Behavioral/Systems/Cognitive

Neural Basis of Adaptive Response Time Adjustment during Saccade Countering

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Humans and macaque monkeys adjust their response time adaptively in stop-signal (countermanding) tasks, responding slower after stop-signal trials than after control trials with no stop signal. We investigated the neural mechanism underlying this adaptive response time adjustment in macaque monkeys performing a saccade countermanding task. Earlier research showed that movements are initiated when the random accumulation of presaccadic movement-related activity reaches a fixed threshold. We found that a systematic delay in response time after stop-signal trials was accomplished not through a change of threshold, baseline, or accumulation rate, but instead through a change in the time when activity first began to accumulate. The neurons underlying movement initiation have been identified with stochastic accumulator models of response time performance. Therefore, this new result provides surprising new insights into the neural instantiation of stochastic accumulator models and the mechanisms through which executive control can be exerted.

Introduction

Stochastic accumulator models provide a powerful explanation of the random and systematic variation of response times (RTs) in speeded decision-making tasks (Smith and Ratcliff, 2004; Bogacz et al., 2006; Gold and Shadlen, 2007). They account for RT and choice probability in terms of three basic parameters: baseline activity before accumulation begins, the rate of accumulation once it begins, and the threshold for terminating the decision process. A fourth parameter, residual processing time, includes the time between the presentation of a stimulus and the beginning of accumulation plus the time between reaching a threshold and executing the response. The neural instantiation of the processes described by these models was first elucidated with the discovery that responses are initiated when the random accumulation of activity in particular movement-related neurons reaches a fixed threshold (Hanes and Schall, 1996). Subsequently, several investigators have shown how the activity before choice responses of particular neurons in posterior parietal cortex, frontal eye field, and the superior colliculus can be identified quantitatively with the stochastic accumulation process described by race and diffusion models (Mazurek et al., 2003; Ratcliff et al., 2007; Purcell et al., 2010).

Stochastic accumulator models account for adaptation of RT to minimize errors and maximize rewards. This adaptation is commonly supposed to be accomplished through changes in the threshold of accumulation that triggers a response (Nakahara et al., 2006; Simen et al., 2006; Schall and Boucher, 2007; Forstmann et al., 2008; Ratcliff and McKoon, 2008). The hypotheses that adaptation of RT is accomplished through changes in the threshold of accumulation has not been tested directly by measurements in neurons identified with the parameters of stochastic accumulator models in monkeys optimizing RT to perform a task.

We addressed this question using data from macaque monkeys performing a saccade stop-signal (countermanding) task (Logan and Cowan, 1984; Hanes et al., 1998; Paré and Hanes, 2003; Brown et al., 2008). Humans and macaque monkeys adapt their performance in stop-signal tasks, producing longer RTs after successfully inhibiting a planned movement (Rieger and Gauggel, 1999; Emeric et al., 2007; Verbruggen and Logan, 2008c; Nelson et al., 2010). These data provide a unique opportunity to determine whether the neural implementation of systematic adjustments of RT is accomplished through a change of threshold, baseline, or accumulation rate or alternatively through a change in the onset time of accumulation. Our analysis is based on the identification of presaccadic movement-related activity with the accumulation process described by a stochastic accumulator model of stop-signal performance (Boucher et al., 2007). Several lines of evidence support this linking proposition (Schall, 2004).
First, the activity of these neurons in frontal eye fields (FEFs) and superior colliculus (SC) is modulated in a manner sufficient to control the initiation of eye movements in the stop-signal task. Saccades are initiated when the activity of these neurons reaches a threshold of discharge rate, and the variability in RT across trials arises from stochastic variation in the rate at which the activity accumulates to the threshold (Sparks, 1978; Hanes and Schall, 1996; Dorris et al., 1997). Thus, neural measures of the onset of accumulation, baseline discharge rate before the onset, rate of accumulation, and threshold can be identified with the corresponding parameters in a stochastic accumulator model.

