sound block) and reward-unavailable trials (green line for light block, light blue line for sound block) for light (top) and sound (bottom) blocks. Fully correct performance corresponds to an approach proportion of 1 for reward-available trials and 0 for reward-unavailable trials. Rats initially approach on both reward-available and reward-unavailable trials, and learn with experience to skip reward-unavailable trials. Red bars indicate days in which a rat statistically discriminated between reward-available and rewardunavailable cues, determined by a chi square test. Dashed line indicates time of electrode implant surgery. C: Proportion of trials approached for each cue, averaged across all recording sessions and shown for each rat. Different columns indicate the different cues (reward-available (red) and reward-unavailable (green) light cues, reward-available (navy blue) and reward-unavailable (light blue) sound cues). Different symbols correspond to individual subjects; horizontal black line shows the mean. All rats learned to discriminate between reward-available and reward-unavailable cues, as indicated by the clear difference of proportion approached between rewardavailable (\sim 90% approached) and reward-unavailable cues (\sim 30% approached), for both blocks (see Results for statistics).

NAc encodes behaviorally relevant and irrelevant cue features

We sought to address which parameters of our task were encoded by NAc activity, specifically whether the NAc encodes aspects of motivationally relevant cues not directly tied to reward, such as the identity 91 and location of the cue, and whether this coding is accomplished by separate or overlapping populations 92 (Figure 1A). We recorded a total of 443 units with > 200 spikes in the NAc from 4 rats over 57 sessions 93 (Table 1; range: 12 - 18 sessions per rat) while they performed a cue discrimination task (Figure 2A). Units that exhibited a drift in firing rate over the course of either block, as measured by a Mann-Whitney U test 95 comparing firing rates for the first and second half of trials within a block, were excluded from further analysis, leaving 344 units for further analysis. The activity of 133 (39%) of these 344 units were modulated 97 by the cue, as determined by comparing 1 s pre- and post-cue activity with a Wilcoxon signed-rank test, with more showing a decrease in firing (n = 103) than an increase (n = 30) around the time of cue-onset 99 (Table 1). Within this group, 24 were classified as putative fast spiking interneurons (FSIs), while 109 were classified as putative medium spiny neurons (MSNs). Upon visual inspection, we observed several patterns 101 of firing activity, including units that discriminated firing upon cue-onset across various cue conditions, 102 showed sustained differences in firing across cue conditions, had transient responses to the cue, showed 103 a ramping of activity starting at cue-onset, and showed elevated activity immediately preceding cue-onset 104 (Figure 3, 3-supplement 1, 3-supplement 2). 105

Task parameter	Total	↑ MSN	\downarrow MSN	↑ FSI	\downarrow FSI
All units	443	155	216	27	45
Rat ID					
R053	145	51	79	4	11
R056	70	12	13	17	28
R057	136	55	75	3	3
R060	92	37	49	3	3
Analyzed units	344	117	175	18	34
Cue modulated units	133	24	85	6	18
GLM aligned to cue-onset					
Cue identity	42 (32%)	9 (38%)	25 (29%)	0 (-)	8 (44%)
Cue location	55 (41%)	11 (46%)	33 (39%)	3 (50%)	8 (44%)
Cue outcome	26 (20%)	5 (21%)	15 (18%)	1 (17%)	5 (28%)
Approach behavior	32 (24%)	8 (33%)	19 (22%)	2 (33%)	3 (17%)
Trial length	22 (17%)	5 (21%)	14 (16%)	0 (-)	3 (17%)
Trial number	42 (32%)	11 (46%)	20 (24%)	1 (17%)	10 (56%)
Trial history	8 (6%)	1 (4%)	5 (6%)	0 (-)	1 (6%)
GLM aligned to nosepoke					
Cue identity	28 (21%)	3 (13%)	17 (20%)	2 (33%)	6 (33%)
Cue location	30 (23%)	2 (8%)	21 (25%)	2 (33%)	5 (28%)
Cue outcome	23 (17%)	2 (8%)	14 (16%)	1 (17%)	6 (33%)
GLM aligned to outcome					
Cue identity	25 (19%)	4 (17%)	15 (18%)	2 (33%)	4 (22%)
Cue location	31 (23%)	5 (21%)	23 (27%)	0 (-)	3 (17%)
Cue outcome	34 (26%)	6 (25%)	15 (18%)	4 (67%)	9 (50%)

Table 1: Overview of recorded NAc units and their relationship to task variables at various time epochs. Percentage is relative to the number of cue-modulated units (n = 133).

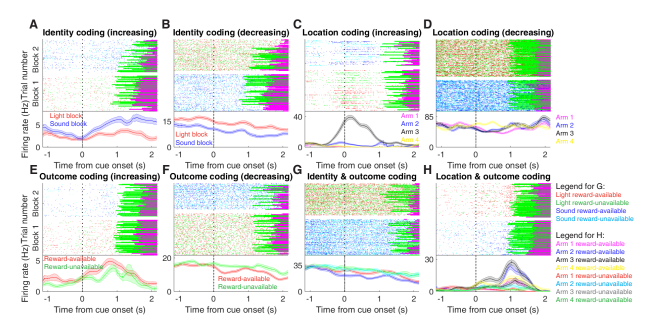


Figure 3: Examples of cue-modulated NAc units influenced by different task parameters. A: Example of a cue-modulated NAc unit that showed an increase in firing following the cue, and exhibited identity coding. Top: rasterplot showing the spiking activity across all trials aligned to cue-onset. Spikes across trials are color-coded according to cue type (red: reward-available light; green: reward-unavailable light; navy blue: reward-available sound; light blue: reward-unavailable sound). Green and magenta bars indicate trial termination when a rat initiated the next trial or made a nosepoke, respectively. White space halfway up the rasterplot indicates switching from one block to the next. Dashed line indicates cue-onset. Bottom: PETHs showing the average smoothed firing rate for the unit for trials during light (red) and sound (blue) blocks, aligned to cue-onset. Lightly shaded area indicates standard error of the mean. Note this unit showed a larger increase in firing to sound cues. B: An example of a unit that was responsive to cue identity as in A, but for a unit that showed a decrease in firing to the cue. Note the sustained higher firing rate during the light block. C-D: Cue-modulated units that exhibited location coding. Each color in the PETHs represents average firing response for a different cue location. C: The firing rate of this unit only changed on arm 3 of the task. D: Firing rate decreased for this unit on all arms but arm 4. E-F: Cue-modulated units that exhibited outcome coding, with the PETHs comparing reward-available (red) and reward-unavailable (green) trials. E: This unit showed a slightly higher response during presentation of reward-available cues. F: This unit showed a dip in firing when presented with reward-available cues. G-H: Examples of cue-modulated units that encoded multiple cue features. G: This unit showed both identity and outcome coding. H: An example of a unit that coded for both identity and location.

To characterize more formally whether these cue-modulated responses were influenced by various aspects of 106 the task, we fit a sliding window generalized linear model (GLM) to the firing rate of each cue-modulated unit 107 surrounding cue-onset, using a forward selection stepwise procedure for variable selection, a bin size of 500 108 ms for firing rate and a step size of 100 ms for the sliding window. Fitting GLMs to all trials within a session 109 revealed that a variety of task parameters accounted for a significant portion of firing rate variance in NAc 110 cue-modulated units (Figure 4A, 4-supplement 1, 4-supplement 2, Table 1). Notably, a significant proportion 111 of units discriminated between the light and sound block (identity coding: \sim 32% of cue-modulated units, 112 accounting for ~5% of firing rate variance) or the arms of the apparatus (location coding: ~41% of cue-113 modulated units, accounting for ~4% of firing rate variance) throughout the entire window surrounding 114 cue-onset. Additionally, a substantial proportion of units discriminating between the common portion of reward-available and reward-unavailable trials (outcome coding: $\sim 20\%$ of cue-modulated units, accounting 116 for \sim 4% of firing rate variance) was not observed until after the onset of the cue (z-score > 1.96 when comparing observed proportion of units to a shuffled distribution obtained when shuffling the firing rates of 118 each unit across trials before running the GLM). Furthermore, our variable selection method ensured that the observed coding was not due to potential confounds from other task variables, such as behavioral response at 120 the choice point (approach behavior; left vs. right), variability in response vigor (trial length; see McGinty et al. 2013), drift due to the passage of time (trial number), and the pseudorandom nature of cue presentation 122 (trial history). In addition to accounting for firing rate variance explained due to whether the rat turned left 123 or right, we ran our cue-onset GLM using only approach trials, and found a similar proportion of outcome 124 coding units (34 units; \sim 26% of cue-modulated units), providing further support that these units were coding 125 the expected outcome of the cue. Taken together, these results from the GLMs suggest that the NAc encodes 126 features of outcome-predictive cues in addition to expected outcome.

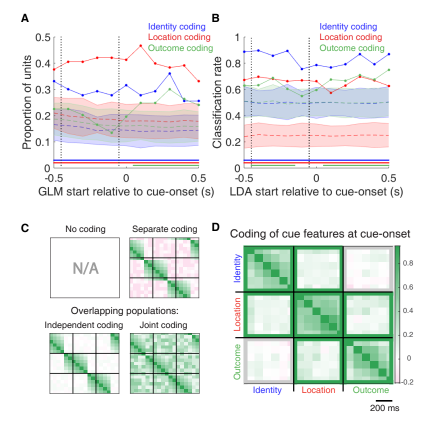


Figure 4: Summary of influence of cue features on cue-modulated NAc units at time points surrounding cue-onset. A: Sliding window GLM (bin size: 500 ms; step size: 100 ms) demonstrating the proportion of cue-modulated units where cue identity (blue solid line), location (red solid line), and outcome (green solid line) significantly contributed to the model at various time epochs relative to cueonset. Dashed colored lines indicate the average of shuffling the firing rate order that went into the GLM 100 times. Error bars indicate 1.96 standard deviations from the shuffled mean. Solid lines at the bottom indicate when the proportion of units observed was greater than the shuffled distribution (z-score > 1.96). Points in between the two vertical dashed lines indicate bins where both pre- and post-cue-onset time periods were used in the GLM. B: Sliding window LDA (bin size: 500 ms; step size: 100 ms) demonstrating the classification rate for cue identity (blue solid line), location (red solid line), and outcome (green solid line) using a pseudoensemble consisting of the 133 cue-modulated units. Dashed colored lines indicate the average of shuffling the firing rate order that went into the cross-validated LDA 100 times. Solid lines at the bottom indicate when the classifier performance greater than the shuffled distribution (z-score > 1.96). Points in between the two vertical dashed lines indicate bins where both pre- and post-cue-onset time periods were used in the classifier. C-D: Correlation matrices testing the presence and overlap of cue feature coding at cue-onset. C: Schematic outlining the possible outcomes for coding across cue features at cue-onset, generated by correlating the recoded beta coefficients from the GLMs and comparing to a shuffled distribution (see text for analysis details). Top left: coding is not present, therefore no comparison is possible. Top right: cue features are coded by separate populations of units. Displayed is a correlation matrix with each of the 9 blocks representing correlations for two cue features across the post-cue-onset time bins from the sliding window GLM, with green representing positive correlations (r > 0), pink representing negative correlations (r < 0), and white representing no correlation (r = 0). X- and y-axis have the same axis labels, therefore the diagonal represents the correlation of a cue feature against itself at that particular time point (r = 1). Here the large amount of pink in the off-diagonal elements suggests that coding of cue features occur separately from one another. Bottom left: Coding of cue features occurs in overlapping but independent populations of units, shown here by the abundance of white and relative lack of green and pink in the off-diagonal elements. Bottom right: Coding of cue features occurs in a joint (correlated) overlapping population, shown here by the large amount of green in the off-diagonal elements. D: Correlation matrix showing the correlation among cue identity, location, and outcome coding surrounding cue-onset. The window of GLMs used in each block is from cue-onset to the 500 ms window post-cue-onset, in 100 ms steps. Each individual value is for a sliding window GLM within that range, with the scale bar contextualizing step size. Color bar displays relationship between correlation value and color. Colored square borders around each block indicate the result of a comparison of the mean correlation of a block to a shuffled distribution, with pink indicating separate populations (z-score < -1.96), grey indicating overlapping but independent populations, and green indicating joint overlapping populations (z-score > 1.96).

