Behavioral/Systems/Cognitive

Moment-to-Moment Tracking of State Value in the Amygdala

Marina A. Belova, Joseph J. Paton,* and C. Daniel Salzman

Departments of *Neuroscience and Psychiatry, W. M. Keck Center on Brain Plasticity and Cognition, Kavli Institute for Brain Sciences, and Mahoney Center for Brain and Behavior, Columbia University Medical Center, and New York State Psychiatric Institute, New York, New York 10032

As an organism interacts with the world, how good or bad things are at the moment, the value of the current state of the organism, is an important parameter that is likely to be encoded in the brain. As the environment changes and new stimuli appear, estimates of state value must be updated to support appropriate responses and learning. Indeed, many models of reinforcement learning posit representations of state value. We examined how the brain mediates this process by recording amygdala neural activity while monkeys performed a trace-conditioning task requiring fixation. The presentation of different stimuli induced state transitions; these stimuli included unconditioned stimuli (USs) (liquid rewards and aversive air puffs), newly learned reinforcement-predictive visual stimuli (conditioned stimuli (CSs)), and familiar stimuli long associated with reinforcement (fixation point (FP)). The FP had a positive value to monkeys, because they chose to foveate it to initiate trials. Different populations of amygdala neurons tracked the positive or negative value of the current state, regardless of whether state transitions were caused by the FP, CSs, or USs. Positive value-coding neurons increased their firing during the fixation interval and fired more strongly after rewarded CSs and rewards than after punished CSs and air puffs. Negative value-coding neurons did the opposite, decreasing their firing during the fixation interval and firing more strongly after punished CSs and air puffs than after rewarded CSs and rewards. This representation of state value could underlie how the amygdala helps coordinate cognitive, emotional, and behavioral responses depending on the value of one’s state.

Key words: amygdala; monkey; state value; reinforcement learning; reward; punishment

Introduction

A common question in our daily lives is, “How are you?” This question assumes that we can evaluate our current situation, or “state.” Am I happy or sad? Thirsty or satiated? Is a reward coming, or is danger lurking? The concept of current state incorporates these features and more, and reinforcement learning theorists often define the concept of state broadly, as a combination of internal and external processes, stimuli, and events (Sutton and Barto, 1998; Dayan and Abbott, 2001).

When animals encounter stimuli in the environment, they often undergo changes in state. Some stimuli directly cause unpleasant or rewarding sensations; other stimuli predict future pleasant or aversive stimuli. Cognitive, behavioral, and physiological responses depend on how a state relates to current and future reinforcement, i.e., the state value. To promote survival, organisms should track state value to choose the best course of action and improve future assessments of state value to accurately predict reinforcement.

The value of states can be controlled through classical conditioning paradigms, in which subjects learn that rewarding or punishing unconditioned stimuli (USs) follow initially neutral conditioned stimuli (CSs) (Pavlov, 1927; Dickinson and Mackintosh, 1978). The amygdala receives inputs from multiple sensory systems carrying information about CSs and USs (Amaral et al., 1992; McDonald, 1998; Stefanacci and Amaral, 2002) and mediates learning about the relationship between environmental cues and reinforcement (LeDoux, 2000; Everitt et al., 2003; Baleine and Killcross, 2006; Paton et al., 2006; Belova et al., 2007). In primate amygdala, different populations of neurons are involved in learning the association between conditioned visual stimuli and rewards or punishments (Paton et al., 2006). We have shown previously that one population of neurons fires more to stimuli paired with liquid reward (positive “value-coding” cells), and another population of neurons fires more to images paired with air puff (negative “value-coding” cells) (Paton et al., 2006).