Materials and Methods

Methods for collecting and analyzing the performance and neurophysiological data have been described in detail previously (Hanes et al., 1998; Paré and Hanes, 2003). Here we will summarize the basic elements of the task and emphasize the new approach taken with this dataset.

Stop-signal task performance measures. On no-stop-signal trials, monkeys were reinforced for shifting gaze to a target that appeared when the fixation spot disappeared (Fig. 1). On stop-signal trials, the fixation spot reappeared after a variable delay following target onset. The stop-signal delays ranged from 25 to 275 ms for the FEF data and from 25 to 230 ms for the SC data; the values were selected from one of four values with intervals of 40, 50, or 60 ms adjusted for each session such that at the shortest delay monkeys withheld the movement on >85% of trials and at the longest delay monkeys withheld the movement on <15% of trials. Monkeys were reinforced if they withheld the planned saccade (referred to as “signal-inhibit” or “canceled” trials) but not if they failed to inhibit the saccade (referred to as “signal-respond” or “noncanceled” trials). The percentage of trials on which a stop signal occurred ranged from 30% to 50%, and none of the findings varied with this quantity. For each data collection session, saccade RT and associated neuronal activity on no-stop-signal trials were sorted as a function of whether the preceding trial was a no-stop-signal or canceled trial. RT was measured in no-stop-signal trials preceded by another no-stop-signal trial or preceded by a stop-signal trial in which monkeys inhibited the response. In previous work, we have found that RT measures longer on trials following successful inhibition but is at all, after unsuccessful inhibition (Emerick et al., 2007; see also Nelson et al., 2010).

Data were selected on the basis of the quality of task performance and the characteristics of the neural activity. Criteria for task performance were sufficient numbers of each type of trial (>10) and probability of failing to stop ranging from <20% at the shortest stop-signal delay to >80% at the longest stop-signal delay.

Neuron functional classification. FEF and SC consist of a diversity of neurons that exhibit qualitatively different patterns of modulation in this task (Hanes et al., 1998; Paré and Hanes, 2003; Ray et al., 2009; see also Murthy et al., 2009). Our analyses focused on movement-related neurons and visually responsive neurons. These different types of neurons were distinguished using multiple criteria. These criteria are quantitative extensions of conventional criteria used to distinguish functional classes of neurons in visuomotor structures like FEF and SC.

One set of classification criteria were applied to data obtained during a memory-guided saccade task in which monkeys direct gaze to a fixation spot and maintain fixation after a peripheral target is flashed. A gaze shift to the remembered location can be initiated only after the fixation spot is removed. This procedure distinguishes visually evoked from movement-related activity. In the memory guided saccade task, movement neurons exhibit an accumulation of activity only before saccade initiation, while visually responsive neurons exhibit a consistent visual response with a latency <100 ms following target onset.

The stop-signal task provides additional classification criteria. Whereas movement neurons exhibit a pronounced modulation of discharge rate on canceled stop-signal trials coincident with stop-signal reaction time (SSRT), visual neurons exhibit either no modulation at all or a modulation well after SSRT on canceled stop-signal trials (Hanes et al., 1998; Paré and Hanes, 2003; Brown et al., 2008). Another recent analysis has demonstrated that visuomovement neurons are clearly distinguished from movement neurons by the timing and pattern of their modulation (Ray et al., 2009). Only movement neurons exhibit the progressive accumulation of activity that corresponds to stochastic accumulation, like that seen in the activation of the GO units in the interactive race model (Boucher et al., 2007).