To assess what information may be encoded at the population level, we trained a classifier on a pseudoensemble of the 133 cue-modulated units (Figure 4B). Specifically, we used the firing rate of each unit for each 129 trial as an observation, and different cue conditions as trial labels (e.g. light block, sound block). A lin-130 ear discriminant analysis (LDA) classifier with 10-fold cross-validation could correctly predict a trial above 131 chance levels for the identity and location of a cue across all time points surrounding cue-onset (z-score > 132 1.96 when comparing classification accuracy of data versus a shuffled distribution), whereas the ability to 133 predict whether a trial was reward-available or reward-unavailable (outcome coding) was not significantly 134 higher than the shuffled distribution for the time point containing 500 ms of pre-cue firing rate, and increased 135 gradually as a trial progressed, providing evidence that cue information is also present in the pseudoensemble 136 level. 137

To quantify the overlap of cue feature coding we correlated recoded beta coefficients from the GLMs, as-138 signing a value of '1' if a cue feature was a significant predictor for that unit and '0' if not, and calculated 139 a z-score comparing the mean of the obtained correlations to the mean and standard deviation of a shuffled 140 distribution, generated by shuffling the unit ordering within an array (Figure 1A,C, 4C,D). This revealed that 141 identity was coded independently from outcome (mean r = .01, z-score = 0.81), and by a joint population 142 with location (mean r = .10, z-score = 6.61), while location and outcome were coded by a joint population of 143 units (mean r = .12, z-score = 8.07). Together, these findings show that various cue features are represented 144 in the NAc at both the single-unit and pseudoensemble level, with location being coded by joint populations 145 with identity and outcome, but that identity is coded independently from outcome.

NAc population activity distinguishes all task phases:

Next, we sought to determine how coding of cue features evolved over time. Two main possibilities can be distinguished (Figure 1B); a unit coding for a feature such as cue identity could remain persistently active, or a progression of distinct units could activate in sequence. To visualize the distribution of responses throughout our task space and test if this distribution is modulated by cue features, we z-scored the firing rate of each unit, plotted the normalized firing rates of all units aligned to cue-onset, and sorted them according to the time of peak firing rate (Figure 5). We did this separately for both the light and sound blocks, and found a nearly uniform distribution of firing fields in task space that was not limited to alignment to the cue (Figure 5A). Furthermore, to determine if this population level activity was similar across blocks, we also organized firing during the sound blocks according to the ordering derived from the light blocks. This revealed that while there was some preservation of order, the overall firing was qualitatively different across the two blocks, implying that population activity distinguishes between light and sound blocks.

To control for the possibility that any comparison of trials would produce this effect, we divided each block 159 into two halves and looked at the correlation of the average smoothed firing rates across various combina-160 tions of these halves across our cue-onset centered epoch to see if the across block comparisons were less 161 correlated than the within block correlations. A linear mixed effects model revealed that within block corre-162 lations (e.g., one half of light trials vs other half of light trials) were higher and more similar than across block 163 correlations (e.g. half of light trials vs half of sound trials) suggesting that activity in the NAc discriminates 164 across light and sound blocks (mean within block correlation = .38; mean across block correlation = .34; p 165 < .001). This process was repeated for cue location (Figure 5B; mean within block correlation = .36; mean 166 across block correlation = .29; p < .001) and cue outcome (Figure 5C; mean within block correlation = .35; 167 mean across block correlation = .25; p < .001). Additionally, given that the majority of our units showed an 168 inhibitory response to the cue, we also plotted the firing rates according to the lowest time in firing, and again 169 found some maintenance of order, but largely different ordering across the two blocks (Figure 5-supplement 170 1). Together, these results illustrate that NAc coding of task space was not confined to salient events such as 171 cue-onset, but was approximately uniformly distributed throughout the task.

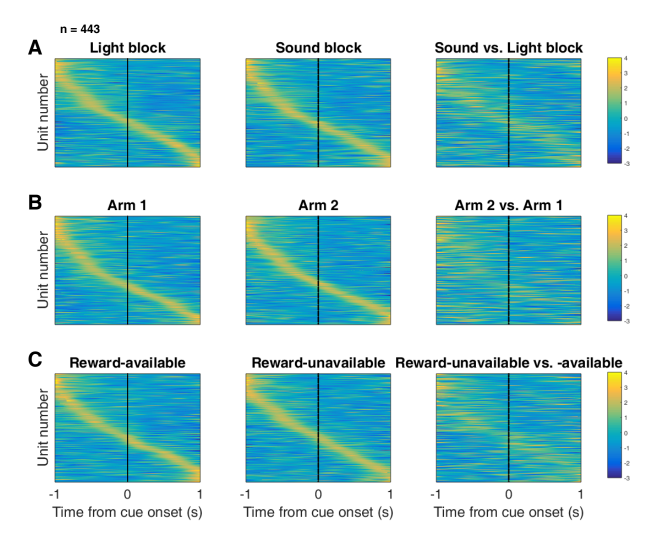


Figure 5: Distribution of NAc firing rates across time surrounding cue-onset. Each panel shows normalized (z-score) peak firing rates for all recorded NAc units (each row corresponds to one unit) as a function of time (time 0 indicates cue-onset), averaged across all trials for a specific cue type, indicated by text labels. A, left: Heat plot showing smoothed normalized firing activity of all recorded NAc units ordered according to the time of their peak firing rate during the light block. Each row is a units average activity across time to the light block. Dashed line indicates cue-onset. Notice the yellow band across time, indicating all aspects of visualized task space were captured by the peak firing rates of various units. A, middle: Same units ordered according to the time of the peak firing rate during the sound block. Note that for both blocks, units tile time approximately uniformly with a clear diagonal of elevated firing rates. A, right: Unit firing rates taken from the sound block, ordered according to peak firing rate taken from the light block. Note that a weaker but still discernible diagonal persists, indicating partial similarity between firing rates in the two blocks. Color bar displays relationship between z-score and color. B: Same layout as in A, except that the panels now compare two different locations on the track instead of two cue modalities. As for the different cue modalities, NAc units clearly discriminate between locations, but also maintain some similarity across locations, as evident from the visible diagonal in the right panel. Two example locations were used for display purposes; other location pairs showed a similar pattern. C: Same layout as in A, except that panels now compare reward-available and reward-unavailable trials. Overall, NAc units coded experience on the task, as opposed to being confined to specific task events only. Units from all sessions and animals were pooled for this analysis.

NAc encoding of cue features persists until outcome:

In order to be useful for credit assignment in reinforcement learning, a trace of the cue must be maintained until the outcome, so that information about the outcome can be associated with the outcome-predictive cue 175 (Figure 1B). Investigation into the post-approach period during nosepoke revealed units that discriminated 176 various cue features, with some units showing discriminative activity at both cue-onset and nosepoke (Figure 177 6, 6-supplement 1, 6-supplement 2). To quantitatively test whether representations of cue features persisted post-approach until the outcome was revealed, we fit sliding window GLMs to the post-approach firing 179 rates of cue-modulated units aligned to both the time of nosepoke into the reward receptacle, and after the 180 outcome was revealed (Figure 7A,B, 7-supplement 1 A-D, Table 1). This analysis showed that a variety 181 of units discriminated firing according to cue identity (~20% of cue-modulated units), location (~25% of 182 cue-modulated units), and outcome (~25% of cue-modulated units), but not other task parameters, showing 183 that NAc activity discriminates various cue conditions well into a trial. 184

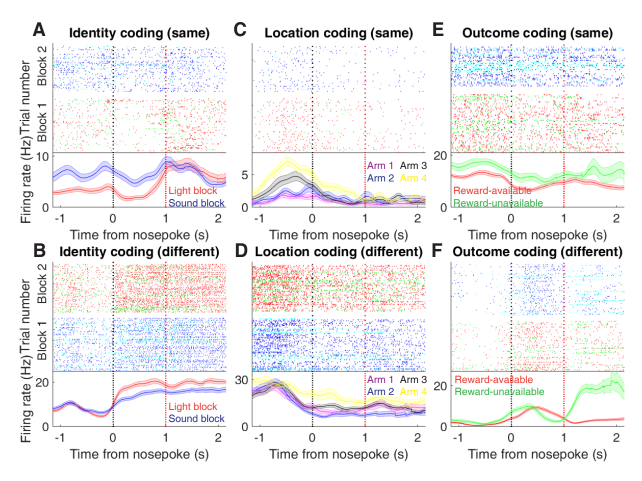


Figure 6: Examples of cue-modulated NAc units influenced by cue features at time of nosepoke. A: Example of a cue-modulated NAc unit that exhibited identity coding at both cue-onset and during subsequent nosepoke hold. Top: rasterplot showing the spiking activity across all trials aligned to nosepoke. Spikes across trials are color coded according to cue type (red: reward-available light; green: reward-unavailable light; navy blue: reward-available sound; light blue: reward-unavailable sound). White space halfway up the rasterplot indicates switching from one block to the next. Black dashed line indicates nosepoke. Red dashed line indicates receipt of outcome. Bottom: PETHs showing the average smoothed firing rate for the unit for trials during light (red) and sound (blue) blocks, aligned to nosepoke. Lightly shaded area indicates standard error of the mean. Note this unit showed a sustained increase in firing to sound cues during the trial. B: An example of a unit that was responsive to cue identity at time of nosepoke but not cue-onset. C-D: Cue-modulated units that exhibited location coding, at both cue-onset and nosepoke (C), and only nosepoke (D). Each color in the PETHs represents average firing response for a different cue location. E-F: Cue-modulated units that exhibited outcome coding, at both cue-onset and nosepoke (E), and only nosepoke (F), with the PETHs comparing reward-available (red) and reward-unavailable (green) trials.

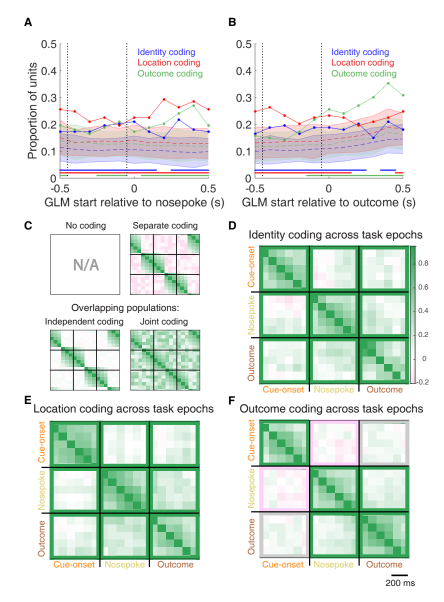


Figure 7: Summary of influence of cue features on cue-modulated NAc units at time points surrounding nosepoke and subsequent receipt of outcome. A-B: Sliding window GLM illustrating the proportion of cue-modulated units influenced by various predictors around time of nosepoke (A), and outcome (B). A: Sliding window GLM (bin size: 500 ms; step size: 100 ms) demonstrating the proportion of cue-modulated units where cue identity (blue solid line), location (red solid line), and outcome (green solid line) significantly contributed to the model at various time epochs relative to when the rat made a nosepoke. Dashed colored lines indicate the average of shuffling the firing rate order that went into the GLM 100 times. Error bars indicate 1.96 standard deviations from the shuffled mean. Solid lines at the bottom indicate when the proportion of units observed was greater than the shuffled distribution (z-score > 1.96). Points in between the two vertical dashed lines indicate bins where both pre- and post-cue-onset time periods were used in the GLM. B: Same as A, but for time epochs relative to receipt of outcome after the rat got feedback about his approach. C-F: Correlation matrices testing the persistence of cue feature coding across points in time. C: Schematic outlining the possible outcomes for coding of a cue feature across various points in a trial, generated by correlating the recoded beta coefficients from the GLMs and comparing to a shuffled distribution (see text for analysis details). Top left: coding is not present, therefore no comparison is possible. Top right: a cue feature is coded by separate populations of units across time. Displayed is a correlation matrix with each of the 9 blocks representing correlations for a cue feature across time bins for two task events from the sliding window GLM, with green representing positive correlations (r > 0), pink negative correlations (r < 0), and white representing significant correlation (r = 0). X- and y-axis have the same axis labels, therefore the diagonal represents the correlation of cue feature against itself at that particular time point (r = 1). Here the large amount of pink in the off-diagonal elements suggests that coding of a cue feature is accomplished by separate populations of units across time. Bottom left: Coding of a cue feature across time occurs in overlapping but independent populations of units, shown here by the abundance of white and relative lack of green and pink in the off-diagonal elements. Bottom right: Coding of a cue feature across time occurs in a joint overlapping population, shown here by the large amount of green in the off-diagonal elements. D: Correlation matrix showing the correlation of units that exhibited identity coding across time points after cue-onset, nosepoke, and outcome receipt. The window of GLMs used in each block is from the onset of the task phase to the \$00 ms window post-onset, in 100 ms steps. Each individual value is for a sliding window GLM within that range, with the scale bar contextualizing step size. Color bar displays relationship between correlation value and color. Colored square borders around each block indicate the result of a comparison of the mean correlation of a block to a shuffled distribution, with pink indicating separate populations (z-score < -1.96), grey indicating overlapping but independent populations, and green indicating joint overlapping populations (z-score > 1.96). E-F: Same as D, but for location and outcome coding, respectively.