Here we determined whether amygdala neurons encode value in relation to sensory events that can induce transitions to different states. Monkeys performed a trace-conditioning task in which three novel CSs were paired with one of three USs: large liquid reward, small liquid reward, or aversive air puff. Three
types of stimuli were presented: a fixation point (FP) that monkeys looked at to initiate trials; the three visual CSs; and three USs, which differed in sensory modality and valence. If a stimulus elicited approach or defensive behavior, we operationally defined it as causing a transition to a state with positive or negative value, respectively. We asked whether amygdala neurons reflect the value of states induced by such sensory events. For this to be true, amygdala neurons must (1) encode value in relation to each sensory event (FP, CS, and US), and (2) integrate information about value across sensory modalities. Evidence that neurons integrate value information across sensory modalities could be obtained by examining the response profile of neurons responding most strongly to a CS when it is associated with aversive air puff. If these neurons also respond differentially to CSs associated with different magnitudes of rewards, it would suggest that these neurons encode the association between a CS and the value of USs, in which USs can differ in valence and sensory modality. Overall, the ability to track value in time across sensory stimuli, even when these stimuli differ in valence and modality, could indicate that amygdala neural activity reflects the value of an organism’s overall state.

Materials and Methods
Experimental paradigm and data collection
Our general methods for experimental control and electrophysiological recording in rhesus monkeys have been described previously (Paton et al., 2006). Monkeys sat in a Plexiglas primate chair (Crist Instruments) with their eyes 57 cm in front of a 21 inch Sony cathode ray tube monitor. Experimenters observed monkeys during experiments using a video camera feed. All animal procedures conformed to National Institutes of Health guidelines and were approved by the Institutional Animal Care and Use Committees at New York State Psychiatric Institute and Columbia University.

Trace-conditioning task. Monkeys learned the reinforcement value of three novel abstract images (fractal patterns) in each experiment through a trace-conditioning procedure in which each CS was associated with a reward delivery, which was followed by a large reward. The weak positive image was never associated with a reward.

We recorded neural activity from 145 neurons in the right amygdala of three rhesus monkeys (Macaca mulatta) weighing 5–10 kg (monkey V, 70 neurons; monkey P, 38 neurons; monkey L, 37 neurons). Based on magnetic resonance imaging (MRI), we positioned recording chambers directly over the amygdala. In each experiment, using a motorized multielectrode drive (NAN Instruments), we individually advanced four epoxide-insulated tungsten microelectrodes (FHG Inc.) into the brain through dummy-puncturing guide tubes positioned within a rigid grid of guide holes. Grids had holes spaced at 1.3 mm. Each guide tube had four inner guide tubes such that four electrodes could be advanced and independently controlled through the larger guide tube. We used the Plexon system for signal amplification, filtering, digitizing of spike waveforms, spike sorting using a principal component analysis platform (on-line with off-line verification), and data storage.

Monkeys either performed a fixation task or no task during the search for well isolated neurons. We included all well isolated neurons in this study.

Reconstruction of recording sites. The localization of the amygdala in each monkey was achieved using MRI. Each monkey was anesthetized with isoflurane, intubated, and imaged in a 1.5 Tesla research magnet in the Columbia University Department of Radiology. We imaged with an electrode inserted in a grid hole in the chamber and directed toward the amygdala. For reconstructing recording sites, we used a two-dimensional inversion recovery (IR) sequence, with 2-mm-thick slices and 0 intergap spacing and 0.234 × 0.468 mm within-slice resolution. These images were used to reconstruct the borders of the amygdala and recording sites. All recordings occurred primarily in the basolateral and central nuclei of the amygdala, in overlapping regions in all three monkeys.

Data analysis
Population analysis of monkeys’ behavior (licking and blinking). To construct population-average licking and blinking curves, we first assigned a 1 or 0 to each sampled time bin within each trial based on whether the monkey licked (blinked) or did not lick (did not blink) in that bin. We averaged the monkey’s responses across trials within each experiment and then averaged across experiments. Each monkey demonstrated graded licking behavior and differential blinking behavior.

Population-level peristimulus time histograms of neural activity. Peristimulus time histograms (PSTHs) were sampled in 10 ms nonoverlapping bins and convolved with a 50 ms at half-width Gaussian kernel for display purposes. Because on average monkeys learn CS–US associations within five or fewer trials, we removed the first four trials from initial and reversal learning sessions in each experiment for the PSTHs shown in Figures 2 and 4.