On the basis of data obtained in the stop-signal task, movement neurons that met the following criteria were selected: (1) differential rate of growth of movement activity for short versus long RT on no-stop-signal trials; (2) negative modulation of discharge rate on canceled stop-signal trials occurring before stop-signal reaction time; (3) progressively higher rate on cancel versus movement neurons with longer stop-signal delays; and (4) no visual responses or only small visual transients followed by a return of discharge rate below or near baseline activity before the beginning of accumulation of movement activity; the average rate at onset time was 10 and 5 spikes/s for SC and FEF neurons, respectively. Visually responsive neurons met the following criteria: (1) consistent visual response with a latency <100 ms and (2) absence of modulation before the RT to the stop signal on canceled stop-signal trials. These neurons included both visual neurons, which were characterized by an absence of modulation of discharge rate before stop-signal reaction time in canceled trials, and visuomovement neurons, which showed no differential rate of growth of movement activity for short versus long RT on no-stop-signal trials (Ray et al., 2009).

Measures of neural activity. We examined the average neuronal activity measured during no-stop-signal trials as a function of whether the preceding trial was a no-stop-signal trial or a canceled stop-signal trial. Neural activity was measured through spike density functions constructed by convolving spike trains with a filter shaped like a postsynaptic potential (Hanes et al., 1998; Paré and Hanes, 2003). Our analyses focused on four characteristics of the activity of movement-related neurons that could contribute to the observed variation of RT: differences in the baseline firing rate, the threshold activation level at which saccades were initiated, the rate of accumulation of activation to the threshold, and the
onset of movement-related activity. Baseline activity was measured as the average discharge rate from −200 ms until the target appeared. Threshold activation was measured as the average discharge rate in the interval 10–20 ms before saccade initiation (Hanes and Schall, 1996). The onset of movement-related activity was measured with a sliding-window algorithm working backward from the peak of activity of movement neurons relative to target onset (Woodman et al., 2008). The onset of activity was defined as the first time point that met the following criteria: the spike density function did not increase significantly according to a Spearman correlation (α = 0.05) over a time window ranging from −10 ms to +10 ms, the correlation between spike density and time remained nonsignificant for 10 ms as the center of the window continued to move backward from the saccade; if any of the values of the spike density function within 10 ms forward of the onset time identified by the correlation algorithm fell below the activity level at that onset time, the onset time was then redefined as the time point at which the spike density function was lowest. For four of the movement neurons recorded in FEF, the number of model trials equaled the total number of observed trials. After the models were fit to behavioral data from each session with FEF or SC recordings, we tested for significant differences in onset, rate, and threshold using a t test on the best-fitting parameter values. Note that whereas the confidence interval for a Spearman correlation (alpha = 0.05), for a Student’s t test of best-fitting parameter values for rate, threshold, and onset were found using the subplex algorithm (Rowan, 1990), a variant of the Nelder–Mead simplex method (Nelder and Mead, 1965) that is well suited for optimizing stochastic models. Like most hill-climbing parameter search algorithms, subplex is susceptible to hitting local minima, so we started the subplex algorithm with at least 160 different starting points in parameter space for each dataset we fit. Following Ratcliff and Tuerlinckx (2002), we optimized the fit of each model to the behavioral data by minimizing a Pearson (χ²) statistic,

\[ \chi^2 = \sum \frac{(o_i - p_i)^2}{p_i}. \]

In this case, observed (o_i) and predicted (p_i) refer to the number of trials with RT’s falling into each quintile bin of the observed and predicted RT distribution. Simulated trials were rescaled so that that total number of model trials equaled the total number of observed trials. After the models were fit to behavioral data from each session with FEF or SC recordings, we tested for significant differences in onset, rate, and threshold using a standard t test on the best-fitting parameter values. Note that whereas the data presented in Table 1 used only trials in which the target fell in the neurons’ movement or receptive field, the data used for the model fits included trials at all target locations.