To determine whether NAc representations of cue features at nosepoke and outcome were encoded by a similar pool of units as during cue-onset, we correlated recoded beta coefficients from the GLMs for a cue 186 feature across time points in the task, and compared the obtained correlations to correlations generated by 187 shuffling unit ordering within a recoded array (Figure 1B,C, 7C-F). This revealed that identity coding was 188 accomplished by a joint population across all three task events (cue-onset and nosepoke: mean r = .05, z-189 score = 3.47; cue-onset and outcome: mean r = .08, z-score = 5.55; nosepoke and outcome: mean r = .15, 190 z-score= 10.91). Applying this same analysis for cue location revealed a similar pattern for location coding 191 (cue-onset and nosepoke: mean r = .06, z-score = 4.15; cue-onset and outcome: mean r = .09, z-score 192 6.40; nosepoke and outcome: mean r = .20, z-score = 14.29). However, outcome coding at cue-onset was 193 separate from coding at nosepoke (mean r = -.04, z-score = -3.10), and independent from coding at outcome (mean r = .03, z-score = 1.65), while joint coding was observed between nosepoke and outcome (mean r =195 .15, z-score = 9.74). Together, these findings show that the NAc maintains representations of cue identity and location by a joint overlapping population throughout a trial, while separate populations of units encode 197 cue outcome before and after a behavioral decision has been made.

To assess overlap among cue features at nosepoke and outcome receipt, we applied the same recoded coefficient analysis (Figure 7-supplement 1 E,F). This revealed joint coding of cue features at the time of nosepoke (cue identity and location: mean r = .12, z-score = 8.26; cue identity and outcome: mean r = .05, z-score = 3.65; mean r = .10, z-score = 6.60); while at outcome, identity was coded by a joint population with both location (mean r = .09, z-score = 5.58), and outcome (mean r = .04, z-score = 2.93), and location and outcome were coded by an independent population of units (mean r = .00, z-score = 0.28).

To assess the distributed coding of units for task space around outcome receipt, we aligned normalized peak firing rates to nosepoke onset (Figure 7-supplement 2). This revealed a clustering of responses around outcome receipt for all cue conditions where the rat would have received reward, in addition to the same pattern of higher within- vs across-block correlations for cue identity (Figure 7-supplement 2 A,C; mean within block correlation = .55; mean across block correlation = .48; p < .001), cue location (Figure 7-

supplement 2 B,E; mean within block correlation = .47; mean across block correlation = .41; p < .001), and cue outcome (Figure 7-supplement 2 C,F; mean within block correlation = .51; mean across block correlation = .41; p < .001), further reinforcing that NAc activity distinguishes all task phases.

Discussion

The main result of the present study is that NAc units encode not only the expected outcome of outcome-214 predictive cues, but also the identity of such cues (Figure 1A). The population of units that coded for cue 215 identity was statistically independent from the population coding for expected outcome at cue-onset (i.e. 216 overlap as expected from chance), while a joint overlapping population coded for identity and outcome 217 at both nosepoke and outcome receipt (i.e. overlap greater than that expected from chance, Figure 1C). 218 Importantly, this identity coding was maintained on approach trials by a similar population of units both 219 during a delay period where the rat held a nosepoke until the outcome was received, and immediately after 220 outcome receipt (Figure 1B,C). Cue identity information was also present at the population level, as indicated by high classification performance based on pseudoensembles. More generally, NAc unit activity profiles 222 were not limited to salient task events such as the cue, nosepoke and outcome, but were distributed more uniformly throughout the task. This temporally distributed activity differed systematically between cue 224 identities, expected outcomes and locations. We discuss these observations and their implications below. 225

226 Identity coding:

Our finding that NAc units can discriminate between different outcome-predictive stimuli with similar motivational significance (i.e. encode cue identity) expands upon an extensive rodent literature examining NAc
correlates of conditioned stimuli (Ambroggi et al., 2008; Atallah et al., 2014; Bissonette et al., 2013; Cooch
et al., 2015; Day et al., 2006; Dejean et al., 2017; Goldstein et al., 2012; Ishikawa et al., 2008; Lansink et al.,

2012; McGinty et al., 2013; Nicola, 2004; Roesch et al., 2009; Roitman et al., 2005; Saddoris et al., 2011; 231 Setlow et al., 2003; Sugam et al., 2014; West & Carelli, 2016; Yun et al., 2004). Perhaps the most comparable 232 work in rodents comes from a study that found a subset of NAc units that modulated their firing for an odor 233 when it predicted distinct but equally valued rewards (Cooch et al., 2015). The present study is complemen-234 tary to such outcome identity coding, in showing that NAc units encode cue identity in addition to the reward 235 it predicts (Figure 1A). Setlow et al. (2003) paired two distinct odor cues with appetitive and aversive odor 236 cues respectively in a Go/NoGo task, such that cue identity and cue outcome were linked. Although reversal 237 sessions were run that uncoupled identity and outcome, the resulting changes in reinforcement history and 238 behavioral performance precluded a clear test of cue identity coding. Thus, our study is distinct in asking 239 how distinct cues encoding the same anticipated outcome are encoded. Our results suggest that the NAc dis-240 sociates cue identity representations at multiple levels of analysis (e.g. single-unit and population) even when 241 the motivational significance of these stimuli is identical. Viewed within the neuroeconomic framework of 242 decision making, functional magnetic resonance imaging (fMRI) studies have found support for NAc rep-243 resentations of offer value, a domain-general common currency signal that enables comparison of different attributes such as reward identity, effort, and temporal proximity (Bartra et al., 2013; Levy & Glimcher, 2012; 245 Peters & Büchel, 2009; Sescousse et al., 2015). Our study adds to a growing body of electrophysiological research that suggests the view of the NAc as a value center, while informative and capturing major aspects 247 of NAc processing, neglects additional contributions of NAc to learning and decision making such as the 248 offer (cue) identity signal reported here. 249

Our analyses were designed to eliminate several potential alternative interpretations to cue identity coding.

Because the different cues were separated into different blocks, units that discriminated between cue identities could instead be encoding time or other slowly-changing quantities. We excluded this possible confound by excluding units that showed a drift in firing between the first and second half within a block. Additionally, we included time as a nuisance variable in our GLMs, to exclude firing rate variance in the remaining units that could be attributed to this confound. Furthermore, we found that the temporally evolving firing rate throughout a trial was more strongly correlated within a block than across blocks. However, the possibility

remains that instead of, or in addition to, stimulus identity, these units encode a preferred context, or even a macroscale representation of progress through the session. Indeed, encoding of the current strategy could be an explanation for the presence of pre-cue identity coding (Figure 4A), as well as for the differential distributed coding of task structure across blocks observed in the current study (Figure 5).

An overall limitation of the current study is that rats were never presented with both sets of cues simul-261 taneously, and were not required to switch strategies between multiple sets of cues (this was attempted in 262 behavioral pilots, however animals took several days of training to successfully switch strategies). Addition-263 ally, our recordings were done during performance on the well-learned behavior, and not during the initial 264 acquisition of the cue-outcome relationships when an eligibility trace would be most useful. Thus, it is 265 unknown to what extent the cue identity encoding we observed is behaviorally relevant, although extrapo-266 lating data from other work (Sleezer et al., 2016) suggests that cue identity coding would be modulated by 267 relevance. Furthermore, NAc core lesions have been shown to impair shifting between different behavioral 268 strategies (Floresco et al., 2006), and it is possible that selectively silencing the units that prefer responding 269 for a given modality or rule would impair performance when the animal is required to use that information, 270 or artificial enhancement of those units would cause them to use the rule when it is the inappropriate strategy. 271

NAc activity provides a rich task representation beyond reward alone:

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Beyond coding of cue identity, we found several other notable features of NAc activity. First, a substantial number of cue-modulated units was differentially active depending on location, consistent with previous reports (Lavoie & Mizumori, 1994; Mulder et al., 2005; Strait et al., 2016; Wiener et al., 2003). However, it is notable that in our task, location is explicitly uninformative about reward, yet coding of this uninformative variable persists. This is unlike previous work of location coding in the dorsolateral striatum, which was present when location was predictive of reward, and absent when it was uninformative (Schmitzer-Torbert and Redish, 2008). Persistent coding of location in the NAc is likely attributable to inputs from the hippocampus (Lansink et al., 2016; Sjulson et al., 2017; Tabuchi et al., 2000; van der Meer & Redish, 2011);

speculatively, this coding may map onto a bias in credit assignment, such that motivationally relevant events are likely to be associated with the locations where they occur.

A second striking feature of NAc activity evident from this task is that NAc units were not only active at 283 salient events such as cue presentation, nosepoking, and feedback about the outcome, but distributed their 284 activity throughout a trial (Figure 5). This observation is consistent with previous work reporting that NAc 285 units can signal progress through a sequence of cues and/or actions (Atallah et al., 2014; Berke et al., 2009; 286 Khamassi et al., 2008; Lansink et al., 2012; Mulder et al., 2004; Shidara et al., 1998) and reminiscent of 287 similar observations in the ventral pallidum (Tingley et al., 2014) to which the NAc projects. Extending this 288 previous work, we show that the specific pattern of NAc units throughout a trial can be modified by task 289 variables such as cue identity. This richer view of NAc activity recalls a dynamical systems perspective, in 290 which different task conditions correspond to different trajectories in a neural state space (e.g. Buonomano 291 and Maass, 2009; Shenoy et al. 2013). In any case, this view of NAc activity provides a substantially richer 292 picture than that expected from encoding of reward-related variables alone. 293

Functional relevance of cue identity coding:

One possible function of cue identity coding is to support contextual modulation of the motivational rel-295 evance of specific cues. A context can be understood as a particular mapping between specific cues and 296 their outcomes: for instance, in context 1 cue A but not cue B is rewarded, whereas in context 2 cue B 297 but not cue A is rewarded. Successfully implementing such contextual mappings requires representation of 298 the cue identities. Indeed, Sleezer et al. (2016) recorded NAc responses during the Wisconsin Card Sort-299 ing Task, a common set-shifting task used in both the laboratory and clinic, and found units that preferred firing to stimuli when a certain rule, or rule category was currently active. Further support for a modula-301 tion of NAc responses by strategy comes from an fMRI study that examined blood-oxygen-level dependent (BOLD) levels during a set-shifting task (FitzGerald et al., 2014). In this task, participants learned two sets 303 of stimulus-outcome contingencies, a visual set and an auditory set. During testing they were presented with 304

both simultaneously, and the stimulus dimension that was relevant was periodically shifted between the two.

It was found that bilateral NAc activity reflected value representations for the currently relevant stimulus
dimension, and not the irrelevant stimulus. Given that BOLD activity is thought to reflect the processing
of incoming and local information, and not spiking output (Logothetis et al., 2001), it is possible that the
relevance-gated value representations observed by FitzGerald et al. (2014) are integrated with the relevant
identity coding in the output of the NAc, as observed in the current study.