Receiver operating characteristic analyses. We used a receiver operating characteristic (ROC) analysis adapted from signal-detection theory (Green and Swets, 1966; Paton et al., 2006) for the following analyses.

Classification of cells as value coding. To classify a cell as encoding the value of conditioned images, we first defined two intervals: the visual stimulus (CS) and trace intervals (90–390 ms after CS onset and from 390 ms after CS onset until US onset, respectively) (Paton et al., 2006). If neural responses to images that underwent reversal during either (or both) the visual stimulus or trace intervals changed significantly and in opposite directions, the cell was classified as encoding value. Statistical significance was established by using an ROC analysis comparing activity in the 20 trials before and after reversal for both images; ROC values significantly different from 0.5 for both images (p < 0.05, permutation test with 1000 permutations, after Bonferroni’s correction) were used as our statistical criterion. Neurons were classified as encoding positive or negative value when both ROC values were greater or less than 0.5, indicating that cells responded more strongly to images associated with rewards (positive value) or air puffs (negative value), respectively. For subsequent analyses (described below), we used the interval (visual stimulus or trace) in which value coding was strongest (ROC values most different from 0.5).

Analysis of graded CS and US neural responses. To assess whether a cell that passed the test described above (see Classification of cells as value coding) encoded value during the CS or trace interval in a graded manner, we used an ROC analysis to construct two neural discrimination indices (NDIs) for CSs: the two NDIs compare activity on weak positive trials with strong positive trials (positive NDI) and to negative trials

Electrophysiological recordings. We recorded neural activity from 145
During these experiments, monkeys performed a trace-conditioning task (Fig. 1A) (see Materials and Methods). To assay learning during conditioning, we measured two behaviors during the trace interval of the conditioning procedure: anticipatory licking and blinking. Licking behavior indicated that monkeys discriminated among conditioned visual stimuli in a graded manner (Fig. 1B): they licked most on large reward trials, less on small reward trials, and almost not at all on negative trials. Blinking behavior indicated that monkeys learned to predict air puff (Fig. 1C).

To test whether amygdala activity provides a neural correlate of state value, tracking value consistently across the sequential presentation of multiple stimuli, we recorded activity from 145 neurons in three monkeys during performance of the trace-conditioning task. We focus first on neural activity during the fixation interval, before CS onset. The FP was an over-trained, positively valued stimulus, because monkeys had learned that centering gaze on the FP can lead to reward from the beginning of their training. Indeed, the fact that monkeys looked at the FP to begin a trial (an approach behavior) indicated that the FP had a positive value to our monkeys. Within these experiments, the FP was associated with reward on two-thirds of all reinforced trials (large and small reward trials). In contrast to FP value, CS value varied within an experiment, which was attributable to the different CSs used and to the reversal of contingencies within the task (see Materials and Methods).

We examined responses during the fixation interval separately in two populations of neurons: those that responded most strongly to either CSs associated with large rewards or CSs associated with aversive air puffs (defined as positive and negative value-coding neurons, respectively). Neurons were classified as encoding positive or negative value by comparing the responses to CSs paired with large rewards with responses to the same CSs when they were paired with air puffs. We used a ROC analysis to evaluate activity from the 20 trials before and after CS–US contingency reversal and assessed statistical significance with a permutation test (Paton et al., 2006). If both CSs elicited significantly higher activity when they were associated with either reward or punishment (in either the CS or trace interval, p < 0.05 after Bonferroni’s correction), we designated the cell as encoding positive or negative value, respectively. Seventy of 145 neurons (48%) met these criteria: 47 cells fired more strongly on large reward trials (positive value-coding cells), and 23 cells fired more strongly on air-puff trials (negative value-coding cells). The proportion of positive and negative value-coding cells did not vary significantly across monkeys ($\chi^2$ test, p > 0.05) (Table 1).