Table 1. Observed parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>No-stop-signal trials preceded by a no-stop-signal trial</th>
<th>No-stop-signal trials preceded by a canceled trial</th>
<th>Average difference ± 95% confidence interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>FEF (movement)</td>
<td>273 ± 29</td>
<td>291 ± 29</td>
<td>18 ± 4 **</td>
</tr>
<tr>
<td>FEF (visual)</td>
<td>268 ± 30</td>
<td>287 ± 37</td>
<td>19 ± 4 **</td>
</tr>
<tr>
<td>SC (movement)</td>
<td>258 ± 27</td>
<td>263 ± 29</td>
<td>5 ± 2 *</td>
</tr>
<tr>
<td>SC (visual)</td>
<td>240 ± 36</td>
<td>248 ± 33</td>
<td>8 ± 2 **</td>
</tr>
<tr>
<td>Threshold (spikes/s)</td>
<td>103.0 ± 44.3</td>
<td>104.2 ± 46.5</td>
<td>1.2 ± 2.3</td>
</tr>
<tr>
<td>Baseline (spikes/s)</td>
<td>132.0 ± 35.4</td>
<td>133.4 ± 37.0</td>
<td>1.4 ± 1.9</td>
</tr>
<tr>
<td>Rate of growth (spikes/s/s)</td>
<td>0.58 ± 0.40</td>
<td>0.61 ± 0.57</td>
<td>0.02 ± 0.05</td>
</tr>
<tr>
<td>Onset time (ms)</td>
<td>106.5 ± 38.7</td>
<td>123.6 ± 43.6</td>
<td>17.1 ± 4.4 **</td>
</tr>
<tr>
<td>SC movement</td>
<td>72.8 ± 9.2</td>
<td>64.4 ± 10.7</td>
<td>1.6 ± 0.9</td>
</tr>
<tr>
<td>SC movement</td>
<td>137.6 ± 24.9</td>
<td>143.0 ± 25.3</td>
<td>5.4 ± 2.8 *</td>
</tr>
<tr>
<td>Initial visual response magnitude (spikes/s)</td>
<td>71.0 ± 7.3</td>
<td>71.5 ± 6.7</td>
<td>0.6 ± 0.4</td>
</tr>
<tr>
<td>SC visual</td>
<td>70.2 ± 26.9</td>
<td>69.7 ± 24.2</td>
<td>−0.5 ± 2.5</td>
</tr>
<tr>
<td>SC visual</td>
<td>80.8 ± 40.4</td>
<td>81.4 ± 50.6</td>
<td>0.6 ± 5.6</td>
</tr>
</tbody>
</table>

All trials were included in the calculation of response time; only trials in which the target fell in the neuron’s response field were used to calculate the indicated parameters. Data in the first two columns are mean ± SD. *p < 0.05, **p < 0.01.

Results

The major innovation of this study was to analyze the temporal pattern of activity contingent on the history of behavioral performance. The results reported here are based on a reanalysis of data...
collected previously. The data for this study were derived from a subset of the single-unit recordings and performance measures from macaque monkeys performing a saccade stop-signal task that have been reported before; data were collected from the FEF of two male monkeys (Hanes et al., 1998) and from the intermediate and deep layers of the SC of two other male monkeys (Paré and Hanes, 2003). In total, 36 movement neurons (22 from FEF and 14 from SC) and 48 visually responsive neurons (26 from FEF and 22 from SC) met the criteria described above for these analyses. The data from the movement neurons in the FEF consisted of a total of 756 trial sequences in which a no-stop-signal trial followed a no-stop-signal trial (nostop-nostop) and 327 trial sequences in which a no-stop-signal trial followed a canceled stop-signal trial (canceled-nostop), with averages of 34 and 15 trials per neuron, respectively. The data from the movement neurons in the SC consisted of a total of 5478 nostop-nostop trial sequences and 1436 canceled-nostop trial sequences, with averages of 391 and 103 trials per neuron, respectively.

