Immediate changes in anticipatory activity of caudate neurons associated with reversal of position-reward contingency

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ABSTRACT

The primate caudate nucleus plays a crucial role in transforming cognitive/motivational information into eye-movement signals. A subset of caudate projection neurons fire before a visual target's onset. This anticipatory activity is sensitive to position-reward contingencies and correlates with saccade latency, which is shorter toward a rewarded position. We recorded single-unit activity of caudate projection neurons to examine the dynamics of change in anticipatory activity immediately after switches of the position-reward contingency. Two monkeys performed a visually-guided saccade task where only one position was associated with reward. The position-reward mapping remained constant within a block, but was reversed frequently between blocks without any indication to the monkey. Therefore, the switch could be detected only by unexpected reward delivery or unexpected lack of reward. After the switch, both saccade latency and anticipatory activity showed reliable changes already in the second trial, whether or not the first trial was rewarded. However, anticipatory activity in the second trial was generally higher if the first trial was rewarded, and the measured saccade latencies could be better explained by the difference in anticipatory activity between the two caudate nuclei. We suggest that anticipatory activity of caudate neurons reflects the reversal set of reward-position contingency.
INTRODUCTION

The control of goal-directed behavior includes the ability to quickly adapt to novel environmental situations. Flexible adaptation requires not only simple stimulus-response-reward associations but also detecting basic events and rules. Many animals show the ability to utilize rules and adjust behavioral repertoires accordingly (Miller et al. 2003). Although there are numerous ways to characterize "rules", the present study concerns about generalizability or transferability. For example, learning becomes progressively easier as a result of previous experience with similar problems. This increased ability to solve similar problems, namely learning set, is thought to reflect the affirmation or rejection of hypotheses or rules (Harlow 1949). In a simplest form, learning sets can be investigated as "reversal sets" (Harlow, 1950; Meyer, 1951; Treichler and Petros, 1983), where animals learn to switch several stimulus-reward associations. The present study aimed at examining neuronal indications of a reversal set in the primate caudate nucleus, more specifically, immediate changes in caudate anticipatory activity upon switches of the position-reward contingency.

Many studies of behavioral switches in primates exist, but investigations of the underlying neural mechanisms have begun only recently, after advances in neuronal recording and experimental paradigms allowed researchers to study single neurons over entire learning episodes within an experimental session (Assad et al. 1998; Chen and Wise 1995a,b; Mitz et al. 1991; Nakamura et al. 1998; Tremblay et al. 1998; Tremblay and Schultz 2000). Neuronal changes paralleling behavioral changes during learning were found in various cortical and subcortical regions (Chen and Wise 1995a,b; Mitz et al. 1991; Nakamura et al. 1998; Niki et al. 1990; Rolls et al. 1996; Thorpe et al. 1983; Tremblay et al. 1998; Tremblay and Schultz 2000; Watanabe 1990; Pasupathy and Miller 2005). Some studies showed rapid neuronal changes following changes in stimulus-response (e.g., Assad et al. 1998) or stimulus-reward contingency (e.g., Tremblay et al. 1998).

It has been suggested that the basal ganglia play a critical role in various selection/switching behaviors (e.g., Rolls 1994; Redgrave et al. 1999). The striatum (caudate nucleus, putamen, and ventral striatum) is the main input station of the basal ganglia and receives glutamate-mediated excitatory inputs from all areas of the cortex, as well as afferents from the thalamus and limbic structures such as the hippocampus and amygdala (Parent 1990; Parthasarathy et al. 1992).
Neurons in the primate striatum show a variety of activity related to rewards, expectation of external events that are behaviorally relevant (reward-predicting or movement-eliciting stimuli), and movement preparation (Alexander et al. 1986; Apicella et al. 1992; Hikosaka et al. 1989a,b,c; Rolls 1994; Schultz et al. 1992; Watanabe et al. 2003a). The striatum is therefore thought to be one of the primary sites where sensory, motor, cognitive, and motivational signals interact. In most cases, the expectation of reward appears to be a central component of the striatal activities (Hollerman et al. 1998; Schultz et al. 1992). We chose the head and body of the caudate nucleus as the target recording sites because these subregions are known to be related to the generation of saccadic eye movements (Hikosaka et al. 1989a; Hikosaka et al. 2000; Watanabe et al. 2003a) as well as the coding of position-reward contingencies (Kawagoe et al. 1998; Lauwereyns et al. 2002; Takikawa et al. 2002).