Neural responses during the fixation interval differed in positive and negative value-coding neurons. Given that monkeys positively value the FP, if neurons encode state value, then positive value-coding cells should tend to increase their firing during the fixation interval, and negative value-coding cells should tend to decrease their firing. Individual amygdala neurons that exemplify these response profiles are shown in Figure 2, A (a positive value-coding cell) and B (a negative value-coding cell). Overall, 30 of 47 positive value-coding neurons significantly increased their firing rate during the fixation interval, eight cells decreased their firing rate, and nine cells did not change their level of activity (Fig. 2D) (Wilcoxon’s test, p < 0.05). Among the 23 negative value-coding cells, 14 significantly decreased their firing rate after...
FP presentation, three increased their firing rate, and the remaining six had no change in activity (Fig. 2D) (Wilcoxon’s test, \( p < 0.05 \)). Thus, at a rate significantly greater than chance, positive value-coding neurons increase firing during the fixation interval and negative value-coding cells decrease firing (\( \chi^2 \) test, \( p < 10^{-5} \)). This \( \chi^2 \) test was also highly significant if we only considered neurons with sharp increases or decreases in responses to the FP (positive value-coding neurons: 23 of 30 neurons with increases; negative value-coding neurons: 9 of 14 neurons with decreases). In contrast, non-value-coding neurons increase or decrease their activity after FP appearance occurred with statistically indistinguishable probabilities (\( \chi^2 \) test, \( p = 0.67 \)) (Fig. 2D). On average, across each population, positive value-coding neurons increased firing during the fixation interval, negative value-coding cells decreased their firing, and non-value-coding cells showed a very weak, transient increase in response (Fig. 2C). These data suggest that amygdala neurons may track the value of states across multiple events during the trial, including FP and CS presentation.

The fact that positive, negative, and non-value-coding neurons respond differently to the fixation point was not attributable to a difference in firing on rewarded compared with punished trials, because responses did not differ across trial types. Indeed, the mean response during the fixation interval on rewarded compared with punished trials was highly correlated across cells (slope of linear fit, 1.0; \( r = 0.99; p < 10^{-10} \)). This is consistent with the fact that monkeys did not know which CS to expect during the fixation interval before CS onset. In addition, fixation interval activity on trials before and after CS–US reversal were highly correlated (slope of linear fit, 0.94; \( r = 0.99; p < 10^{-10} \)), ruling out the possibility that the differential responses during the fixation interval could be accounted for by different responses during initial and reversal learning.

Neural responses during the fixation interval also do not appear to be related to the uncertainty of reinforcement; unpredictability has been shown to modulate amygdala responses to stimuli (Belova et al., 2007; Herry et al., 2007; Whalen, 2007). Essentially, the fixation point appearance indicated the onset of a trial in which, on average, trials had a 0.53 probability of reward (half of which were large reward trials and half small reward trials), a 0.27 probability of punishment, and a 0.20 probability of no reinforcement. Uncertainty decreased after CS appearance, when a large reward, small reward, or punishment was specified, each occurring with a probability of 0.8. If neurons encode uncertainty in a monotonically increasing manner, responses should decrease during the CS interval compared with during the fixation interval. However, on average, positive value-coding neurons increase their firing compared with the fixation interval after a rewarded CS appears (mean increase, 5.12 spikes/s; \( p < 0.001, t \) test) and decrease their firing after a punished CS appears (mean decrease, 4.53 spikes/s; \( p < 0.01, t \) test) (Fig. 3A). In contrast, negative value-coding neurons do the opposite, decreasing their firing after rewarded CSs compared with fixation interval responses (mean decrease, 2.95 spikes/s; \( p < 0.05, t \) test) and increasing their firing after punished CSs appear (mean increase, 3.61 spikes/s; \( p < 0.05, t \) test) (Fig. 3B). Overall, therefore, neural activity does not appear to be correlated with reinforcement uncertainty.