**Systematic variation of RT with trial history**

Overall across the 82 sessions in which both movement and visual neurons were recorded from the FEF and the SC, saccade RTs on no-stop-signal trials preceded by a canceled trial were significantly longer than RTs on no-stop-signal trials preceded by a no-stop-signal trial (260 vs 274 ms, t test \( p < 0.001 \)) (Fig. 2, Table 1). This RT difference happened to be larger in sessions when FEF neurons were recorded than in sessions when SC neurons were recorded (FEF monkey 1–276 vs 297 ms, \( p < 0.001 \); FEF monkey 2–252 vs 266 ms, \( p = 0.025 \); SC monkey 1–229 vs 236 ms, \( p < 0.001 \); SC monkey 2–268 vs 274 ms, \( p = 0.03 \)). The data from the two structures were obtained in different monkeys in different laboratories. The RT difference was smaller for the SC data, probably because these monkeys performed many more trials than did the monkeys providing the FEF data. Ultimately, the same qualitative optimization of performance was observed during collection of both FEF and SC data, constituting independent replications of the phenomenon. The quantitative differences in RT between FEF and SC sessions are not pertinent to this study, for it does not aim to compare FEF and SC neuronal activity.

Because RT measured on no-stop-signal trials following unsuccessful inhibition trials was elevated little if at all, neurophysiological data from these trials is not useful for evaluating the mechanism of adaptive adjustment. Moreover, because stop trials in which inhibition was unsuccessful were not reinforced, RT analyses of the sequences involving these trials could be confounded.

**Pattern of activity of movement neurons**

Individually (Fig. 3) and collectively (Fig. 4) across the population of movement neurons in FEF and SC, saccades were initiated when the activity reached a threshold that did not vary significantly with RT. The random variability of RT within an experimental condition arose from stochastic variation in the rate of accumulation of the discharge rate to the threshold (Sparks, 1978; Hanes and Schall, 1996; Dorris et al., 1997). On successful inhibition trials, activity in the same movement neurons begins to increase and then abruptly decreases before reaching threshold, so no response is triggered (Hanes et al., 1998; Paré and Hanes, 2003). The shift from increasing to decreasing firing rate occurs before the SSRT, so these neurons can be in the causal chain controlling initiation of the response.

To accomplish the systematic delay of RT after successful inhibition, the presaccadic movement activity could vary in one of four respects: increasing threshold, decreasing baseline, reducing rate of accumulation, or delaying the onset of accumulation after target appearance. We measured each of these parameters in the
The change in RT, then assuming no change of any other parameter, the observed difference in threshold we measured would produce a 2 ms increase in RT for the FEF sessions and a 1 ms increase for the SC sessions. To account for the difference of RT measured during FEF recordings, the threshold of FEF movement neurons must increase eightfold more than it actually does. To account for the difference of RT measured during SC recordings, the threshold of SC movement neurons must increase fourfold more than it actually does. This analysis demonstrates conclusively that adjustments of threshold necessary to produce the observed changes of RT do not occur in FEF and SC movement neurons. The same argument holds for changes of baseline.

We also found no significant decrease in average rate of accumulation (Fig. 5c). The rate of accumulation was measured in multiple ways. Because RT = onset + (threshold − baseline)/rate, each of the quantities we report are related. Table 1 reports a calculation of rate that afforded self-consistent values of RT, onset, baseline, and threshold.

If a change in rate of accumulation is the basis for the change in RT, then assuming no change of any other parameter, the observed difference in rate we measured would produce a 5 ms decrease of RT for the FEF sessions and a 3 ms increase for the SC sessions. To account for the difference of RT measured during FEF recordings, the rate of accumulation of FEF movement neurons must decrease by 10% when the measured value was an increase of 3%. To account for the difference of RT measured during SC recordings, the rate of accumulation of SC movement neurons must decrease twofold more than it actually does. This analysis demonstrates that adjustments of accumulation rate necessary to produce the observed changes of RT do not occur in FEF and SC movement neurons.

Because rate is in the denominator, small measurement errors can translate into large variation in RT. We measured the rate of accumulation in two other ways. First, we calculated rate = (activity at onset time)/(RT − onset time). Second, we calculated rate = (activity at 70% of threshold − activity at 30% of threshold)/(time when activity reached 70% of threshold − time when activity reached 30% of threshold). Table 2 lists the central tendencies of each of these values. Neither of the measures exhibited a significant difference that could account for the systematic change of RT with trial history. To summarize, these data illustrate the challenge in measuring the rate of accumulation in neural data and demonstrate that, in these data, no measurement provided evidence that the rate of accumulation could explain the systematic variation in RT.