The main purpose of the present study was to find manifestations of immediate and robust changes of neural activity. To this end, we examined changes in oculomotor behavior (saccade latency) following switches of position-reward contingency and recorded neuronal activity from the primate caudate nucleus. We focused on the spatially-tuned anticipatory activity of primate caudate neurons. In many caudate neurons, the anticipatory activity is tuned to the position-reward contingency, not simply to the position or the reward expectation. Some caudate neurons exhibit reward associations with other stimulus features such as color (Lauwereyns, Takikawa, Kawagoe, Kobayashi, Koizumi, Coe, Sakagami and Hikosaka 2001), indicating that the caudate anticipatory activity can represent various types of reward association and expectation. It also shows a clear relationship with saccade latency, presumably reflecting the animal's motivational state or response bias (movement preparation) toward the rewarded position (Lauwereyns et al. 2002). Our current hypothesis for the functional significance of spatially-tuned anticipatory activity is that it represents motivationally-biased motor signals, which may be projected to the superior colliculus (Hikosaka et al. 2000), where both bias-type and gain-type modulations are observed (Ikeda and Hikosaka 2003). Yet, importantly, the anticipatory activity itself does not trigger eye movements but modulates subthreshold activity in the superior colliculus. This way, they eventually modify the goal-directed behavior (eye-movements in this case) in a reward dependent manner. Thus, the caudate anticipatory activity appears to be suitable for examining the relationship between internal motivational sets (e.g., position-reward associations) and oculomotor behaviors.

To efficiently study changes in behavior and neuronal activity, we employed a
reversal-learning paradigm (Assad et al. 1998; Rolls et al. 1996). Within a block of 20 completed trials, the reward was mapped consistently onto one target position (Figure 1). The position-reward association remained constant within a block but was reversed frequently and automatically between blocks. The switch of the position-reward contingency occurred without any indication to the monkey. Consequently, the animal had to learn the switched contingency by trial and error, specifically by detecting an unexpected reward delivery or an unexpected reward omission. This feature allowed us to investigate the role of previous experiences of position-reward associations. Specifically, we were interested in whether changes in saccadic latency and neuronal activity depend on the reward history (i.e., whether the first trial was rewarded or not, and whether the first and second trials were in same or different reward conditions).

**MATERIALS AND METHODS**

**Subjects and Surgery.** Two adult male Japanese monkeys (monkey A and monkey B; *Macaca fuscata*) were used (body weight 6.0 - 7.5 kg). The monkeys received dry pellets and small amounts of fresh fruit or vegetables in their home cages. During periods of training and experiments, the monkeys' access to water in the cage was controlled and monitored.

We implanted a head-holding device, a chamber for unit recording, and a scleral search coil under general anesthesia. The monkey was sedated with ketamine (4.6-6.0 mg/kg) and xylazine (1.8-2.4 mg/kg) given intramuscularly, and then general anesthesia was induced by intravenous injection of pentobarbital (4.5-6.0 mg/kg/hr) with butorphanol tartrate (0.02 mg/kg/hr). After the skull was exposed, 10-15 acrylic screws were bolted into it. The screws acted as anchors to which a plastic head holder and chamber were fixed to the skull with dental acrylic resin. A recording chamber (antero-posterior: 42 mm; lateral: 30 mm; depth: 10 mm) was placed over the fronto-parietal cortices, tilted laterally by 35 deg in the coronal plane and was aimed at the head and the body of the caudate nucleus based on magnetic resonance imaging (Hitachi, AIRIS, 0.3T, Tokyo). A scleral eye coil was implanted in one eye for monitoring eye position (Judge et al. 1980). The monkey received antibiotics (sodium ampicillin 25-40 mg/kg intramuscularly each day) after the operation. All surgical and experimental procedures conformed to the NIH Principles of Laboratory Animal Care (NIH publication no. 86-23, revised 1985) and were approved by the Juntendo University Animal Care and Use Committee.
**Behavioral task.** The monkey sat in a primate chair inside a dark sound-attenuated room with his head being immobilized. The visual stimuli were small red spots of light, 0.2 degree in diameter, back-projected onto a tangent screen by LED projectors. In each trial the monkey was required to direct and maintain his gaze at a central fixation spot during a first fixation ("pre-target") period of 1500 ms. After the pre-target period of 1500 ms, the fixation spot disappeared and a peripheral target appeared at 20 degrees to the left or to the right. The monkey had to make a saccade within 500 ms to within 3 degrees of the target position. An auditory tone of 800-Hz rectangular waveform followed each completed saccade. If the monkey made a fixation break or a late or inaccurate saccade, the same trial was repeated.