A different potential role for cue identity coding is in learning to associate rewards with reward-predictive 311 features of the environment, a process referred to as *credit assignment* in the reinforcement learning literature 312 (Sutton & Barto, 1998). Maladaptive decision making, as occurs in schizophrenia, addiction, Parkinson's 313 Disease and others can result from dysfunctional reward prediction errors (RPEs) and value signals (Frank et 314 al., 2004; Gradin et al., 2011; Maia & Frank, 2011). This view has been successful in explaining both positive 315 and negative symptoms in schizophrenia, and deficits in learning from feedback in Parkinson's (Frank et al., 316 2004; Gradin et al., 2011). However, the effects of RPE and value updating are contingent upon encoding 317 of preceding action and cue features, the eligibility trace (Lee et al., 2012; Sutton & Barto, 1998). Value 318 updates can only be performed on these aspects of preceding experience that are encoded when the update 319 occurs. Therefore, maladaptive learning and decision making can result from not only aberrant RPEs but also 320 from altered cue feature encoding. For instance, on this task the environmental stimulus that signaled the 321 availability of reward was conveyed by two distinct cues that were presented in four locations. Although in 322 our current study, the location and identity of the cue did not require any adjustments in the animals behavior, 323 we found coding of these features alongside the expected outcome of the cue that could be the outcome of 324 credit assignment computations computed upstream (Akaishi et al., 2016; Asaad et al., 2017; Chau et al., 325 2015; Noonan et al., 2017). Identifying neural coding related to an aspect of credit assignment is important 326 as inappropriate credit assignment could be a contributor to conditioned fear overgeneralization seen in 327 disorders with pathological anxiety such as generalized anxiety disorder, post-traumatic stress disorder, and 328 obsessive-compulsive disorder (Kaczkurkin et al., 2017; Kaczkurkin & Lissek, 2013; Lissek et al., 2014), 329 and delusions observed in disorders such as schizophrenia, Alzheimer's and Parkinson's (Corlett et al., 2010; 330

Kapur, 2003). Thus, our results provide a starting point for studies of the neural basis of credit assignment, and the extent and specific manner in which this process fails in syndromes such as schizophrenia, obsessivecompulsive disorder, and others.

Methods

Subjects:

A sample size of 4 adult male Long-Evans rats (Charles River, Saint Constant, QC) from an a priori determined sample of 5 were used as subjects (1 rat was excluded from the data set due to poor cell yield).

Rats were individually housed with a 12/12-h light-dark cycle, and tested during the light cycle. Rats were
food deprived to 85-90% of their free feeding weight (weight at time of implantation was 440 - 470 g), and
water restricted 4-6 hours before testing. All experimental procedures were approved by the the University
of Waterloo Animal Care Committee (protocol# 11-06) and carried out in accordance with Canadian Council
for Animal Care (CCAC) guidelines.

Overall timeline:

Each rat was first handled for seven days during which they were exposed to the experiment room, the sucrose solution used as a reinforcer, and the click of the sucrose dispenser valves. Rats were then trained on the behavioral task (described in the next section) until they reached performance criterion. At this point they underwent hyperdrive implantation targeted at the NAc. Rats were allowed to recover for a minimum of five days before being retrained on the task, and recording began once performance returned to presurgery levels. Upon completion of recording, animals were gliosed, euthanized and recording sites were histologically confirmed.

Behavioral task and training:

The behavioral apparatus was an elevated, square-shaped track (100 x 100 cm, track width 10 cm) containing 352 four possible reward locations at the end of track "arms" (Figure 2A). Rats initiated a trial by triggering a 353 photobeam located 24 cm from the start of each arm. Upon trial initiation, one of two possible light cues (L1, 354 L2), or one of two possible sound cues (S1, S2), was presented that signaled the presence (reward-available 355 trial, L1+, S1+) or absence (reward-unavailable trial, L2-, S2-) of a 12% sucrose water reward (0.1 mL) at the upcoming reward site. A trial was classified as an approach trial if the rat turned left at the decision point 357 and made a nosepoke at the reward receptacle (40 cm from the decision point), while trials were classified as 358 a skip trial if the rat instead turned right at the decision point and triggered the photobeam to initiate the next 359 trial. A trial was labeled *correct* if the rat approached (i.e. nosepoked) on reward-available trials, and skipped 360 (i.e. did not nosepoke) on reward-unavailable trials. On reward-available trials there was a 1 second delay 361 between a nosepoke and subsequent reward delivery. Trial length was determined by measuring the length 362 of time from cue-onset until nosepoke (for approach trials), or from cue-onset until the start of the following 363 trial (for skip trials). Trials could only be initiated through clockwise progression through the series of arms, 364 and each entry into the subsequent arm on the track counted as a trial. Cues were present until 1 second after 365 outcome receipt on approach trials, and until initiating the following trial on skip trials. 366

Each session consisted of both a light block and a sound block with 100 trials each. Within a block, one cue signaled reward was available on that trial (L1+ or S1+), while the other signaled reward was not available 368 (L2- or S2-). Light block cues were a flashing white light, and a steady yellow light. Sound block cues 369 were a 2 kHz sine wave (low) and a 8 kHz sine wave (high) whose amplitude was modulated from 0 to 370 maximum by a 2 Hz sine wave. Outcome-cue associations were counterbalanced across rats, e.g. for some 371 rats L1+ was the flashing white light, and for others L1+ was the steady yellow light. The order of cue 372 presentation was pseudorandomized so that the same cue could not be presented more than twice in a row. 373 Block order within each day was also pseudorandomized, such that the rat could not begin a session with 374 the same block for more than two days in a row. Each session consisted of a 5 minute pre-session period

on a pedestal (a terracotta planter filled with towels), followed by the first block, then the second block, 376 then a 5 minute post-session period on the pedestal. For approximately the first week of training, rats were 377 restricted to running in the clockwise direction by presenting a physical barrier to running counterclockwise. 378 Cues signaling the availability and unavailability of reward, as described above, were present from the start 379 of training. Rats were trained for 200 trials per day (100 trials per block) until they discriminated between 380 the reward-available and reward-unavailable cues for both light and sound blocks for three consecutive days, 381 according to a chi-square test rejecting the null hypothesis of equal approaches for reward-available and 382 reward-unavailable trials, at which point they underwent electrode implant surgery. 383

384 Surgery:

Surgical procedures were as described previously (Malhotra et al., 2015). Briefly, animals were administered 385 analgesics and antibiotics, anesthetized with isoflurane, induced with 5% in medical grade oxygen and main-386 tained at 2% throughout the surgery (\sim 0.8 L/min). Rats were then chronically implanted with a "hyperdrive" 387 consisting of 20 independently drivable tetrodes, with 4 designated as references tetrodes, and the remaining 388 16 either all targeted for the right NAc (AP +1.4 mm and ML +1.6 mm relative to bregma; Paxinos & Watson 389 1998), or 12 in the right NAc and 4 targeted at the mPFC (AP +3.0 mm and ML +0.6 mm, relative to bregma; 390 only data from NAc tetrodes was analyzed). Following surgery, all animals were given at least five days to 391 recover while receiving post-operative care, and tetrodes were lowered to the target (DV -6.0 mm) before 392 being reintroduced to the behavioral task. 393

Data acquisition and preprocessing:

After recovery, rats were placed back on the task for recording. NAc signals were acquired at 20 kHz with a RHA2132 v0810 preamplifier (Intan) and a KJE-1001/KJD-1000 data acquisition system (Amplipex). Signals were referenced against a tetrode placed in the corpus callosum above the NAc. Candidate spikes for sorting into putative single units were obtained by band-pass filtering the data between 600-9000 Hz, thresh-

olding and aligning the peaks (UltraMegaSort2k, Hill et al., 2011). Spike waveforms were then clustered with KlustaKwik using energy and the first derivative of energy as features, and manually sorted into units (MClust 3.5, A.D. Redish et al., http://redishlab.neuroscience.umn.edu/MClust/MClust.html). Isolated units containing a minimum of 200 spikes within a session were included for subsequent analysis. Units were classified as FSIs by an absence of interspike intervals (ISIs) > 2 s, while MSNs had a combination of ISIs > 2 s and phasic activity with shorter ISIs (Atallah et al., 2014; Barnes et al., 2005).

405 Data analysis:

Behavior. To determine if rats distinguished behaviorally between the reward-available and reward-unavailable 406 cues (cue outcome), we generated linear mixed effects models to investigate the relationships between cue type and the proportion of trials approached, with cue outcome (reward available or not) and cue identity 408 (light or sound) as fixed effects, and the addition of an intercept for rat identity as a random effect. For each 409 cue, the average proportion of trials approached for a session was used as the response variable. Contribu-410 tion of cue outcome to behavior was determined by comparing the full model to a model with cue outcome 411 removed. To assess within session learning we divided each block into two halves, and compared a model 412 including a block half variable to a null model excluding this variable, to see if adding block half improved 413 prediction of overall behavioral performance. 414

Neural data. Given that some of our analyses compare firing rates across time, particularly comparisons across blocks, we sought to exclude units with unstable firing rates that would generate spurious results reflecting a drift in firing rate over time unrelated to our task. We used a multipronged strategy to address this potential confound. As a first step, we ran a Mann-Whitney U test comparing the cue-modulated firing rates for the first and second half of trials within a block, and excluded 99 of 443 units from analysis that showed a significant change for either block, leaving 344 units for further analyses by our GLM. Furthermore, we included time (trial number) as a nuisance variable in our GLMs to control for firing rate variance account for by this confound (see below). To investigate the contribution of different cue features (cue identity, cue

location and cue outcome) on the firing rates of NAc single units, we first determined whether firing rates for a unit were modulated by the onset of a cue by collapsing across all cues and comparing the firing rates for the 1 s preceding cue-onset with the 1 s following cue-onset. Single units were considered to be cue-modulated if a Wilcoxon signed-rank test comparing pre- and post-cue firing was significant at p < .01. Cue-modulated units were then classified as either increasing or decreasing if the post-cue activity was higher or lower than the pre-cue activity, respectively.

To determine the relative contribution of different task parameters to firing rate variance (as in Figure 4A, 429 4-supplement 1), a forward selection stepwise GLM using a Poisson distribution for the response variable 430 was fit to each cue-modulated unit, using data from every trial in a session. Cue identity (light block, sound 431 block), cue location (arm 1, arm 2, arm 3, arm 4), cue outcome (reward-available, reward-unavailable), 432 behavior (approach, skip), trial length, trial number, and trial history (reward availability on the previous 2 433 trials) were used as predictors, with firing rate as the response variable. The GLMs were fit using a 500 ms 434 sliding window moving in 100 ms steps centered at 250 ms pre-cue (so no post-cue activity was included) 435 to centered at 750 ms post-cue, such that 11 different GLMs were fit for each unit, tracking the temporal 436 dynamics of the influence of task parameters on firing rate around the onset of the cue. Units were classified 437 as being modulated by a given task parameter if addition of the parameter significantly improved model fit 438 using deviance as the criterion (p < .01), and the total proportion of cue-modulated units influenced by a 439 task parameter was counted for each time bin. A comparison of the R-squared value between the final model 440 and the final model minus the predictor of interest was used to determine the amount of firing rate variance 441 explained by the addition of that predictor for a given unit. To control for the amount of units that would 442 be affected by a predictor by chance, we shuffled the trial order of firing rates for a particular unit within a 443 time bin, ran the GLM with the shuffled firing rates, counted the proportion of units encoding a predictor, 444 and took the average of this value over 100 shuffles. We then calculated how many z-scores the observed 445 proportion was from the mean of the shuffled distribution. For this and all subsequent shuffle analyses, we 446 used a z-score of greater than 1.96 or less than -1.96 as a marker of significance. To further control for whether outcome coding could be attributed to subsequent behavioral variability at the choice point, we ran 448

our cue-onset GLM for approach trials only.