In contrast, the average onset of accumulation for no-stop-signal trials preceded by no-stop-signal trials occurred significantly earlier than the average onset of accumulation in no-stop-signal trials preceded by canceled trials (Fig. 5d).

If a change in the onset of accumulation is the basis for the change in RT, then assuming no change of any other parameter, the observed difference in onset that we measured would produce a 17 ms increase of RT for the FEF sessions and a 5 ms increase for the SC sessions. These values correspond precisely to the observed 17 ms increase in RT for the FEF sessions and the 5 ms increase for the SC sessions.

These results indicate that the systematic variation in RT can be accounted for by variation of the onset of accumulation, but the data cannot rule out conclusively that trade-offs between parameters do not contribute some measure of variation as well. However, if such trade-offs occur, then it is hard to understand how the accumulation onset time could correspond so well to the observed difference in RT.
The difference in onset time was smaller for the SC neurons than for the FEF neurons. However, the difference in RT was also smaller during sessions when the SC data were collected than in sessions when the FEF data were collected. Thus, the qualitative patterns of behavioral and neural data were the same in SC and FEF datasets; both showed slowing of RT after canceled stop trials and both showed a corresponding shift in the onset of accumulation and not in baseline, rate of accumulation, or threshold. The quantitative differences between the FEF and SC datasets should not obscure the qualitative similarity in the relationships. Indeed, the combination of data across laboratories and neural structures increases confidence in the reliability of the general conclusions.

**Ballistic accumulator model**

The activity of movement neurons in FEF and SC has been described as instantiating a stochastic accumulation process (Carpenter and Williams, 1995; Hanes and Schall, 1996; Schall, 2004; Boucher et al., 2007; Ratcliff et al., 2007; Carpenter et al., 2009). To investigate this linking proposition further, we fit a linear ballistic accumulator model (Brown and Heathcote, 2008). The behavioral data consisted of the RT distributions of no-stop-signal trials preceded by either a canceled stop-signal trial or a no-stop-signal trial. The model was fit separately to the RTs from each of the 36 sessions during which the movement neuron data were being collected from the frontal eye fields or superior colliculus. RT on a simulated trial was given by the time at which threshold was reached plus a fixed ballistic movement time of 10 ms; this interval corresponds to the time before saccade initiation when omnipause neurons in the brainstem release inhibition on the medium lead burst neurons (Scudder et al., 2002).

Table 3 lists the values of the rate, onset, and threshold, plus predicted RTs for the two trial sequences when all parameters were free to vary. This model accounted for the RTs and their difference between the two trial sequences. It fit the difference of RTs in the FEF data through a significant difference in the onset time with no significant difference in threshold or rate of accumulation. No significant difference was observed for the SC fits, but as discussed below, the RT difference for these data was rather small.

**Pattern of activity of visual neurons**

One plausible explanation for the change in onset time with the increase of RT after canceled stop trials is a strategic adjustment in a stage of processing preceding the movement neurons. One earlier stage strongly implicated in this task is encoding the location of the target, which is represented in the activity of visually responsive neurons in FEF and SC. If RT slowing resulted from a strategic adjustment in an earlier stage of processing, then we should see different response latencies or different discharge rates in visually responsive neurons on no-stop-signal trials preceded by canceled trials compared to no-stop-signal trials preceded by no-stop-signal trials. However, in both the FEF and SC data, we found no systematic differences in the latency or the magnitude of the evoked responses of visual neurons measured in the different trial sequences (Fig. 6, Table 1). These results are inconsistent with the hypothesis that the RT slowing observed in this task was due to an adaptive adjustment in visual processing preceding the movement neurons in FEF or SC.