To investigate the influence of incentive on eye-movement behavior and caudate neuronal activity, we used an asymmetrical reward schedule (biased-saccade task: Figure 1). Within a block of 20 completed trials, reward (= a drop of water) was mapped consistently onto one target position and never on the other position. Since the position of the target was randomized and counterbalanced within a block, the monkey was rewarded in only half of the trials in a given block. The position-reward contingency remained constant within a block but was reversed frequently (22 - 51 times during behavioral sessions, 6 - 16 times during recording sessions) and automatically, without any indication to the monkey or any pause between blocks. Consequently, the switch of the position-reward contingency could be detected only on the basis of the unexpected reward delivery and the unexpected lack of reward. During behavioral sessions, the inter-trial interval (ITI) was fixed within a session but varied among sessions (1500, 3000, 6000 ms).

**Electrophysiological recording.** Eye position was measured with a standard magnetic search-coil (MEL-25, Enzanshi-Kogyo, Tokyo) technique (Judge et al. 1980), digitized at 500 Hz, and stored with event times for offline analysis. During recording sessions, action potentials of single neurons were recorded with tungsten electrodes (FHC, Inc. Bowdoinham, Maine, impedance 1.5 - 3 M ohm). Microelectrodes were advanced perpendicularly to the cortical surface, using an oil-driven micro-manipulator (MO-95, Narishige, Tokyo). The action potentials were amplified, filtered (500 Hz to 2kHz) and processed by a window discriminator (MDA-4 and DDIS-1, BAK Electronics, Germantown, MD). We selected extra-cellular neural activity of presumed projection neurons, which show very low spontaneous activity (0.01-1 Hz), but not of presumed interneurons, which show irregular tonic activity (2-10 Hz; Aosaki et al. 1994; Kimura...
For the purpose of the present study, we searched selectively for neurons that showed anticipatory activity during the pre-target period while the monkey performed the task. When we encountered such a projection neuron by visual inspection, we proceeded with recording the neuron in as many trials as possible. During neuronal-recording sessions, the ITI was fixed at 3000 ms. We conducted behavioral and recording sessions separately because it was impractical to keep single neurons isolated for multiple sessions and a comparison between neural activity and saccade latency within the same session was practically difficult.

**Data analysis.** We used the following procedure to determine the time of saccade initiation. An eye movement was judged as a possible saccade if its velocity and acceleration exceeded predetermined threshold values (30°/s and 90°/s², respectively). To be accepted as a saccade, (1) the velocity must exceed 45°/s after the onset, (2) this suprathreshold velocity must be maintained for at least 10 ms, and (3) the total duration must be longer than 25 ms. The end of the eye movement was determined when the velocity became lower than 40°/s. These threshold values were determined empirically by applying them to sample saccades, which led to almost perfect detection of saccades in the trained monkeys.

Neurons were classified as 'anticipatory neurons' (see Figure 2) if they showed a statistically reliable increase in the average number of spikes in the window of -1500 ms to 0 ms from target onset ('anticipatory activity') as compared to the activity before the onset of the fixation spot (from -1000 ms to 0 ms from fixation onset). All comparisons of average firing rates were evaluated by two-tailed t-tests, using a significance level of \( P < 0.05 \). After selecting anticipatory neurons, we examined whether their anticipatory activity systematically changed depending on the position-reward contingency and showed a significant difference between the two position-reward contingencies (unpaired t-test between contralateral versus ipsilateral reward conditions, \( P < 0.05 \)).

For both saccade latency and anticipatory activity, we performed reward-history analyses in order to examine how changes in saccade latency and anticipatory activity depended on the history of reward delivery in the preceding trial. The following paragraph presents our reasoning.

After the switch of the position-reward contingency, the animal would experience a 'surprising' position-reward event. For instance, in a left-reward block, the animal would make quick saccades to the left position and delayed saccades to the right position (Watanabe et al. 2003b). In the first trial after the contingency switch (now in a right-reward block), if the target
appears on the left, the animal would make a quick saccade to it (because the animal could not know about the contingency switch) but receive no reward (unexpected omission of reward). In the second trial, if the target again appears on the left, the saccade latency would become longer. However, what would the saccade latency be if the target appears on the right in the second trial? After the contingency switch, the animal had not encountered an event where a rightward saccade was associated with reward. If the latency of the saccade to the right position becomes significantly shorter than those in the previous block, this suggests that the reversal set for this particular task is effectively learned (Harlow, 1950; Meyer, 1951; Treichler and Petros, 1983). Observing these behavioral results, we then asked the same question about the anticipatory activity of caudate projection neurons (Figure 2). Would caudate anticipatory neurons change their activity immediately after the first ('surprising') trial?