To get a sense of the predictive power of these cue feature representations we trained a classifier using firing 450 rates from a pseudoensemble comprised of our 133 cue-modulated units (Figure 4B). We created a matrix 451 of firing rates for each time epoch surrounding cue-onset where each row was an observation representing 452 the firing rate for a trial, and each column was a variable representing the firing rate for a given unit. Trial 453 labels, or classes, were each condition for a cue feature (e.g. light and sound for cue identity), making sure to align trial labels across units. We then ran LDA on these matrices, using 10-fold cross validation to train 455 the classifier on 90% of the trials and testing its predictions on the held out 10% of trials, and repeated this 456 approach to get the classification accuracy for 100 iterations. To test if the classification accuracy was greater 457 by chance, we shuffled the order of firing rates for each unit before we trained the classifier. We repeated 458 this for 100 shuffled matrices for each time point, and calculated how many z-scores the mean classification 459 rate of the observed data was from the mean of the shuffled distribution. 460

To determine the degree to which coding of cue identity, cue location, and cue outcome overlapped within 461 units we correlated the recoded beta coefficients from the GLMs for the cue features (Figure 4C,D). Specifi-462 cally, we generated an array for each cue feature at each point in time where for all cue-modulated units we 463 coded a '1' if the cue feature was a significant predictor in the final model, and '0' if it was not. We then 464 correlated an array of the coded 0s and 1s for one cue feature with a similar array for another cue feature, repeating this process for all post cue-onset sliding window combinations. The NAc was determined as coding 466 a pair of cue features in a) separate populations of units if there was a significant negative correlation (r < 1467 0), b) an independently coded overlapping population of units if there was no significant correlation (r = 0), 468 or c) a jointly coded overlapping population of units if there was a significant positive correlation (r > 0). 469 To summarize the correlation matrices generated from this analysis, we shuffled the unit ordering for each 470 array 100 times, took the mean of the 36 correlations for a block comparison for each of the 100 shuffles 471 for an analysis window, and used the mean and standard deviation of these shuffled correlation averages to 472 compare to the mean of the comparison block for the actual data.

To better visualize responses to cues and enable subsequent population level analyses (as in Figures 3, 5), 474 spike trains were convolved with a Gaussian kernel ($\sigma = 100 \text{ ms}$), and peri-event time histograms (PETHs) 475 were generated by taking the average of the convolved spike trains across all trials for a given task condition. 476 To visualize NAc representations of task space within cue conditions, normalized spike trains for all units 477 were ordered by the location of their maximum or minimum firing rate for a specified cue condition (Figure 478 5). To compare representations of task space across cue conditions for a cue feature, the ordering of units 479 derived for one condition (e.g. light block) was then applied to the normalized spike trains for the other 480 condition (e.g. sound block). To assess whether the task distributions were different across cue conditions, 481 we split each cue condition into two halves, controlling for the effects of time by shuffling trial ordering 482 before the split, and calculated the correlation of the temporally evolving smoothed firing rate across each of 483 these halves, giving us 6 correlation values for each unit. We then concatenated these 6 values across all 443 484 units to give us an array of 2658 correlation coefficients. We then fit a linear mixed effects model, trying to 485 predict these block comparison correlations with comparison type (e.g. 1st half of light block vs. 1st half of 486 sound sound) as a fixed-effect term, and unit number as a random-effect term. Comparison type is nominal, so dummy variables were created for the various levels of comparison type, and coefficients were generated 488 for each condition, referenced against one of the within-within comparison types (e.g. 1st half of light block 489 vs. 2nd half of light block). The NAc was considered to discriminate across cue conditions if across-block 490 correlations were lower than within-block correlations. Additionally, we ran a model comparison between 491 the above model and a null model with just unit number, to see if adding comparison type improved model 492 fit. 493

To identify the responsivity of units to different cue features at the time of nosepoke into a reward receptacle, and subsequent reward delivery, the same cue-modulated units from the cue-onset analyses were analyzed at the time of nosepoke and outcome receipt using identical analysis techniques for all approach trials (Figures 6, 7). To compare whether coding of a given cue feature was accomplished by the same or distinct population of units across time epochs, we ran the recoded coefficient correlation that was used to assess the degree of overlap among cue features within a time epoch. All analyses were completed in MATLAB R2015a, the code is available on our public GitHub repository (http://github.com/vandermeerlab/papers), and the data can be accessed through DataLad.

Histology:

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Upon completion of the experiment, recording channels were gliosed by passing $10 \mu A$ current for 10 seconds and waiting 5 days before euthanasia, except for rat R057 whose implant detached prematurely. Rats were anesthetized with 5% isoflurane, then asphyxiated with carbon dioxide. Transcardial perfusions were performed, and brains were fixed and removed. Brains were sliced in $50 \mu m$ coronal sections and stained with thionin. Slices were visualized under light microscopy, tetrode placement was determined, and electrodes with recording locations in the NAc were analyzed (Figure 8).

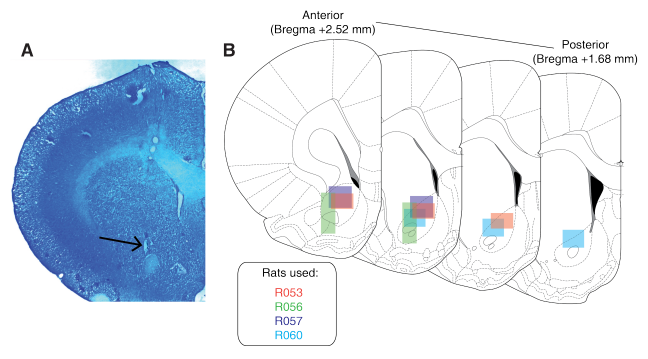


Figure 8: Histological verification of recording sites. Upon completion of experiments, brains were sectioned and tetrode placement was confirmed. **A:** Example section from R060 showing a recording site in the NAc core just dorsal to the anterior commissure (arrow). **B:** Schematic showing recording areas for all subjects.

Figure supplements

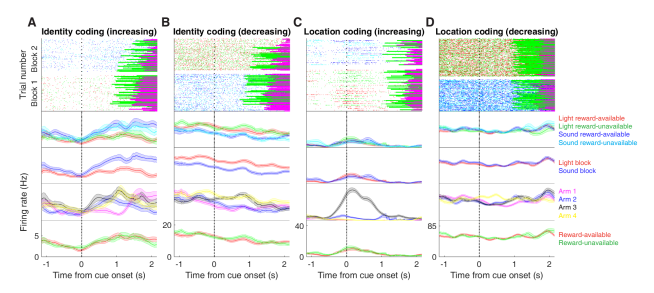


Figure 3 supplement 1: Expanded examples of cue-modulated NAc units influenced by different task parameters for Figure 3A-D, showing firing rate breakdown by: cue type (top PETH), cue identity (top-middle PETH), cue location (bottom-middle PETH), and cue outcome (bottom PETH).

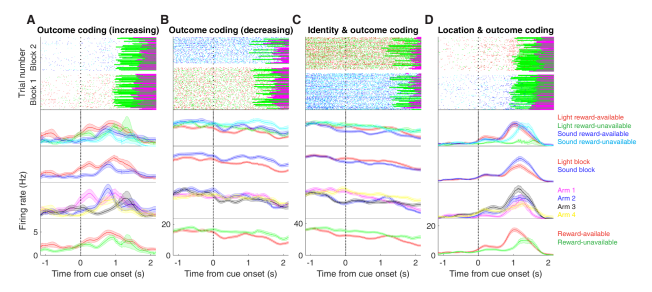


Figure 3 supplement 2: Expanded examples of cue-modulated NAc units influenced by different task parameters for Figure 3E-H, showing firing rate breakdown by: cue type (top PETH), cue identity (top-middle PETH), cue location (bottom-middle PETH), and cue outcome (bottom PETH).

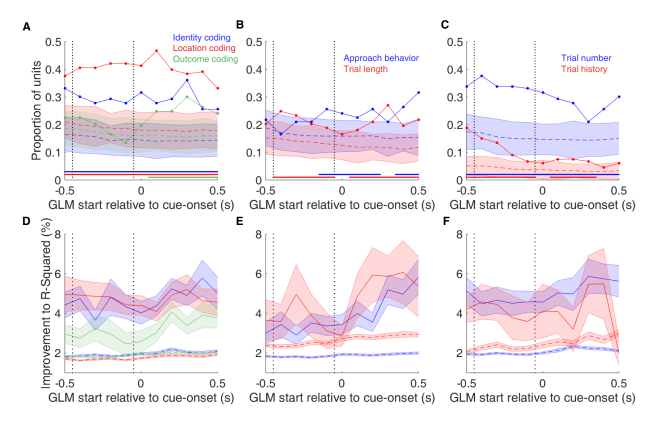


Figure 4 supplement 1: Summary of influence of various task parameters on cue-modulated NAc units at time points surrounding cue-onset. A-C: Sliding window GLM illustrating the proportion of cue-modulated units influenced by various predictors around time of cue-onset. A: Sliding window GLM (bin size: 500 ms; step size: 100 ms) demonstrating the proportion of cue-modulated units where cue identity (blue solid line), location (red solid line), and outcome (green solid line) significantly contributed to the model at various time epochs relative to cue-onset. Dashed colored lines indicate the average of shuffling the firing rate order that went into the GLM 100 times. Error bars indicate 1.96 standard deviations from the shuffled mean. Solid lines at the bottom indicate when the proportion of units observed was greater than the shuffled distribution (z-score > 1.96). Points in between the two vertical dashed lines indicate bins where both pre- and post-cue-onset time periods were used in the GLM. B: Same as A, but for approach behavior and trial length. C: Same as A, but for trial number and trial history. D-F: Average improvement to model fit. D: Average percent improvement to R-squared for units where cue identity, location, or outcome were significant contributors to the final model for time epochs surrounding cue-onset. Shaded area around mean represents the standard error of the mean. E: Same as D, but for approach behavior and trial length. F: Same D, but for trial number and trial history.

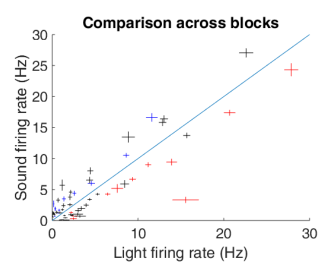


Figure 4 supplement 2: Scatter plot depicting comparison of firing rates for cue-modulated units across light and sound blocks. Crosses are centered on the mean firing rate, range represents the standard error of the mean. Colored crosses represents units that had cue identity as a significant predictor of firing rate variance in the GLM centered at cue-onset (blue are sound block preferring, red are light block preferring), whereas black crosses represent units where cue identity was not a significant predictor of firing rate variance. Diagonal dashed line indicates point of equal firing across blocks.

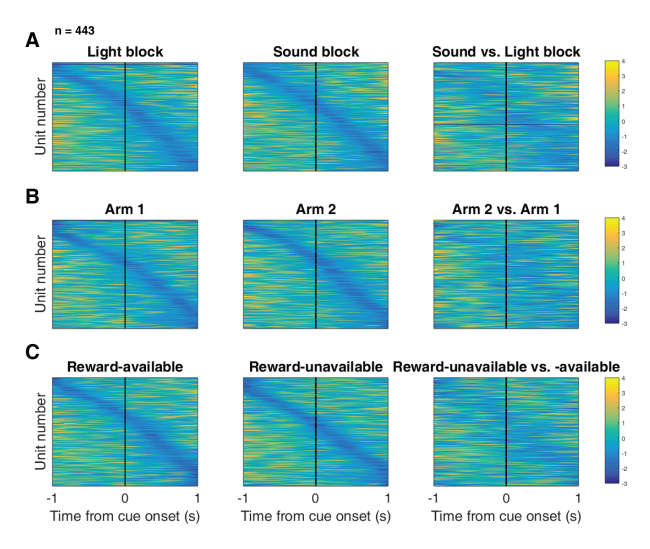


Figure 5 supplement 1: Distribution of NAc firing rates across time surrounding cue-onset. Each panel shows normalized (z-score) minimum firing rates for all recorded NAc units (each row corresponds to one unit) as a function of time (time 0 indicates cue-onset), averaged across all trials for a specific cue type, indicated by text labels. A: Responses during different stimulus blocks as in Figure 5A, but with units ordered according to the time of their minimum firing rate. B: Responses during trials on different arms as in Figure 5B, but with units ordered by their minimum firing rate. C: Responses during cues signaling different outcomes as in Figure 5C, but with units ordered by their minimum firing rate. Overall, NAc units coded experience on the task, as opposed to being confined to specific task events only. Units from all sessions and animals were pooled for this analysis.