Within the context of the trace-conditioning task, the value of the state initiated by CS presentation may simply be determined by the value of the US associated with that CS, i.e., the predictive value of the CS. Can neural responses to CSs therefore be described as encoding the value of the associated USs? If this is the case, then responses to CSs associated with small rewards should consistently be intermediate to responses to those associated with large rewards or punishments. Figure 4A–D displays PSTHs of single neurons that demonstrate this response property. In these
and the following PSTHs, we combined only postlearning trials to remove the effect of trials when monkeys had not yet learned the CS–US associations. The neurons shown in Figure 4, A and C, fire at a high rate on large reward trials, a low rate on punished trials, and an intermediate rate on small reward trials. In contrast, the neurons displayed in Figure 4, B and D, show a graded response pattern in reverse order: they fire most strongly on punished trials, least strongly on large reward trials, and again intermediate on small reward trials.

We next sought to quantify the degree to which neuronal responses to weak positive CSs were consistently intermediate to the other two CSs. By definition, all 70 value-coding cells fired differentially on large reward and air-puff trials. To characterize the responses to small reward trials compared with the other two trial types, we used ROC analyses to compute two NDIs. One index quantified the difference in responses to CSs on large and small reward trials (positive NDI for CSs). The other index quantified the difference between responses to CSs on small reward and air-puff trials (negative NDI for CSs). Figure 4E plots these two indices against each other. The vast majority of positive value-coding cells fall into the top right quadrant of the plot, whereas the majority of negative value-coding cells fall into the bottom left quadrant. The preferential distribution of neurons into these two quadrants, as opposed to being randomly distributed in all quadrants, was statistically significant (p < 0.005, χ² test). Thus, neuronal responses on small reward trials tend to fall between responses on large reward trials and negative trials in both positive and negative value-coding neurons. Encoding of value in individual amygdala neurons, therefore, can be described as a graded representation that spans positive and negative valences.

Of particular importance, negative value-coding neurons, which responded most strongly when a CS predicted an air puff, often responded differentially to CSs associated with different amounts of liquid reward. These amygdala neurons therefore did not simply represent the association of the CS with the sensory properties of a preferred US (in this case, the air puff). Instead, the responses to the three CSs reflected an integration of information about multiple reinforcers with different sensory properties. This response profile, therefore, may be best described as representing the association between the CS and the value of the corresponding US, in which USs can differ in sensory modality, valence, and magnitude.

To gain insight into the anatomical location of these amygdala neurons, we reconstructed recording sites using MRI (Fig. 5). We used a permutation test to identify neurons with positive and/or negative NDIs that are significantly different from 0.5 (p < 0.05). In these neurons, responses to CSs associated with small rewards differ from the responses to CSs associated with either large rewards or punishments. We found that neurons with positive and negative NDIs that are different from 0.5 could be found in a range of locations within the amygdala, including some locations that were deep and some locations that were in the anterior portion of the amygdala. According to published atlases of rhesus monkey anatomy, the central nucleus is located approximately in the dorsal 30% and the posterior 60% of the amygdala, whereas the basal, accessory basal, and lateral nuclei comprise a large portion of the remaining amygdala, including areas ventral to the central nucleus (Paxinos et al., 2000). Although the MRI does not permit delineation of amygdala nuclei, the reconstruction shows that neurons with positive and negative NDIs significantly different from 0.5 are unlikely to be restricted to the central nucleus, which is located in the dorsal and posterior aspect of the amygdala. At least some of these neurons are likely to lie within the basolateral complex.
Within the context of a classical conditioning procedure, the value of a state induced by presenting a CS is dictated by the value of the associated US. Therefore, the data presented so far are consistent with the notion that amygdala neural responses to the FP and visual CSs encode state values. We next examined whether this conceptual framework can also describe responses to the USs. Here we show that amygdala activity during the US interval does in fact track three different values of primary reinforcement that can differ in sensory modality. Examination of Figure 6, A and B, reveals that the responses to CSs associated with different USs is correlated with response levels after US delivery. These plots show the relationship among four indices computed for each value-coding cell: positive and negative CS NDIs and corresponding US NDIs. The CS NDIs were identical to those in Figure 4E, and the US NDIs were computed in the same manner, except calculated using activity during a reinforcement period 50–600 ms after US delivery. The positive NDIs are significantly correlated (r = 0.37; p < 0.005), as are the negative NDIs (r = 0.62; p < 0.005). These data indicate that amygdala neurons respond in a sustained and graded manner across both the CS and US intervals, although USs differ in sensory modality.

