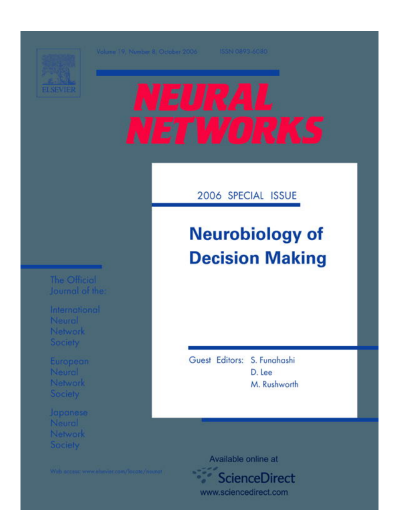
Provided for non-commercial research and educational use only.

Not for reproduction or distribution or commercial use.



This article was originally published in a journal published by Elsevier, and the attached copy is provided by Elsevier for the author's benefit and for the benefit of the author's institution, for non-commercial research and educational use including without limitation use in instruction at your institution, sending it to specific colleagues that you know, and providing a copy to your institution's administrator.

All other uses, reproduction and distribution, including without limitation commercial reprints, selling or licensing copies or access, or posting on open internet sites, your personal or institution's website or repository, are prohibited. For exceptions, permission may be sought for such use through Elsevier's permissions site at:

http://www.elsevier.com/locate/permissionusematerial



Available online at www.sciencedirect.com



Neural Networks

Neural Networks 19 (2006) 1013-1026

www.elsevier.com/locate/neunet

2006 Special Issue

Rapid decision threshold modulation by reward rate in a neural network

Patrick Simen^{a,b,*}, Jonathan D. Cohen^{a,c,1}, Philip Holmes^{b,d,2}

^a Center for the Study of Brain, Mind and Behavior, Princeton University, United States

Received 29 November 2005; accepted 1 May 2006

Abstract

Optimal performance in two-alternative, free response decision-making tasks can be achieved by the drift—diffusion model of decision making – which can be implemented in a neural network – as long as the threshold parameter of that model can be adapted to different task conditions. Evidence exists that people seek to maximize reward in such tasks by modulating response thresholds. However, few models have been proposed for threshold adaptation, and none have been implemented using neurally plausible mechanisms. Here we propose a neural network that adapts thresholds in order to maximize reward rate. The model makes predictions regarding optimal performance and provides a benchmark against which actual performance can be compared, as well as testable predictions about the way in which reward rate may be encoded by neural mechanisms. © 2006 Elsevier Ltd. All rights reserved.

Keywords: Reinforcement learning; Drift-diffusion; Decision making; Stochastic optimization

1. Introduction and background

A tradeoff between speed and accuracy is one of the hallmarks of human performance in cognitive tasks. Typically observed in controlled behavioral experiments in which participants are encouraged to respond quickly, the concept formalizes the common sense notion that rushing produces more mistakes. Despite the pervasive nature of this phenomenon and the long-standing recognition of it, relatively little research has addressed how organisms address the balance between speed and accuracy. Nevertheless, any successful model of the physical mechanisms underlying decision making will ultimately need to account for the speed–accuracy tradeoff: why does a tradeoff occur at all, and how do organisms change that tradeoff as conditions change?

In abstract models of decision making, especially those addressing two-alternative forced choice (TAFC) tasks, the

speed-accuracy tradeoff has traditionally been explained in terms of a task-dependent decision criterion, or threshold for termination of the decision-making process. This process is typically described as a progressive accumulation of evidence for each of the alternatives, the decision being made when the evidence in favor of one alternative versus the other exceeds the threshold (Laming, 1968; Luce, 1986; Ratcliff, 1978). If the threshold is low, the decision will be made quickly, but will be subject to noise. If the threshold is high, the decision process will take longer but will have greater time to 'average out' the effects of noise and therefore be more accurate. While a number of neural network models have addressed the mechanisms underlying TAFC task performance (e.g., Botvinick, Braver, Barch, Carter, and Cohen (2001), Brody, Hernandez, Zainos, and Romo (2003), Grossberg and Gutowski (1987), Usher and McClelland (2001) and Wang (2002)), thresholds in these models have typically been modeled simply as assigned parameters rather than as neural mechanisms in their own right.