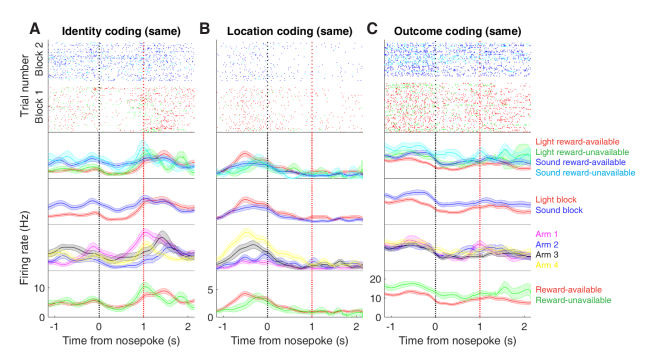


Figure 6 supplement 1: Expanded examples of cue-modulated NAc units influenced by different cue features at both cue-onset and during subsequent nosepoke hold for Figure 6A,C,E, showing firing rate breakdown by: cue type (top PETH), cue identity (top-middle PETH), cue location (bottom-middle PETH), and cue outcome (bottom PETH).

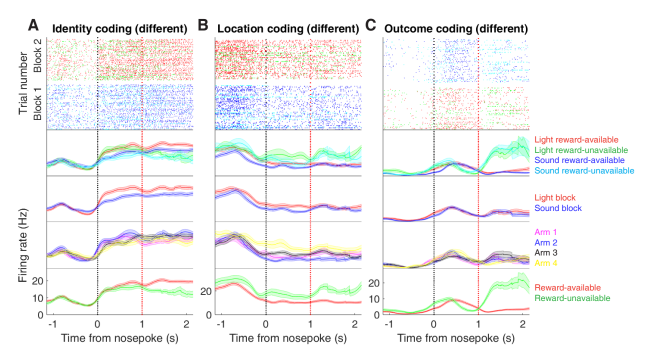


Figure 6 supplement 2: Expanded examples of cue-modulated NAc units influenced by different cue features at time of nosepoke for Figure 6B,D,F, showing firing rate breakdown by: cue type (top PETH), cue identity (top-middle PETH), cue location (bottom-middle PETH), and cue outcome (bottom PETH).

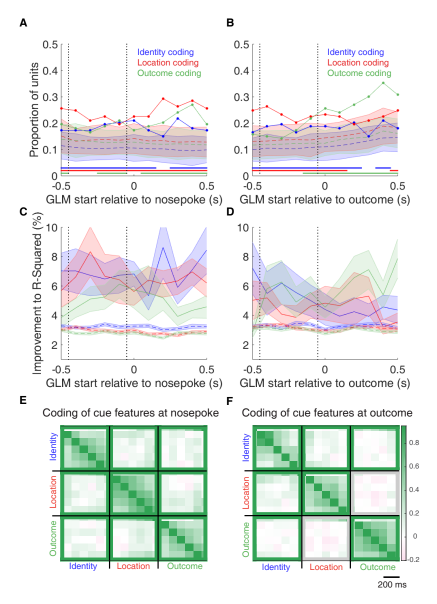


Figure 7 supplement 1: Summary of influence of cue features on cue-modulated NAc units at time points surrounding nosepoke and subsequent receipt of outcome. A-B: Sliding window GLM illustrating the proportion of cue-modulated units influenced by various predictors around time of nosepoke (A), and outcome (B). A: Sliding window GLM (bin size: 500 ms; step size: 100 ms) demonstrating the proportion of cue-modulated units where cue identity (blue solid line), location (red solid line), and outcome (green solid line) significantly contributed to the model at various time epochs relative to when the rat made a nosepoke. Dashed colored lines indicate the average of shuffling the firing rate order that went into the GLM 100 times. Error bars indicate 1.96 standard deviations from the shuffled mean. Solid lines at the bottom indicate when the proportion of units observed was greater than the shuffled distribution (z-score > 1.96). Points in between the two vertical dashed lines indicate bins where both pre- and post-cue-onset time periods were used in the GLM. B: Same as A, but for time epochs relative to receipt of outcome after the rat got feedback about his approach. C-D: Average improvement to model fit. C: Average percent improvement to R-squared for units where cue identity (blue solid line), location (red solid line), or outcome (green solid line) were significant contributors to the final model for time epochs relative to nosepoke. Dashed colored lines indicate the average of shuffling the firing rate order that went into the GLM 100 times. Shaded area around mean represents the standard error of the mean. D: Same C, but for time epochs relative to receipt of outcome. E-F: Correlation matrices testing the presence and overlap of cue feature coding at nosepoke (E) and outcome (F). E: Correlation matrix showing the correlation among identity, location, and outcome coding at nosepoke. Each of the 9 blocks represents correlations for two cue features across various nosepoke-centered time bins from the sliding window GLM, with green representing positive correlations (r > 0), pink negative correlations (r < 0), and grey representing no significant correlation (r = 0). X- and y-axis have the same axis labels, therefore the diagonal represents the correlation of a cue feature against itself at that particular time point (r = 1). The window of GLMs used in each block is from the onset of the task phase to the 500 ms window post-onset, in 100 ms steps. Each individual value is for a sliding window GLM within that range, with the scale bar contextualizing step size. Colored square borders around each block indicate the result of a comparison of the mean correlation to a shuffled distribution, with pink indicating separate populations (z-score < -1.96), grey indicating overlapping but independent populations, and greehlindicating joint overlapping populations (z-score > 1.96). F: Same as E, but for time bins following outcome receipt. Color bar displays relationship between correlation value and color.

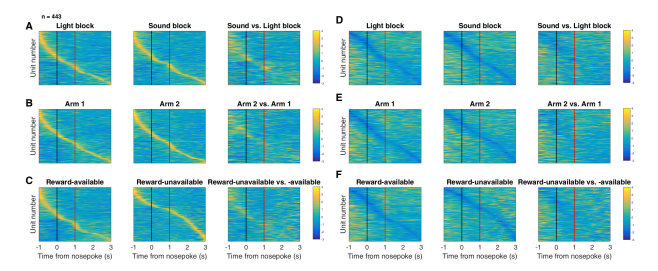


Figure 7 supplement 2: Distribution of NAc firing rates across time surrounding nosepoke for approach trials. Each panel shows normalized (z-score) firing rates for all recorded NAc units (each row corresponds to one unit) as a function of time (time 0 indicates nosepoke), averaged across all approach trials for a specific cue type, indicated by text labels. A-C: Heat plots aligned to normalized peak firing rates. A, far left: Heat plot showing smoothed normalized firing activity of all recorded NAc units ordered according to the time of their peak firing rate during the light block. Each row is a units average activity across time to the light block. Black dashed line indicates nosepoke. Red dashed line indicates reward delivery occurring 1 s after nosepoke for reward-available trials. Notice the yellow band across time, indicating all aspects of visualized task space were captured by the peak firing rates of various units. A, middle: Same units ordered according to the time of the peak firing rate during the sound block. Note that for both blocks, units tile time approximately uniformly with a clear diagonal of elevated firing rates, and a clustering around outcome receipt. A, right: Unit firing rates taken from the sound block, ordered according to peak firing rate taken from the light block. Note that a weaker but still discernible diagonal persists, indicating partial similarity between firing rates in the two blocks. Color bar displays relationship between z-score and color. B: Same layout as in A, except that the panels now compare two different locations on the track instead of two cue modalities. As for the different cue modalities, NAc units clearly discriminate between locations, but also maintain some similarity across locations, as evident from the visible diagonal in the right panel. Two example locations were used for display purposes; other location pairs showed a similar pattern. C: Same layout as in A, except that panels now compare correct reward-available and incorrect reward-unavailable trials. The disproportionate coding around outcome receipt for reward-available, but not reward-unavailable trials suggests encoding of reward receipt by NAc units. D-F: Heat plots aligned to normalized minimum firing rates. D: Responses during different stimulus blocks as in A, but with units ordered according to the time of their minimum firing rate. E: Responses during trials on different arms as in B, but with units ordered by their minimum firing rate. F: Responses during cues signaling different outcomes as in C, but with units ordered by their minimum firing rate. Overall, NAc units coded experience on the task, as opposed to being confined to specific task events only. Units from all sessions and animals were pooled for this analysis.

References

- 511 Akaishi, R., Kolling, N., Brown, J. W., & Rushworth, M. (2016). Neural Mechanisms of Credit Assignment in a Multicue
- 512 Environment. *Journal of Neuroscience*, 36(4), 1096–1112. doi: 10.1523/JNEUROSCI.3159-15.2016
- 513 Ambroggi, F., Ishikawa, A., Fields, H. L., & Nicola, S. M. (2008). Basolateral Amygdala Neurons Facilitate Reward-Seeking
- Behavior by Exciting Nucleus Accumbens Neurons. Neuron, 59(4), 648–661. doi: 10.1016/j.neuron.2008.07.004
- Asaad, W. F., Lauro, P. M., Perge, J. A., & Eskandar, E. N. (2017). Prefrontal Neurons Encode a Solution to the Credit-Assignment
- 516 Problem. The Journal of Neuroscience, 37(29), 6995–7007. doi: 10.1523/JNEUROSCI.3311-16.2017
- Atallah, H. E., McCool, A. D., Howe, M. W., & Graybiel, A. M. (2014). Neurons in the ventral striatum exhibit cell-type-specific
- representations of outcome during learning. *Neuron*, 82(5), 1145–1156. doi: 10.1016/j.neuron.2014.04.021
- Averbeck, B. B., & Costa, V. D. (2017). Motivational neural circuits underlying reinforcement learning. *Nature Neuroscience*,
- 520 20(4), 505–512. doi: 10.1038/nn.4506
- Barnes, T. D., Kubota, Y., Hu, D., Jin, D. Z., & Graybiel, A. M. (2005). Activity of striatal neurons reflects dynamic encoding and
- recoding of procedural memories. *Nature*, 437(7062), 1158–1161. doi: 10.1038/nature04053
- 523 Bartra, O., McGuire, J. T., & Kable, J. W. (2013). The valuation system: A coordinate-based meta-analysis
- of BOLD fMRI experiments examining neural correlates of subjective value. NeuroImage, 76, 412-427. doi:
- 525 10.1016/J.NEUROIMAGE.2013.02.063
- 526 Berke, J. D., Breck, J. T., & Eichenbaum, H. (2009). Striatal Versus Hippocampal Representations During Win-Stay Maze Perfor-
- 527 mance. Journal of Neurophysiology, 101(3), 1575–1587. doi: 10.1152/jn.91106.2008
- 528 Berridge, K. C. (2012). From prediction error to incentive salience: Mesolimbic computation of reward motivation. European
- 529 Journal of Neuroscience, 35(7), 1124–1143. doi: 10.1111/j.1460-9568.2012.07990.x
- Bissonette, G. B., Burton, A. C., Gentry, R. N., Goldstein, B. L., Hearn, T. N., Barnett, B. R., ... Roesch, M. R. (2013). Separate
- Populations of Neurons in Ventral Striatum Encode Value and Motivation. PLoS ONE, 8(5), e64673. doi: 10.1371/jour-
- nal.pone.0064673
- Bouton, M. E. (1993). Context, time, and memory retrieval in the interference paradigms of Pavlovian learning. Psychological
- *Bulletin*, 114(1), 80–99.doi: 10.1371/journal.pone.0064673
- 535 Carelli, R. M. (2010). Drug Addiction: Behavioral Neurophysiology. In Encyclopedia of neuroscience (pp. 677-682). Elsevier.
- doi: 10.1016/B978-008045046-9.01546-1
- 537 Chang, S. E., & Holland, P. C. (2013). Effects of nucleus accumbens core and shell lesions on autoshaped lever-pressing. Be-
- 538 havioural Brain Research, 256, 36–42. doi: 10.1016/j.bbr.2013.07.046
- 539 Chang, S. E., Wheeler, D. S., & Holland, P. C. (2012). Roles of nucleus accumbens and basolateral amygdala in autoshaped lever
- 540 pressing. Neurobiology of Learning and Memory, 97(4), 441–451. doi: 10.1016/j.nlm.2012.03.008