RESULTS

Behavioral results. For behavioral sessions, monkey A completed 469 blocks, and monkey B completed 319 blocks. For both monkeys, saccade latency clearly changed after the switches of the position-reward contingency (Figure 3, upper panels). Learning curves were stable during the experiment after the extensive training (more than 6 months). After learning reached asymptote levels (trial 6 through trial 20), saccade latency was significantly shorter in rewarded trials than unrewarded trials for both monkeys (unpaired t-test, \( P < 0.001 \)). The changes in saccade latency occurred mainly within the first few trials.

Neuronal database. In recording sessions, we encountered a total of 426 neurons in three caudate nuclei of the two monkeys. Among them, 338 neurons were judged as projection neurons by their low spontaneous activity, and the remaining 88 neurons were classified as tonically active neurons (Aosaki et al. 1994). Of the 338 putative projection neurons, 104 neurons (104/338, 31 \%) appeared to have task-related activity, with 46 neurons (46/104; 44 \%) that were judged to show elevated anticipatory activity as compared to inter-trial activity. Forty-one of these neurons (16 neurons from monkey A, 25 neurons from monkey B) sufficient data for statistical analyses (at least 160 trials, i.e., 8 reversals). All of these 41 neurons showed a statistically reliable increase in the neuronal activity in the pre-target period as compared to the control period (two-tailed paired \( t \)-test, \( P < 0.05 \)), and so were classified as anticipatory neurons (see Figure 2). Thirty-one of the 41
anticipatory neurons (76 %) systematically changed their anticipatory activity depending on the position-reward contingency. Twenty-five of the 31 contingency-sensitive anticipatory neurons showed stronger anticipatory activity when the contralateral position was associated with reward than when the ipsilateral position was associated with reward (25/31; 81 %; Contra-bias neurons). The remaining six neurons (6/31; 19 %) will be referred to as Ipsi-bias neurons.

Neuronal results. We focused on the neuronal data from the 31 anticipatory neurons that showed sensitivity to the position-reward contingency, and analyzed the changing dynamics of anticipatory activity after contingency switches and its dependency on previous reward history. Neuronal activity was normalized within each neuron with respect to the neuron's maximum firing rate, and the data from all appropriate trials (in error-free blocks) were combined. The lower panels of Figure 3 show the mean (normalized) firing rate as a function of the number of trials after the contingency switch. The blue line shows mean anticipatory activity in blocks where reward was associated with the neurons' preferred positions (preferred blocks; contralateral for Contra-bias neurons, and ipsilateral for Ipsi-bias neurons). The red line represents the mean anticipatory activity in blocks where reward was associated with positions opposite to the neurons' preferred positions (non-preferred blocks; ipsilateral for Contra-bias neurons, and contralateral for Ipsi-bias neurons). The anticipatory activity showed a remarkable plasticity contingent upon the position-reward mapping (lower panels of Figure 3), which appeared to parallel the changes in saccade latency (upper panels of Figure 3), consistent with Lauwereyns et al. (2002). The changes in anticipatory activity occurred mainly within the first few trials.

Reward-history analyses. For simplicity, only blocks in which the monkeys made correct responses for the next two trials after a contingency change were included in the reward-history analyses.

For behavioral data: Obviously, in the first trial after a contingency switch, the monkey would expect reward based on the previous (opposite) contingency. We refer to rewarded and unrewarded trials after the contingency change as R and U, respectively (Figure 4). The second trials after the contingency change consisted of 4 types: RR (first rewarded --> second rewarded),

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1 Plotting saccade latency and neural activity as a function of trial number (Figure 3) partly showed the rapid behavioral and neural changes but did not unquestionably demonstrate an immediate and robust change. This was because there were four possible trial types for both behavioral measure (saccade latency) and neural measure (anticipatory activity), and the data shown in Figure 3 were from combinations and means of these trial types.
UR (first unrewarded -> second rewarded), RU (first rewarded -> second unrewarded), UU (first unrewarded -> second unrewarded). Mean saccade latencies in the initial two trials after the contingency switch are shown in the upper panels of Figure 5. In the first trials after the contingency switch (R and U), the saccade latency was shorter for unrewarded directions (U) than for reward directions (R) (unpaired t-test $P < 0.01$). This indicates that the monkeys followed the previous (opposite) position-reward contingency. In the second trials, the saccade latency already showed significant changes; rewarded trials tended to produce shorter saccade latencies than unrewarded trials (unpaired t-test $P < 0.01$). This held true for all four trial types: RR, UR, RU, and UU. In other words, the saccade latency changed just after completing a single (surprising reward/unrewarded) trial following the position-reward contingency switch.