Here we propose an explicit set of neural mechanisms by which thresholds may be implemented and adapted to maximize the reward rate. We do this by building on existing models of TAFC decision making that involve competing accumulators of evidence, one for each possible action. To each

^b Program in Applied and Computational Mathematics, Princeton University, United States

^c Department of Psychology, Princeton University, United States

^d Department of Mechanical and Aerospace Engineering, Princeton University, United States

^{*} Corresponding address: 209 Fine Hall, Washington Road, Princeton, NJ 08544, United States. Tel.: +1 609 258 6155; fax: +1 609 258 1735.

E-mail addresses: psimen@math.princeton.edu (P. Simen), jdc@princeton.edu (J.D. Cohen), pholmes@Math.Princeton.EDU (P. Holmes).

¹ Tel.: +1 609 258 2696; fax: +1 609 258 2574.

² Tel.: +1 609 258 2958; fax: +1 609 258 1735.

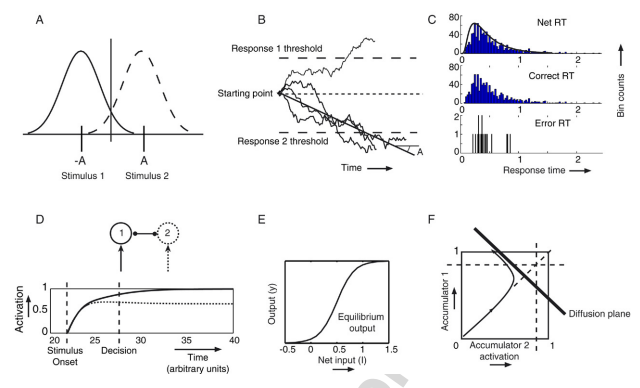


Fig. 1. A: The two stimulus distributions. B: Sample paths of a drift-diffusion process. C: Long-tailed analytical RT density (solid curve) and simulated RT histogram (top), correct RT histogram (middle), error RT histogram (bottom). D: Time courses of noise-free, mutually inhibitory evidence accumulation units with sigmoid activation functions. E: The sigmoid activation function. F: A smoothed sample path of mutually inhibitory accumulator activations in the (y_1, y_2) -phase space showing rapid attraction to a line (the 'diffusion plane') followed by drift and diffusion in its neighborhood.

accumulator we add a mechanism for implementing a response threshold: a unit with a high-gain, sigmoidal activation function that approximates a step function. We propose that such units control response initiation, and that they are triggered by critical levels of accumulated evidence in the accumulator units. We then define an algorithm for modulating thresholds and describe its implementation using a neurally plausible mechanism. The model consists of a set of stochastic differential equations that is equivalent to a classic connectionist recurrent neural network with five units. In Sections 2 and 3 we review relevant background before describing the model in Sections 4–6. We conclude with a discussion in Section 7.

2. The drift-diffusion model (DDM)

Sequential sampling models of decision making have long provided accounts of many regularities in response time (RT) and accuracy data in choice–reaction experiments (Luce, 1986). In sequential sampling, the stimulus is assumed to be a sequence of samples from one of two possible distributions, as, for example, in Fig. 1A. To determine which distribution is actually generating the stimulus, sampling is repeated and evidence in favor of one or another hypothesis is accumulated until a response criterion has been reached. Speed–accuracy tradeoffs can be explained in such models by shifts in the response threshold toward or away from the starting points of the decision variables: closer thresholds mean shorter RTs and higher error rates on average (Laming, 1968; Ratcliff, 1978).

In random walk versions of TAFC sequential sampling models, each evidential increment for one hypothesis reduces the evidence in favor of the other so that there is only a single decision variable: the difference in accumulated evidence for each hypothesis. (Fig. 1B shows this variable plotted against time for four different decisions. Fig. 1C shows the resulting response time distributions over many decisions.) Steps in the random walk are equivalent to increments of the total loglikelihood ratio for one hypothesis over the other, making the model equivalent to the sequential probability ratio test (SPRT) (Laming, 1968). This is theoretically appealing as a starting point for investigating the role of reward in decision making, because the SPRT is optimal in the sense that no other test can achieve higher expected accuracy in the same expected time, or, conversely, reach a decision faster for a given level of accuracy (Wald & Wolfowitz, 1948). However, the SPRT does not specify the optimal threshold for maximizing other values of potential interest, such as reward rate.