- Chau, B. K. H., Sallet, J., Papageorgiou, G. K., Noonan, M. A. P., Bell, A. H., Walton, M. E., & Rushworth, M. F. S. (2015).
- 542 Contrasting Roles for Orbitofrontal Cortex and Amygdala in Credit Assignment and Learning in Macaques. *Neuron*, 87(5),
- 543 1106–1118. doi: 10.1016/j.neuron.2015.08.018
- 544 Cheer, J. F., Aragona, B. J., Heien, M. L. A. V., Seipel, A. T., Carelli, R. M., & Wightman, R. M. (2007). Coordi-
- nated Accumbal Dopamine Release and Neural Activity Drive Goal-Directed Behavior. Neuron, 54(2), 237-244. doi:
- 546 10.1016/j.neuron.2007.03.021
- Cohen, J. D., & Servan-Schreiber, D. (1992). Context, cortex, and dopamine: a connectionist approach to behavior and biology in
- schizophrenia. Psychological Review, 99(1), 45.doi: 10.1016/j.neuron.2007.03.021
- Cooch, N. K., Stalnaker, T. A., Wied, H. M., Bali-Chaudhary, S., McDannald, M. A., Liu, T. L., & Schoenbaum, G. (2015).
- Orbitofrontal lesions eliminate signalling of biological significance in cue-responsive ventral striatal neurons. *Nature Com-*
- *munications*, 6, 7195. doi: 10.1038/ncomms8195
- 552 Corbit, L. H., & Balleine, B. W. (2011). The General and Outcome-Specific Forms of Pavlovian-Instrumental Transfer Are
- 553 Differentially Mediated by the Nucleus Accumbens Core and Shell. *Journal of Neuroscience*, 31(33), 11786–11794. doi:
- 554 10.1523/JNEUROSCI.2711-11.2011
- 555 Corlett, P. R., Taylor, J. R., Wang, X.-J., Fletcher, P. C., & Krystal, J. H. (2010). Toward a neurobiology of delusions. Progress in
- *Neurobiology*, 92, 345–369. doi: 10.1016/j.pneurobio.2010.06.007
- 557 Cromwell, H. C., & Schultz, W. (2003). Effects of Expectations for Different Reward Magnitudes on Neuronal Activity in Primate
- 558 Striatum. *Journal of Neurophysiology*, 89(5), 2823–2838. doi: 10.1152/jn.01014.2002
- 559 Day, J. J., Roitman, M. F., Wightman, R. M., & Carelli, R. M. (2007). Associative learning mediates dynamic shifts in dopamine
- signaling in the nucleus accumbens. Nature Neuroscience, 10(8), 1020–1028. doi: 10.1038/nn1923
- Day, J. J., Wheeler, R. A., Roitman, M. F., & Carelli, R. M. (2006). Nucleus accumbens neurons encode Pavlovian approach behav-
- iors: Evidence from an autoshaping paradigm. European Journal of Neuroscience, 23(5), 1341–1351. doi: 10.1111/j.1460-
- 563 9568.2006.04654.x
- Dejean, C., Sitko, M., Girardeau, P., Bennabi, A., Caillé, S., Cador, M., . . . Le Moine, C. (2017). Memories of Opiate Withdrawal
- Emotional States Correlate with Specific Gamma Oscillations in the Nucleus Accumbens. *Neuropsychopharmacology*,
- 566 42(5), 1157–1168. doi: 10.1038/npp.2016.272
- du Hoffmann, J., & Nicola, S. M. (2014). Dopamine Invigorates Reward Seeking by Promoting Cue-Evoked Excitation in the
- Nucleus Accumbens. *Journal of Neuroscience*, 34(43), 14349–14364. doi: 10.1523/JNEUROSCI.3492-14.2014
- Estes, W. K. (1943). Discriminative conditioning. I. A discriminative property of conditioned anticipation. Journal of Experimental
- 570 Psychology, 32(2), 150–155. doi: 10.1037/h0058316
- FitzGerald, T. H. B., Schwartenbeck, P., & Dolan, R. J. (2014). Reward-Related Activity in Ventral Striatum Is Action Contingent
- and Modulated by Behavioral Relevance. Journal of Neuroscience, 34(4), 1271–1279. doi: 10.1523/JNEUROSCI.4389-
- 573 13.2014

- Flagel, S. B., Clark, J. J., Robinson, T. E., Mayo, L., Czuj, A., Willuhn, I., ... Akil, H. (2011). A selective role for dopamine in stimulus-reward learning. *Nature*, 469(7328), 53–59. doi: 10.1038/nature09588
- Floresco, S. B. (2015). The Nucleus Accumbens: An Interface Between Cognition, Emotion, and Action. *Annual Review of Psychology*, 66(1), 25–52. doi: 10.1146/annurev-psych-010213-115159
- Floresco, S. B., Ghods-Sharifi, S., Vexelman, C., & Magyar, O. (2006). Dissociable roles for the nucleus accumbens core and shell in regulating set shifting. *Journal of Neuroscience*, 26(9), 2449–2457. doi: 10.1523/JNEUROSCI.4431-05.2006
- Frank, M. J., Seeberger, L. C., & O'Reilly, R. C. (2004). By carrot or by stick: Cognitive reinforcement learning in Parkinsonism.

 Science, 306(5703), 1940–1943. doi: 10.1126/science.1102941
- Giertler, C., Bohn, I., & Hauber, W. (2004). Transient inactivation of the rat nucleus accumbens does not impair guidance of instrumental behaviour by stimuli predicting reward magnitude. *Behavioural Pharmacology*, *15*(1), 55–63.doi: 10.1126/sci-ence.1102941
- Goldstein, B. L., Barnett, B. R., Vasquez, G., Tobia, S. C., Kashtelyan, V., Burton, A. C., ... Roesch, M. R. (2012). Ventral Striatum
 Encodes Past and Predicted Value Independent of Motor Contingencies. *Journal of Neuroscience*, 32(6), 2027–2036. doi:
- 587 10.1523/JNEUROSCI.5349-11.2012
- Goto, Y., & Grace, A. A. (2008). Limbic and cortical information processing in the nucleus accumbens. *Trends in Neurosciences*,
 31(11), 552–558. doi: 10.1016/j.tins.2008.08.002
- Gradin, V. B., Kumar, P., Waiter, G., Ahearn, T., Stickle, C., Milders, M., ... Steele, J. D. (2011). Expected value and prediction
 error abnormalities in depression and schizophrenia. *Brain*, *134*(6), 1751–1764. doi: 10.1093/brain/awr059
- Grant, D. A., & Berg, E. (1948). A behavioral analysis of degree of reinforcement and ease of shifting to new responses in a
 Weigl-type card-sorting problem. *Journal of Experimental Psychology*, 38(4), 404–411. doi: 10.1037/h0059831
- Hamid, A. A., Pettibone, J. R., Mabrouk, O. S., Hetrick, V. L., Schmidt, R., Vander Weele, C. M., ... Berke, J. D. (2015).
 Mesolimbic dopamine signals the value of work. *Nature Neuroscience*, 19(1), 117–126. doi: 10.1038/nn.4173
- Hart, A. S., Rutledge, R. B., Glimcher, P. W., & Phillips, P. E. M. (2014). Phasic Dopamine Release in the Rat Nucleus Accumbens Symmetrically Encodes a Reward Prediction Error Term. *The Journal of Neuroscience*, 34(3), 698–704. doi: 10.1523/JNEUROSCI.2489-13.2014
- Hassani, O. K., Cromwell, H. C., & Schultz, W. (2001). Influence of Expectation of Different Rewards on Behavior-Related
 Neuronal Activity in the Striatum. *Journal of Neurophysiology*, 85(6), 2477–2489. doi: 10.1152/jn.2001.85.6.2477
- Hearst, E., & Jenkins, H. M. (1974). Sign-tracking: the stimulus-reinforcer relation and directed action. Psychonomic Society.doi: 10.1152/jn.2001.85.6.2477
- Hill, D. N., Mehta, S. B., & Kleinfeld, D. (2011). Quality Metrics to Accompany Spike Sorting of Extracellular Signals. *Journal of Neuroscience*, 31(24), 8699–8705. doi: 10.1523/JNEUROSCI.0971-11.2011
- Holland, P. C. (1992). Occasion setting in pavlovian conditioning. *Psychology of Learning and Motivation*, 28(C), 69–125. doi: 10.1016/S0079-7421(08)60488-0

- Hollerman, J. R., Tremblay, L., & Schultz, W. (1998). Influence of Reward Expectation on Behavior-Related Neuronal Activity in
 Primate Striatum. *Journal of Neurophysiology*, 80(2), 947–963. doi: 10.1152/jn.1998.80.2.947
- Honey, R. C., Iordanova, M. D., & Good, M. (2014). Associative structures in animal learning: Dissociating elemental and configural processes. *Neurobiology of Learning and Memory*, *108*, 96–103. doi: 10.1016/j.nlm.2013.06.002
- Hyman, S. E., Malenka, R. C., & Nestler, E. J. (2006). NEURAL MECHANISMS OF ADDICTION: The Role of Reward-Related
 Learning and Memory. *Annual Review of Neuroscience*, 29(1), 565–598. doi: 10.1146/annurev.neuro.29.051605.113009
- Ikemoto, S. (2007). Dopamine reward circuitry: Two projection systems from the ventral midbrain to the nucleus accumbensolfactory tubercle complex. *Brain Research Reviews*, 56(1), 27–78. doi: 10.1016/j.brainresrev.2007.05.004
- Ishikawa, A., Ambroggi, F., Nicola, S. M., & Fields, H. L. (2008). Dorsomedial Prefrontal Cortex Contribution to Behavioral and Nucleus Accumbens Neuronal Responses to Incentive Cues. *Journal of Neuroscience*, 28(19), 5088–5098. doi:

10.1523/JNEUROSCI.0253-08.2008

617

- Joel, D., Niv, Y., & Ruppin, E. (2002). Actor-critic models of the basal ganglia: new anatomical and computational perspectives.

 Neural Networks, 15(4-6), 535–547. doi: 10.1016/S0893-6080(02)00047-3
- Kaczkurkin, A. N., Burton, P. C., Chazin, S. M., Manbeck, A. B., Espensen-Sturges, T., Cooper, S. E., ... Lissek, S. (2017).
 Neural substrates of overgeneralized conditioned fear in PTSD. American Journal of Psychiatry, 174(2), 125–134. doi:
 10.1176/appi.ajp.2016.15121549
- Kaczkurkin, A. N., & Lissek, S. (2013). Generalization of Conditioned Fear and Obsessive-Compulsive Traits. *Journal of Psychology & Psychotherapy*, 7, 3. doi: 10.4172/2161-0487.S7-003
- Kalivas, P. W., & Volkow, N. D. (2005). The neural basis of addiction: A pathology of motivation and choice. *American Journal of Psychiatry*, 162(8), 1403–1413. doi: 10.1176/appi.ajp.162.8.1403
- Kapur, S. (2003). Psychosis as a state of aberrant salience: A framework linking biology, phenomenology, and pharmacology in schizophrenia. *American Journal of Psychiatry*, 160(1), 13–23. doi: 10.1176/appi.ajp.160.1.13
- Khamassi, M., & Humphries, M. D. (2012). Integrating cortico-limbic-basal ganglia architectures for learning model-based and model-free navigation strategies. *Frontiers in Behavioral Neuroscience*, *6*, 79. doi: 10.3389/fnbeh.2012.00079
- Khamassi, M., Mulder, A. B., Tabuchi, E., Douchamps, V., & Wiener, S. I. (2008). Anticipatory reward signals in ventral striatal neurons of behaving rats. *European Journal of Neuroscience*, 28(9), 1849–1866. doi: 10.1111/j.1460-9568.2008.06480.x
- Lansink, C. S., Jackson, J. C., Lankelma, J. V., Ito, R., Robbins, T. W., Everitt, B. J., & Pennartz, C. M. A. (2012). Reward Cues in Space: Commonalities and Differences in Neural Coding by Hippocampal and Ventral Striatal Ensembles. *Journal of Neuroscience*, 32(36), 12444–12459. doi: 10.1523/JNEUROSCI.0593-12.2012
- Lansink, C. S., Meijer, G. T., Lankelma, J. V., Vinck, M. A., Jackson, J. C., & Pennartz, C. M. A. (2016). Reward Expectancy

 Strengthens CA1 Theta and Beta Band Synchronization and Hippocampal-Ventral Striatal Coupling. *Journal of Neuro-*science, 36(41), 10598–10610. doi: 10.1523/JNEUROSCI.0682-16.2016
- 639 Lavoie, A. M., & Mizumori, S. J. (1994). Spatial, movement- and reward-sensitive discharge by medial ventral striatum neurons of