For neuronal data: The reward-history analysis was performed on the neuronal data from the 31 neurons. The data of the two monkeys were combined since the results were similar between them (lower panels of Figure 5). The reward-history analysis for neuronal data was not so straightforward as that for behavioral data because anticipatory caudate neurons are sensitive to position-reward contingency and activated before the target presentation (Lauwereyns et al. 2002; Takikawa et al. 2002; see also Figure 2). That is, the reward condition (and target position) in the ongoing trial is irrelevant for the analysis of the neuronal data. If the change in neural activity is based on simple position-reward association, anticipatory activity in the second trial would depend on the particular reward condition and target position of the first trial. On the other hand, if the reversal set has been well established, anticipatory activity would be determined mainly by the reward-position contingency of the current block ($p =$ in the neuron's preferred block, or $n =$ in the neuron's non-preferred block). In the first trial of a preferred block ($n \to p$), the neuron's anticipatory activity occurs before the current trial is known to be R from U. Thus, the neuron's state is denoted as $0-p$. After receiving the unexpected reward in the first trial (R), the neuron’s state becomes $R-p$; if the first trial was unrewarded, the neuron’s state becomes $U-p$. Similarly, in the first trial of the non-preferred block ($p \to n$), the neuron’s state becomes $0-n$. The state in the second trial is $R-n$ and $U-n$ if the first trial is rewarded (R) or unrewarded (U), respectively.

The main question we addressed in the reward-history analysis was: Is a single trial sufficient to induce the change in neuronal activity, irrespective of the trial type? In the first trials after the contingency switch, anticipatory activity was high in non-preferred blocks ($0-n$; red lines) and low in preferred blocks ($0-p$; blue lines; unpaired t-test $P < 0.01$). This was expected because there was
no explicit indication of the switch of the position-reward contingency between blocks. There were significant changes in neuronal activity in all the transitions from the first to the second trials (0-n to R-n; 0-n to U-n; 0-p to R-p; 0-p to U-p; unpaired t-test $P < 0.01$). Thus, both the unexpected reward delivery and the unexpected lack of reward delivery had immediate effects on the anticipatory activity of the caudate projection neurons. To summarize, the results supported the established reversal set at the level of caudate anticipatory activity.

Additionally, an inspection of the neuronal results (lower panels of Figure 5) suggested that the anticipatory neural activity in the second trials tended to be larger when the previous (first) trial was rewarded (R-p > U-p; R-n > U-n; i.e., solid lines > hatched lines). A two-way ANOVA on the data of the second trials (R-p, R-n, U-p, U-n; R-U vs. p-n) confirmed this observation; both main effects of position-reward contingency ($F(1,167) = 11.14$, $p < 0.01$) and of reward condition in the first trials ($F(1,167) = 3.98$, $p < 0.05$) were significant, with no significant interaction ($F(1,167) = .029$, $p = 0.86$). In short, the position-reward contingency and the presence of reward seemed to have independent influences on the anticipatory activity.

**DISCUSSION**

The present study investigated the dynamics of changes in saccade latency and in caudate anticipatory activity while the monkey experienced frequent uncued switches of the position-reward contingency. In general, the saccade latency was sensitive to the reward expectation; the expectation of reward facilitates the generation of a saccade toward the rewarded position (Lauwereyns et al. 2002; Watanabe et al. 2003b). In the present experiment, there was no cue that the monkey could use to predict the occurrence of the contingency switch (unless the monkeys explicitly counted successfully completed trials, of which we found no indication). Therefore, in the first trial after the contingency switch, as the monkey followed the position-reward contingency of the previous block, the saccade latency was longer for a rewarded position and shorter for an unrewarded position. Then, contrary to expectation, the monkey received the water reward or did not receive it. After those 'surprising' reward events, the saccade latencies showed significant changes, compared with those in the first trials. This is not so surprising when the first and the subsequent trials were in the same reward condition (RR and UU). However, the significant changes in saccade latency were observed even when the previous reward
conditions were different from that in the current trial (UR and RU). In these conditions, the monkey had not encountered the position-reward association of the current trial after the contingency switch. This suggests that the monkeys established a reversal set and employed it effectively following each position-reward contingency switch (Harlow, 1950). The reversal set is related to the rule of the asymmetrical reward paradigm (i.e., if one position was rewarded, the other position was not rewarded).