The drift-diffusion model (DDM) (Ratcliff, 1978; Smith & Ratcliff, 2004; Stone, 1960) is a version of the SPRT in which stimuli are sampled continuously rather than at discrete intervals. The difference between the means of the two possible stimulus distributions (see Fig. 1A) imposes a constant drift of net evidence toward one threshold, and the variance imposes a Brownian motion that may lead to errors. In monkeys, the continuously evolving firing rates of neurons in the lateral intraparietal sulcus (area LIP) have been related to competing accumulators that approximate the drift-diffusion process in oculomotor tasks (Gold & Shadlen, 2000, 2001; Roitman & Shadlen, 2002; Shadlen & Newsome, 2001). Similar findings have been reported for frontal structures

responsible for controlling eye movements (Hanes & Schall, 1996). Importantly, as shown in Section 3, the expected reward rate can be computed for the DDM (Gold & Shadlen, 2002), allowing learning mechanisms to be analyzed in terms of a well-defined optimization problem (Bogacz, Brown, Moehlis, Holmes, & Cohen, in press).

The DDM is defined as the stochastic differential equation (SDE)

$$dx = Adt + cdW, (1)$$

where A is the signal strength, $\mathrm{d}W$ denotes increments of an independently and identically distributed (i.i.d.) Wiener (white noise) process and the factor c weights the effect of noise. At any given moment, the distribution of possible positions of a particle moving in one dimension and governed purely by a Wiener process is given by a Gaussian distribution whose mean is the starting point of the particle (in our case, the decision variable), and whose variance is equal to the time elapsed since the start of the process. Brownian motion of this type causes diffusion of a substance within a liquid, from whence comes the term 'diffusion' in the name of the model. Nonzero drift A contributes a tendency for trajectories to move in the direction of the drift, producing a corresponding linear movement in the mean of the particle position distribution over time. Below, we will use the terms 'drift' and 'signal' interchangeably.

For models of this type to explain effects that are observed in human performance – including the speed–accuracy tradeoff, sequential effects such as post-error slowing, and speeded response to frequent stimulus alternations and repetitions (Luce, 1986) – their parameters must be adaptive on a short time-scale. Traditionally, parameters have been inferred by fits to behavioral data, and additional degrees of freedom have been added to models to explain different behavioral phenomena (e.g., Ratcliff and Rouder (1998)). However, in order to go beyond identifying the relevant degrees of freedom and toward the principles that govern the selection of specific parameter values, several questions left unanswered by this approach must be addressed. For example, on what basis do subjects select particular parameter values of the decision process (e.g., starting point or initial value of the decision variable, and threshold for termination)? Are subjects behaving so as to minimize errors, to maximize reward rate, or to do something else altogether? How are their parameters modified in response to ongoing experience? Only a limited number of studies have addressed these questions and, to our knowledge, none have addressed the question of neural implementation.

Below, we propose a neural network model that explains how the decision threshold can be adapted (and a tradeoff between speed and accuracy chosen) in order to maximize the reward rate over multiple trials. This is based on a simple neural network model that implements the DDM, as described in the next section.

2.1. Neural implementation of the DDM

The accumulation of evidence in the DDM can be approximated by a simple neural network with two 'decision'

units, each of which is assumed to be preferentially sensitive to one of the stimuli and also to be subject to inhibition from the other unit (see Fig. 1D), as proposed by Usher and McClelland (2001) (cf. Bogacz et al. (in press) and Gold and Shadlen (2002)). Each unit has a leak term, and therefore accumulates evidence for its corresponding stimulus subject to decay over time, while competing with the unit representing the other decision alternative. We and others have shown that, with suitable parameter choices, this model closely approximates the DDM.