Thus far, we have described neural responses that are correlated with state values induced by FPs, CSs, and USs. To what extent is this neural representation correlated with behavior? We examined the relationship between neural activity in the amygdala and behavior by asking whether the variability in value-related encoding by amygdala neurons was correlated with the variability in monkeys’ behavior from day to day. Overall, monkeys’ behavior indicates that they value trials in a graded manner (Fig. 1B). However, in some sessions, the licking rate on small reward trials was more similar to licking on punished trials. Figure 7A–F shows plots of CS response NDIs (positive and negative, as defined above) scattered against BDIs for both licking (A–D) and blinking (E, F). BDIs were constructed the same way as NDIs, except that they compare the amounts of time spent licking and blinking during the last 500 ms of the trace epoch. A positive licking BDI > 0.5, for example, indicates that more time was spent licking before a large reward than before a small reward or before an air puff. For positive value-coding cells, positive CS response NDIs were correlated with positive licking BDIs (Fig. 7A, r = 0.55, p < 0.0001). This relationship did not hold for negative value-coding neurons (Fig. 7B, r = 0.08, p > 0.05). Conversely, in negative value-coding cells, negative CS response NDIs were correlated with negative licking BDIs (Fig. 7D, r = 0.78, p < 0.0001), and this relationship did not apply for positive value-coding neurons (Fig. 7C, r = 0.14, p > 0.05). Negative blinking BDIs were correlated with negative CS response NDIs in both positive and negative value-coding neurons (Fig. 7E, r = 0.37, p < 0.01; Fig. 7F, r = 0.52, p < 0.05). Taken as a whole, the correlation between positive NDIs and positive BDIs was strongest in positive value-coding cells, and the correlation between negative NDIs and negative BDIs was strongest in negative value-coding cells. These observations raise the intriguing possibility that positive value-coding cells may preferentially contribute to discriminative ability toward the more positive end of the stimulus-value spectrum, whereas negative value-coding neurons may preferentially contribute to discriminative ability toward the more negative end of the stimulus-value spectrum.

**Discussion**

We have described neural activity suggesting that primate amygdala continuously represents the value of a monkey’s current state. We used a classical (trace) conditioning procedure in which state transi-
encoding the value of states. State values play a prominent role in theories of reinforcement learning, because they are critical elements of learning algorithms and of posited decision-making and action control mechanisms (Sutton and Barto, 1998; Dayan and Abbott, 2001; Dayan and Balleine, 2002).

The representation of state value, evident across the population of neurons that we have studied in the amygdala, may arise from the convergence in the amygdala of sensory signals from multiple modalities and inputs from neuromodulatory systems and parts of the cerebral cortex involved in more complex, higher-order functions (Amaral et al., 1992; McDonald, 1998; Ghashghaei and Barbas, 2002; Stefanacci and Amaral, 2002; Höistad and Barbas, 2008). Consistent with this, we have shown previously that amygdala neurons, in addition to having signals correlated with value, can represent the sensory properties of visual stimuli and of reinforcing stimuli (Paton et al., 2006, their Fig. 3a, Table 1). In principle, this representation of sensory stimuli may be transformed into a representation of value through computational steps implemented within the amygdala, and this may explain why some neurons, but not others, show value-related responses to CSs but not FPs or vice versa. In addition, of course, value-related signals could also reflect input to the amygdala that already carries value-related information.