**Discussion**

We found that a systematic elevation of RT after successful stop-signal trials was accomplished through a change in the time when
the activity of movement neurons in FEF and SC first began to accumulate, and not through a change of threshold, baseline, or accumulation rate. We will argue that this new result provides new insights into the neural instantiation of stochastic accumulator models and the mechanisms through which executive control can be exerted.

Relation to previous work on RT adjustments
Numerous studies of speed–accuracy adjustments have been conducted with human participants but none using neurophysiological measures from macaque monkeys. Human brain imaging studies have implicated medial frontal cortical areas and the basal ganglia (e.g., Forstmann et al., 2008; see also Frank, 2006). ERP results obtained in various tasks including stop signal demonstrate changes both preceding and following the onset of the lateralized readiness potential (e.g., Osman et al., 2000; Rinckenaar et al., 2004; Upton et al., 2010). However, fMRI and ERP data provide insufficient temporal or spatial information to resolve alternative hypotheses about the mechanisms responsible for accomplishing the adjustment of RT that have been formulated through stochastic accumulator models of RT.

Movement neurons and stochastic accumulation
Stochastic accumulator models provide a powerful explanation of the random and systematic variation of RT in speeded response tasks (e.g., Ratcliff and Smith, 2004; Bogacz et al., 2006; Ratcliff and McKoon, 2008). We suggest that post-stop-signal slowing may rely on the same mechanisms as speed–accuracy adjustments. Typically, RT and choice probability are accounted for in terms of three parameters: baseline level before the accumulation begins, the rate of accumulation, and the threshold for terminating the accumulation process. A fourth parameter, residual processing time, is not considered formally part of the accumulation process. This residual time includes the time between the presentation of a stimulus and the beginning of accumulation of the decision variable (onset time) plus the time between terminating the accumulation and executing the response (motor time).

Stochastic accumulator models formulated to minimize errors and maximize rewards accomplish the adaptation through changes in the threshold to terminate accumulation (Bogacz et al., 2006; Nakahara et al., 2006; Simen et al., 2006; Ratcliff and McKoon, 2008). As yet, no model has proposed that the adaptation of RT involves changing the time when accumulation begins or the efferent delay time, although Verbruggen and Logan (2009) observed changes in onset in proactive adjustments of RT in the stop-signal task. The hypothesis that adjustment of RT is accomplished through changes in the threshold or rate of accumulation has not been tested directly by measuring those quantities in neurons that can instantiate the process described by stochastic accumulator models in monkeys optimizing RT to perform a task.

The neural instantiation of the processes described by stochastic accumulator models was elucidated with the discovery that responses are initiated when the random accumulation of activity in movement neurons in FEF reaches a fixed threshold (Hanes and Schall, 1996). Several lines of evidence support this linking proposition (Schall, 2004). (1) The temporal pattern of the stochastic accumulation process resembles the form of movement neuron activity (Hanes and Schall, 1996). (2) Models of stochastic accumulation processes map onto the temporal pattern of activity of movement neurons for tasks requiring categorization of a single stimulus (Ratcliff et al., 2007) and for visual search (Purcell et al., 2010). (3) The activity of movement neurons in FEF and SC modulates in a manner sufficient to control the initiation of eye movements in the stop-signal task. Saccades are produced despite the stop signal if the activity of these neurons reaches threshold too rapidly, and saccades are not produced on stop-signal trials when the activity of these movement neurons is inhibited before it reaches threshold (Hanes et al., 1998; Paré and Hanes, 2003). (4) Behavior in the stop-signal paradigm can be modeled very accurately as a race between stochastic GO and STOP processes (Logan and Cowan, 1984), and a neural network model with interacting GO and STOP accumulator units replicates the time course of the activation of the movement neurons (Boucher et al., 2007; Lo et al., 2009; Wong-Lin et al., 2010). Thus, neural measures of the onset of accumulation, baseline discharge rate before the onset, rate of accumulation, and threshold can be identified with the corresponding parameters in stochastic accumulator models.