- rats. Brain Research, 638(1-2), 157–168. doi: 10.1016/0006-8993(94)90645-9
- Lee, D., Seo, H., & Jung, M. W. (2012). Neural Basis of Reinforcement Learning and Decision Making. *Annual Review of Neuroscience*, 35(1), 287–308. doi: 10.1146/annurev-neuro-062111-150512
- Levy, D. J., & Glimcher, P. W. (2012). The root of all value: a neural common currency for choice. *Current Opinion in Neurobiology*, 22, 1027–1038, doi: 10.1016/j.conb.2012.06.001
- Lissek, S., Kaczkurkin, A. N., Rabin, S., Geraci, M., Pine, D. S., & Grillon, C. (2014). Generalized anxiety disor-
- der is associated with overgeneralization of classically conditioned fear. Biological Psychiatry, 75(11), 909–915. doi:
- 647 10.1016/j.biopsych.2013.07.025
- 648 Logothetis, N. K., Pauls, J., Augath, M., Trinath, T., & Oeltermann, A. (2001). Neurophysiological investigation of the basis of the 649 fMRI signal. *Nature*, 412(6843), 150–157. doi: 10.1038/35084005
- Maia, T. V. (2009). Reinforcement learning, conditioning, and the brain: Successes and challenges. *Cognitive, Affective and Behavioral Neuroscience*, *9*(4), 343–364. doi: 10.3758/CABN.9.4.343
- Maia, T. V., & Frank, M. J. (2011). From reinforcement learning models to psychiatric and neurological disorders. *Nature* Neuroscience, 14(2), 154–162. doi: 10.1038/nn.2723
- Malhotra, S., Cross, R. W., Zhang, A., & Van Der Meer, M. A. A. (2015). Ventral striatal gamma oscillations are highly variable
- from trial to trial, and are dominated by behavioural state, and only weakly influenced by outcome value. *European Journal*
- of Neuroscience, 42(10), 2818–2832.doi: 10.1038/nn.2723
- 657 McDannald, M. A., Lucantonio, F., Burke, K. A., Niv, Y., & Schoenbaum, G. (2011). Ventral Striatum and Orbitofrontal Cortex
- Are Both Required for Model-Based, But Not Model-Free, Reinforcement Learning. Journal of Neuroscience, 31(7), 2700–
- 659 2705. doi: 10.1523/JNEUROSCI.5499-10.2011
- McGinty, V. B., Lardeux, S., Taha, S. A., Kim, J. J., & Nicola, S. M. (2013). Invigoration of reward seeking by cue and proximity encoding in the nucleus accumbens. *Neuron*, 78(5), 910–922. doi: 10.1016/j.neuron.2013.04.010
- Mulder, A. B., Shibata, R., Trullier, O., & Wiener, S. I. (2005). Spatially selective reward site responses in tonically active neurons of the nucleus accumbens in behaving rats. *Experimental Brain Research*, *163*(1), 32–43. doi: 10.1007/s00221-004-2135-3
- Mulder, A. B., Tabuchi, E., & Wiener, S. I. (2004). Neurons in hippocampal afferent zones of rat striatum parse routes into
- multi-pace segments during maze navigation. European Journal of Neuroscience, 19(7), 1923–1932. doi: 10.1111/j.1460-
- 9568.2004.03301.x
- Nicola, S. M. (2004). Cue-Evoked Firing of Nucleus Accumbens Neurons Encodes Motivational Significance During a Discriminative Stimulus Task. *Journal of Neurophysiology*, *91*(4), 1840–1865. doi: 10.1152/jn.00657.2003
- Nicola, S. M. (2010). The Flexible Approach Hypothesis: Unification of Effort and Cue-Responding Hypotheses for the Role of
- Nucleus Accumbens Dopamine in the Activation of Reward-Seeking Behavior. Journal of Neuroscience, 30(49), 16585–
- 671 16600. doi: 10.1523/JNEUROSCI.3958-10.2010
- 672 Niv, Y., Daw, N. D., Joel, D., & Dayan, P. (2007). Tonic dopamine: Opportunity costs and the control of response vigor. Psy-

- 673 chopharmacology, 191(3), 507–520. doi: 10.1007/s00213-006-0502-4
- Noonan, M. P., Chau, B. K., Rushworth, M. F., & Fellows, L. K. (2017). Contrasting Effects of Medial and Lateral Orbitofrontal
- 675 Cortex Lesions on Credit Assignment and Decision-Making in Humans. *The Journal of Neuroscience*, 37(29), 7023–7035.
- doi: 10.1523/JNEUROSCI.0692-17.2017
- Paxinos, G., & Watson, C. (1998). The Rat Brain in Stereotaxic Coordinates (4th ed.). San Diego: Academic Press.doi:
- 678 10.1523/JNEUROSCI.0692-17.2017
- Peters, J., & Büchel, C. (2009). Overlapping and distinct neural systems code for subjective value during intertemporal and risky
- decision making. The Journal of neuroscience: the official journal of the Society for Neuroscience, 29(50), 15727–34. doi:
- 681 10.1523/JNEUROSCI.3489-09.2009
- Rescorla, R. A., & Solomon, R. L. (1967). Two-Process Learning Theory: Relationships Between Pavlovian Conditioning and
- Instrumental Learning. *Psychological Review*, 74(3), 151–182. doi: 10.1037/h0024475
- Robinson, T. E., & Flagel, S. B. (2009). Dissociating the Predictive and Incentive Motivational Properties of Reward-Related Cues
- Through the Study of Individual Differences. Biological Psychiatry, 65(10), 869–873. doi: 10.1016/j.biopsych.2008.09.006
- Roesch, M. R., Singh, T., Brown, P. L., Mullins, S. E., & Schoenbaum, G. (2009). Ventral Striatal Neurons Encode the Value
- of the Chosen Action in Rats Deciding between Differently Delayed or Sized Rewards. Journal of Neuroscience, 29(42),
- 688 13365–13376. doi: 10.1523/JNEUROSCI.2572-09.2009
- Roitman, M. F., Wheeler, R. A., & Carelli, R. M. (2005). Nucleus accumbens neurons are innately tuned for reward-
- ing and aversive taste stimuli, encode their predictors, and are linked to motor output. Neuron, 45(4), 587–597. doi:
- 691 10.1016/j.neuron.2004.12.055
- 692 Saddoris, M. P., Stamatakis, A., & Carelli, R. M. (2011). Neural correlates of Pavlovian-to-instrumental transfer in the nucleus ac-
- cumbens shell are selectively potentiated following cocaine self-administration. European Journal of Neuroscience, 33(12),
- 694 2274–2287. doi: 10.1111/j.1460-9568.2011.07683.x
- 695 Salamone, J. D., & Correa, M. (2012). The Mysterious Motivational Functions of Mesolimbic Dopamine. Neuron, 76(3), 470–485.
- doi: 10.1016/j.neuron.2012.10.021
- 697 Schultz, W. (2016). Dopamine reward prediction error coding. Dialogues in Clinical Neuroscience, 18(1), 23-32. doi:
- 698 10.1038/nrn.2015.26
- 699 Schultz, W., Apicella, P., Scarnati, E., & Ljungberg, T. (1992). Neuronal activity in monkey ventral striatum related to the
- expectation of reward. The Journal of neuroscience: the official journal of the Society for Neuroscience, 12(12), 4595–
- 701 610. doi: 10.1523/JNEUROSCI.12-12-04595.1992
- Schultz, W., Dayan, P., & Montague, P. R. (1997). A neural substrate of prediction and reward. Science, 275(5306), 1593–1599.
- 703 doi: 10.1126/science.275.5306.1593
- Sescousse, G., Li, Y., & Dreher, J.-C. (2015). A common currency for the computation of motivational values in the human striatum.
- 705 Social Cognitive and Affective Neuroscience, 10(4), 467–473. doi: 10.1093/scan/nsu074

- Setlow, B., Schoenbaum, G., & Gallagher, M. (2003). Neural encoding in ventral striatum during olfactory discrimination learning.
- 707 Neuron, 38(4), 625–636. doi: 10.1016/S0896-6273(03)00264-2
- Shidara, M., Aigner, T. G., & Richmond, B. J. (1998). Neuronal signals in the monkey ventral striatum related to progress through
- 709 a predictable series of trials. *Journal of Neuroscience*, 18(7), 2613–25.doi: 10.1016/S0896-6273(03)00264-2
- 710 Sjulson, L., Peyrache, A., Cumpelik, A., Cassataro, D., & Buzsáki, G. (2017). Cocaine place conditioning strengthens location-
- specific hippocampal inputs to the nucleus accumbens. bioRxiv, 1–10. doi: 10.1101/105890
- 712 Sleezer, B. J., Castagno, M. D., & Hayden, B. Y. (2016). Rule Encoding in Orbitofrontal Cortex and Striatum Guides Selection.
- 713 Journal of Neuroscience, 36(44), 11223–11237. doi: 10.1523/JNEUROSCI.1766-16.2016
- 714 Strait, C. E., Sleezer, B. J., Blanchard, T. C., Azab, H., Castagno, M. D., & Hayden, B. Y. (2016). Neuronal selectivity
- for spatial positions of offers and choices in five reward regions. *Journal of Neurophysiology*, 115(3), 1098–1111. doi:
- 716 10.1152/jn.00325.2015
- 717 Sugam, J. A., Saddoris, M. P., & Carelli, R. M. (2014). Nucleus accumbens neurons track behavioral preferences and reward
- outcomes during risky decision making. Biological Psychiatry, 75(10), 807–816. doi: 10.1016/j.biopsych.2013.09.010
- 719 Sutton, R., & Barto, A. (1998). Reinforcement Learning: An Introduction (Vol. 9) (No. 5). MIT Press, Cambridge, MA. doi:
- 720 10.1109/TNN.1998.712192
- Tabuchi, E. T., Mulder, A. B., & Wiener, S. I. (2000). Position and behavioral modulation of synchronization of hippocam-
- pal and accumbens neuronal discharges in freely moving rats. Hippocampus, 10(6), 717-728. doi: 10.1002/1098-
- 723 1063(2000)10:6;717::AID-HIPO1009;3.0.CO;2-3
- Takahashi, Y. K., Langdon, A. J., Niv, Y., & Schoenbaum, G. (2016). Temporal Specificity of Reward Prediction Errors
- Signaled by Putative Dopamine Neurons in Rat VTA Depends on Ventral Striatum. Neuron, 91(1), 182-193. doi:
- 726 10.1016/j.neuron.2016.05.015
- 727 Tingley, D., Alexander, A. S., Kolbu, S., de Sa, V. R., Chiba, A. A., & Nitz, D. A. (2014). Task-phase-specific dynamics of basal
- forebrain neuronal ensembles. Frontiers in Systems Neuroscience, 8, 174. doi: 10.3389/fnsys.2014.00174
- van der Meer, M. A. A., & Redish, A. D. (2011). Theta Phase Precession in Rat Ventral Striatum Links Place and Reward
- 730 Information. Journal of Neuroscience, 31(8), 2843–2854. doi: 10.1523/JNEUROSCI.4869-10.2011
- West, E. A., & Carelli, R. M. (2016). Nucleus Accumbens Core and Shell Differentially Encode Reward-Associated Cues after
- 732 Reinforcer Devaluation. Journal of Neuroscience, 36(4), 1128–1139. doi: 10.1523/JNEUROSCI.2976-15.2016
- Wiener, S. I., Shibata, R., Tabuchi, E., Trullier, O., Albertin, S. V., & Mulder, A. B. (2003). Spatial and behavioral correlates in
- nucleus accumbens neurons in zones receiving hippocampal or prefrontal cortical inputs. *International Congress Series*,
- 735 1250(C), 275–292. doi: 10.1016/S0531-5131(03)00978-6
- Yun, I. A., Wakabayashi, K. T., Fields, H. L., & Nicola, S. M. (2004). The Ventral Tegmental Area Is Required for the Behavioral
- and Nucleus Accumbens Neuronal Firing Responses to Incentive Cues. *Journal of Neuroscience*, 24(12), 2923–2933. doi:
- 738 10.1523/JNEUROSCI.5282-03.2004