Other investigators have proposed that the amygdala may support two functions in parallel: encoding CSs with respect to the sensory properties of the USs they predict (which, in rodents, is thought to involve the basolateral amygdala) and encoding the overall motivational value of events or stimuli (which is thought to involve the central nucleus) (Balleine and Killcross, 2006). Indeed, some theoretical frameworks developed to explain conditioning emphasize that, during learning, the brain can form several different associations; these include associations between the CS and the sensory properties of the USs, the CS and the unconditioned response to the US (stimulus–response learning), or the CS and a more abstract quantity, such as the value of the US (Dickinson, 1980; Mackintosh, 1983; Everitt et al., 2003) (but see Gallistel and Gibbon, 2000). In the current study, the same substance, water, was used as a US in both large reward and small reward trials. Single amygdala neurons often discriminate between the value of CSs on these trials. Moreover, for negative-value-coding neurons, the presumed preferred US is an air puff, but many neurons still respond differentially to CSs associated with different magnitudes of liquid rewards. Therefore, many amygdala neurons do not appear to simply report the sensory properties of an associated preferred US. In addition, we have shown previously that amygdala neurons do not fire in relation to the behavioral responses on this task, licking and blinking (Paton et al., 2006), so neural activity does not appear to represent the association between the CS and the measured behavioral responses. Consequently, we conclude that some amygdala neurons appear to encode the value of USs associated with CSs. In primates, at least, these neurons are located throughout the amygdala, including the basal and accessory basal nuclei. Many neurons in the basal and accessory basal nuclei project to sensory and prefrontal cortices (Amaral et al., 1992, 2003; Ghashghaei and Barbas, 2002; Ghashghaei et al., 2007; Höistad and Barbas, 2008). Thus, the extent to which parallel processing of different aspects of CS–US associations occurs in discrete amygdala nuclei in primates, rather than intermixed within nuclei, remains an important target for future investigation.

Most previous studies investigating the representation of value in the brain have used operant tasks, have typically investigated the value of only one type of stimulus or action and have investigated only one valence of reinforcement, i.e., reward (Kawagoe et al., 1998; Platt and Glimcher, 1998; Leon and Shadlen, 1999; Tremblay and...
Schultz, 1999; Cromwell and Schultz, 2003; McCoy et al., 2003; Dorris and Glimcher, 2004; Roesch and Olson, 2004; Sugrue et al., 2004; Samejima et al., 2005; Tobler et al., 2005; Padova-Schioppa and Assad, 2006). Therefore, it remains unclear which the brain areas studied, which include the parietal, cingulate, dorsolateral prefrontal, and orbitofrontal cortices, as well as the basal ganglia and dopamine neurons in the midbrain, encode the ongoing value of states. In many of these studies, specific behavioral responses were required to obtain reinforcments that differed in reward value. Notably, even during such operant tasks, pavlovian associations may be formed as subjects learn to make specific adaptive responses based on sensory stimuli. For example, in operant tasks, monkeys often make saccadic eye movements to a visual target to indicate their choice. Because the selection of different target stimuli may result in different types or amounts of reinforcement, the target stimuli themselves may be conditioned. Thus, the value of specific actions could be built in part from computations of the value of states arising after presentation of action targets. The current data indicate that this computation is likely to be reflected in amygdala activity representing state value.

One outstanding question concerns whether the amygdala represents state value across a wider range of conditions. As noted in Introduction, the value of one’s state can be influenced by both internal and external factors. In the current paper, we investigated neural activity in relation to manipulations of external variables (the presentation of FFs, CSs, and USs), whereas internal variables presumably remained relatively constant during the brief recording sessions. Given the role of the amygdala in processes resulting from changes in internal state, such as satiation induced by reward devaluation procedures (Baxter and Murray, 2002), it will be important to determine whether and how amygdala neurons track state value when it is manipulated by factors such as more distant reward history.

A neural representation of state value in the amygdala could subserve multiple functions. First, decisions and subsequent actions, as well as a variety of physiological, cognitive, and behavioral responses that constitute what we call emotions, may in part be based on a representation of state value. Second, state values play a prominent role in learning algorithms, which posit that a computation of the difference in value of two successive states generates prediction error signals that drive learning (Schultz et al., 1997; Sutton and Barto, 1998; Dayan and Abbott, 2001). In principle, a representation of state value in the amygdala could play a role in all of these processes, albeit through different anatomical pathways (Amaral et al., 1992). Elucidating the specific manner by which neural circuits involving the amygdala perform these functions remains a critical goal for understanding how the brain implements learning, reasoning, and feeling.

References