Evidence to evaluate this linking proposition was sought by fitting a ballistic accumulator model to the performance. A significant change in the onset parameter accounted for the systematic difference in RT regardless of the architecture. Thus, the pattern of activation of the movement neurons is consistent with the form of activation of stochastic and ballistic accumulators fit to the behavioral data. The data obtained in this experiment provide evidence that subjects adapt by changing the onset time of the accumulation process, not the threshold, as commonly believed.

Visual processing and strategic adjustment
One could argue that the FEF and SC movement neurons are not part of the decision process that is adjusted strategically because the accumulation is delayed by preceding visual processing. In fact, when visual search is less efficient, resulting in systematically longer RT, the movement neurons in FEF exhibit a delay in accumulation (Woodman et al., 2008) that corresponds to the delay in time when the FEF visual neurons select the less salient target (Sato et al., 2001). It is well known that the responses of visual cortical neurons have longer latency and lower magnitude for stimuli with lower contrast or a nonpreferred feature. However, in the current task the stimulus on every trial was a single suprathreshold spot of light. While we do not have data from extrastriate cortical areas, there is no reason to believe that any of these neurons respond systematically differently to the target stimulus in different trial contexts. Visual neurons in FEF and SC receive converging inputs from several areas in which perceptual and cognitive processing occurs (Lui et al., 1995; Schall et al., 1995; May, 2006). Thus, strategic adjustments of visual processing should be evident in the activity of the visually responsive neurons in FEF and SC. No changes were seen. This seems to rule out the possibility that the changes in RT following a stop signal were produced in earlier visual processes.

The semantic, logical, and mechanistic distinctions between categorization (“decide that”) and response selection (“decide to”) (Schall, 2001) suggests that a sequence of stochastic accumulators instantiates successive stages of processing. If so, then different accumulators will limit performance in different task conditions depending on the difficulty of categorization and response preparation. In the saccade stop-signal task, visual encoding and stimulus-response mapping were simple and direct, so response preparation must be the rate-limiting process. We recently demonstrated how movement neuron activity can be understood as a gated accumulation of target salience evidence from FEF visual neurons (Purcell et al., 2010). We are now exploring...
the possibility that a gate between visually responsive neurons and movement neurons can be a source of executive control. Increasing the gate that sensory evidence must exceed to merit accumulation will delay the onset of accumulation of movement neuron activity and thereby increase RT. It can also increase accuracy, because delaying the initiation of a response allows the categorization process more time to arrive at correct description.

Sources of executive control

Different sources of executive control have been described that can delay movement activity onset time by extending the holding period of the top-down control module as in the model of Lo et al. (2009). First, the role of medial frontal cortex in the endogenous control of action is beyond dispute (e.g., Ridderinkhof et al., 2004; Nachev et al., 2008; Rushworth, 2008). Indeed, neurons in the supplementary eye field of macaques performing saccade countermanding produce activity that predicts but does not control whether monkeys will cancel successfully (Stuphorn et al., 2010; see also Chen et al., 2010; Scangos and Stuphorn, 2010). Also, weak electrical stimulation of sites in the supplementary eye field improves stop-signal task performance by increasing RT (Stuphorn and Schall, 2006).

Second, previous work has also suggested memory processes are involved in response adjustments in the stop-signal task (Verbruggen and Logan, 2008a,b; Verbruggen et al., 2008). Such a role for memory is consistent with the observation that the amplitude of the P3b event-related potential component is modulated with stop trial history (Upton et al., 2010) and that modulation in this ERP component has been linked to memory processes (see for a review Polich, 2007). The posterior source of this component suggests contributions from parietal or temporal cortex. Whatever their source, adaptive signals from these cortical areas could in turn be facilitated and effected through the basal ganglia circuit, so that premature accumulation of movement activity is prevented (Frank, 2006).

In summary, these new results indicate that accumulation threshold is not the exclusive locus of strategic control and demonstrate how the stochastic process producing responses can enact strategic adjustments to optimize performance. This result provides new insights into the neural instantiation of stochastic accumulator models and the mechanisms through which executive control is exerted.

References