Anticipatory activity in many caudate neurons is sensitive to position-reward contingency (Lauwereyns et al. 2002; Takikawa et al. 2002). Similar to the changes in saccade latency, the changes in the anticipatory activity required only a single trial after the contingency switch. Importantly, the present study showed that the two types of unexpected reward events were both effective in inducing the significant changes in the anticipatory activity. In other words, in order to induce the change in the caudate anticipatory activity, it was not required for the monkey to experience the two possible position-reward associations. This is again a clear sign of the application of the reversal set. Evidently, the roles of the primate caudate nucleus are not limited to simple stimulus-response or stimulus-reward associations. These findings lead us to propose that the primate caudate nucleus can reflect cognitive sets (exemplified by reversal sets), to induce immediate changes in neural activity and oculomotor behaviors.

Another interesting observation is that the anticipatory neural activity in the second trials tended to be larger when the previous (first) trial was rewarded (Figure 5). For both monkeys, the experience of reward in the first trials heightens caudate anticipatory activity, irrespective of the neuron's preferred position. However, such overall activities in neuronal activity were not consistently related to the saccade latencies (Figure 6 left panel). On the other hand, the neuronal bias (expressed as difference between mean activity in the preferred and non-preferred conditions) can reliably be related to the bias in saccade latency (expressed as difference between mean latency of rewarded and that of unrewarded saccades; see Figure 6 right panel). The neuronal bias roughly corresponds to the difference in anticipatory activity between the two caudate nuclei, since the preferred condition for most neurons in the right caudate (i.e., left-rewarded) corresponds to the non-preferred condition for most neurons in the left caudate. That is, the bias in saccade behavior could be comprehended more reliably in terms of a neuronal competition between two opposing motor preparation processes in the caudate nuclei on the opposite sides.
Comparison with other studies

Researchers have investigated neural plasticity in various brain areas by using learning sets, where single neurons can be studied over entire learning episodes (Assad et al. 1998; Chen and Wise 1995a,b; Mitz et al. 1991; Nakamura et al. 1998; Tremblay et al. 1998; Tremblay and Schultz 2000). For example, Schultz and colleagues examined how reward expectation-related activity is modulated during learning in the monkey striatum, showing how the striatum neurons learn novel stimulus-reward associations (Schultz et al. 2003). Although the present study has similarities with previous learning set experiments, it has at least two new and important features.

First, the present study investigated the effect of an uncued switch of the stimulus(position)-reward contingency on striatal, in particular caudate, neurons. For example, in previous learning set experiments on striatal neurons (e.g., Tremblay et al. 1998), there were cues indicating the beginning of a new session. In previous studies of caudate neurons in our laboratory (Kawagoe et al. 1998; 2004; Takikawa et al. 2002), there were long breaks between blocks, which could signal the change of the position-reward contingency. Several studies have employed paradigms similar to ours (i.e., no transition cue between blocks) to force monkeys to learn stimulus-response or stimulus-reward contingencies by trial and error (Assad et al. 1998, 2000; Rolls et al. 1996; Thorpe et al. 1983), but these studies focused mainly on prefrontal neurons and did not take into account possible roles of reward history (type of feedback) in behavioral switch and neural change.

Second, the present study is the first to examine the effect of reward-history on both saccade latency and caudate neural activity during reversal learning. Effects of previous trials on saccade latency and neural activity have been reported in the superior colliculus by using simple sensorimotor tasks (Dorris et al. 1999, 2000; Fecteau and Munoz 2003) and in the caudate nucleus by using a memory-guided saccade task (Itoh et al. 2003). These studies demonstrated the effects of reward history on established performance, which could be traces of simple residual or priming effects. Our study focused on the learning process before the reward-based performance was established. During associative learning, behavioral and neuronal changes associated with the contingency switch involve more than such passive effects of reward-history (Assad et al. 2000; Miller et al. 2003; Wallis et al. 2001). This is particularly true when changes in behavior and
neural activity are based partly on rules rather than simple stimulus-response associations.

**Origins of immediate neuronal changes in caudate neurons**

The immediate neuronal changes could be due to afferent modulatory inputs to caudate projection neurons. One possible brain region that may provide such modulatory input is the substantia nigra pars compacta (SNc) (Kawagoe et al. 2004; Takikawa et al. 2004). Midbrain dopamine neurons in primates respond to unexpected reward and conditioned reward-predicting stimuli by phasic activation. They also show depression of activity when reward is unexpectedly omitted (Schultz et al. 1997) or shifted (Hollerman and Schultz 1998). Dopamine neurons thus appear to signal the extent to which the rewarding outcome deviates from the prediction (prediction error; Schultz et al. 1997; Schultz 2002) and have the formal characteristics of reinforcement signals for acquiring new behavioral reactions (Barto 1994; Montague et al. 1996; Schultz, 2002; Waelti et al. 2001). In the present study, both unpredicted reward delivery and unpredicted reward omission were effective in inducing the immediate neural plasticity of the caudate anticipatory neurons. Since the dopamine response transmits a reward prediction error with a short latency through diverging connections to the striatum and frontal cortex (Schultz 2002), it can serve as "broadcasting" signal for quick changes in multiple brain regions, which might induce the immediate change in caudate neuronal activity (Kawagoe et al. 2004; Takikawa et al. 2004).

Another candidate brain region that participates in the immediate change of caudate anticipatory activity is the prefrontal cortex. The primate prefrontal cortex has massive anatomical connections with the basal ganglia, forming several distinct parallel functional loops (Alexander et al. 1986). The striatum, including the caudate nucleus, is the primary region that receives the input from the prefrontal cortex. Recent investigations on prefrontal functions have pointed to their role in the adaptive changes of stimulus representation. During behavioral reversal, neurons in the prefrontal cortex show remarkable sensitivity to stimulus-reward (Rolls et al. 1996; Watanabe 1990; Watanabe et al. 2002) and stimulus-response (Assad et al. 1998; Niki et al. 1990) associations. Alternations of these associations produce rapid gain and loss of response to the alternate stimuli, which are accompanied by equally rapid changes in the animal's behavior. Such changes in prefrontal neural activity may signify changes in the rules that the animal follows. Moreover, recent studies have shown that the prefrontal cortex is the key site where abstract rules
are acquired and represented. Some sensory and motor-related neurons in the prefrontal cortex change their activity on the basis of the abstract rule (or behavioral context) that the animal is presently employing (Assad et al. 2000; Miller et al. 2003; Sakagami et al., 2001; Wallis et al. 2001; White and Wise 1999). This rule-sensitive neural activity in the prefrontal cortex could be involved in changing the neural processing that underlies the behavioral change (e.g., in the premotor cortex, Wallis and Miller 2003; in the anterior cingulate, Shima and Tanji 1998) and could signal changes of the behavioral context to caudate projection neurons.

In line with the possible involvement of the prefrontal cortex, anticipatory activation for task specific events has also been reported in the prefrontal and premotor cortex in primates (Coe et al. 2002; Kobayashi et al. 2002; Sakagami and Niki 1994; Tremblay and Schultz 2000; Watanabe 1996). Since the central part of the caudate nucleus (from which we recorded the projection neurons) receives massive projections from these cortical regions (Parthasarathy et al. 1992; Selemon and Goldman-Rakic 1985; Yeterian and Pandya 1991), it is possible that caudate anticipatory activity is derived from cortical anticipatory inputs. In fact, anticipatory activity in the frontal eye field shows (Bruce and Goldberg 1985) characteristics similar to that in the caudate nucleus. The anatomical connection between these brain structures (monosynaptic projections from the frontal eye field to the caudate) implies a possible functional relationship between them. Also, Kobayashi et al (2002) have shown that some neurons in the dorsolateral prefrontal cortex show similar anticipatory activity before the explicit instruction. Furthermore, a recent study have reported that, during associative learning in monkeys, neuronal activity in the striatum shows more rapid, almost bistable, changes than that in the prefrontal cortex (Pasupathy and Miller 2005). Therefore, it is possible that the basal ganglia lead the prefrontal cortex in learning new position-reward associations (Houk and Wise 1995; Bar-Gad, Morris, and Bergman 2003). Given these findings, it would be particularly interesting to record neuronal activity simultaneously in the frontal eye field and the caudate in future investigations.

The above mentioned hypotheses of afferent modulatory inputs partly and implicitly assume that motor preparation or bias in the caudate nuclei on the opposite sides independently influence on the activity of the superior colliculus. Another, not mutually exclusive, possibility is that the immediate changes in caudate activity (and consequently changes in saccadic latency) are implemented by a competition between two opposing motor preparation processes that are mutually inhibitory (as implied by Figure 6). When one of this population of neurons experiences
a "surprise" event on the first trial after the switch, its activity either increases or decreases depending on whether it was a rewarded or unrewarded trial. This increased or decreased activity causes the activity in the opposing network to decrease or increase accordingly. Yet, it is unclear such a mutually-inhibitory network is based on pre-existing anatomical connections or it is a consequence of long-term experimental training. It would be interesting to investigate how neuronal networks that enable the animal to form reversal sets are established.

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REFERENCES


Dorris MC, Taylor TL, Klein RM, and Munoz DP. Influence of previous visual stimulus or


FIGURE CAPTIONS

Figure 1  Biased saccade task. The monkey performed a visually-guided saccade task, where only one position was associated with water reward (Lauwereyns et al. 2002).

Figure 2  Anticipatory activity of a caudate projection neuron. This neuron was recorded from the left caudate nucleus of monkey B. Neural activity increased before the target presentation only in blocks where the right target (preferred position) was rewarded. Neural activity started just after the fixation onset, ramping up until the onset of the target, and then disappeared immediately. The anticipatory activities were present only in blocks where the right (contralateral) position was rewarded, confirming the sensitivity to position-reward contingency (Lauwereyns et al. 2002).

Figure 3  Upper panels: Saccade latency as a function of trial number after the contingency switch. Results from the two monkeys are shown separately. Vertical bar = 1 standard error of mean. Lower panels: Changes in anticipatory activity (normalized) of caudate projection neurons as a function of the trial number after the contingency switch. Changes in both saccade latency and anticipatory activity occurred within the initial few trials (shaded area).

Figure 4.  Possible trial combinations of two trials after the position-reward contingency switch (characters in ellipses; e.g., RR) and associated states of a subject and caudate neuron during the pre-cue period (characters in boxes; e.g., R-p). 'R' and 'U' designate 'reward trial' and 'unrewarded trial', respectively. The last characters show the current trial. These classifications were used in the history analyses of saccade latency and anticipatory activity. In the history analysis of anticipatory activity, it should be noted that caudate neurons had no information regarding the reward condition in the ongoing trial. Therefore, the last character must be either 'p' (in the neuron's preferred block) or 'n' (in the neuron's non-preferred block) for anticipatory caudate activity (i.e., during the pre-cue period).

Figure 5  Upper panels: Change in saccade latency in the initial switching phase of the position-reward contingency switch (two trials after a switch). Trials are categorized as described
in Figure 4. On the first trial after the contingency switch, rewarded trials led to longer saccade latencies and unrewarded trials led to shorter latencies. This was because the monkeys did not know the switch of position-reward contingency in the first trials. In the second trials, the saccade latencies already showed significant changes from those in the first trials. This held true even when the monkeys had not experienced the target position and the reward condition of the current trial after the contingency change (RU and UR). Lower panels: Changes in caudate anticipatory activity (normalized) in the initial learning phase of the position-reward contingency switch. In the first trials after the contingency switch, anticipatory activity was high in non-preferred blocks (0-n; red lines) and low in preferred blocks (0-p; blue lines) because there was no explicit indication of the switch of the position-reward contingency between blocks. In the second trials, the anticipatory activity already showed significant changes from those in the first trials, regardless of the reward condition in preceding trials.

**Figure 6**  Left panel: Mean saccade latency (combined rewarded and unrewarded trials) against mean neuronal activity (combined preferred and non-preferred conditions) in the second trials after the contingency switch. The neuronal activity was generally higher when the first trial was rewarded, but such elevation of anticipatory activity did not consistently lead to a difference in saccade latency. Right panel: *Difference* in mean saccade latency between rewarded and unrewarded trials (saccade latency bias) against *difference* in mean anticipatory neuronal activity between preferred and non-preferred conditions (neuronal bias). The neuronal bias is a good predictor of saccade latency bias. Note that both saccade latency bias and neuronal bias are hypothetical measures. There was no individual data point for saccade latency or neuronal bias. This was because the behavioral and recording sessions were conducted separately. Also, neuronal bias was determined by comparing blocks, whereas saccade latency bias was determined by comparing trials.
Figure 1
Figure 3
Figure 4
Figure 5
Figure 6