Specifically, the evolving activation of each unit (indexed by i) is determined by an SDE, the deterministic part of which is

$$\dot{y_i} = -y_i - \beta y_i + I_i, \tag{2}$$

where I_i is the input, usually assumed to be a step function of time (corresponding to stimulus onset) and $-\beta y_j$ represents inhibition from the other unit(s). With the stochastic component of the activation function included, the pair of units is governed by

$$dy_1 = (-y_1 - \beta y_2 + I_1)dt + cdW_1,$$
(3)

$$dy_2 = (-y_2 - \beta y_1 + I_2)dt + cdW_2.$$
(4)

Here we assume linear or piecewise linear activation functions for ease of analysis (we will abandon this when we discuss threshold-crossing detectors, a case in which nonlinearity is critical). This assumption provides a useful approximation to a more realistic, sigmoid function (Cohen & Grossberg, 1983; Freeman, 1979) and is also consistent with the idea that attention acts to place processing units in their central, approximately linear range, where they are most sensitive to afferent input (Cohen, Dunbar, & McClelland, 1990). Sigmoids saturate near 0 at a small, positive baseline value, and also near a finite maximum which is typically rescaled to 1, thus avoiding the implausibilities of potentially unbounded or negative activations. For such units each equation of (3) and (4) takes the form

$$\dot{y}_i = -y_i + \sigma(-\beta y_j + I_i) + c dW_i, \text{ where}$$

$$\sigma(s) = \frac{1}{1 + e^{-\lambda(s - \gamma)}}.$$
(5)

Summing the noise-free linearized equations (3) and (4), we find that solutions approach an attracting line $y_1 + y_2 = (I_1 + I_2)/(1 + \beta)$ exponentially fast at rate $1 + \beta$. Differencing them yields an Ornstein–Uhlenbeck process for the net accumulated evidence $x = y_1 - y_2$:

$$dx = [(\beta - 1)x + I_1 - I_2]dt + cdW.$$
 (6)

If leakage and inhibition are *balanced* ($\beta = 1$) this becomes the DDM (Eq. (1)) with $A = I_1 - I_2$ representing the difference in inputs. See Bogacz et al. (in press), Brown et al. (2005) and Holmes et al. (2005) for further details and verification that the nonlinear system approximates this behavior.

Fig. 1F shows the evolution of the activations $y_1(t)$ and $y_2(t)$ over time. After stimulus onset, the system state (y_1, y_2) approaches the attracting line along which slower, diffusive behavior occurs as the state approaches one or another

boundary under the influence of the noisy signal. Projection of the state (y_1, y_2) onto this line yields the net accumulated evidence x(t), which approximates the DDM as shown in Fig. 1B.

3. A free-response, two-alternative forced choice task

To investigate the hypothesis that speed–accuracy tradeoffs are driven by a process of reward maximization, we consider an experiment in which subjects try to determine the direction of motion of moving dots on a screen, as in Roitman and Shadlen (2002). They are free to respond at any time after stimulus presentation, and a response terminates the stimulus. Initial results from such an experiment suggest that people adapt their speed–accuracy tradeoffs in a manner consistent with the goal of maximizing reward, and that this adaptation can happen quite quickly (Bogacz et al., in press; Simen, Holmes, & Cohen, 2005). People can also adapt their speed–accuracy tradeoffs in similar tasks in response to explicit instructions (Palmer, Huk, & Shadlen, 2005).

To provide a benchmark against which to measure evidence of adaptive behavior, we first describe optimal performance under this experimental paradigm. Assuming a constant rate of trial presentation, the expected reward rate over a sequence of trials in which correct responses are rewarded by 1 unit and errors by 0 can be expressed as follows (Gold & Shadlen, 2002):

$$RR = \frac{1 - ER}{DT + T_0 + RSI}. (7)$$

Here ER is the expected error rate (proportion of errors), DT is the decision time, T_0 is the residual latency (non-decision-making component of response time comprising stimulus encoding and motor execution times), and RSI is the response–stimulus interval (wait time from the last response to the next stimulus onset).

For the DDM, ER and DT, and hence RR, depend only on the signal-to-noise ratio A/c and threshold-to-signal ratio z/A, and we shall assume that these two parameters, as well as T_0 and the RSI, are held fixed within each block of trials. The following analytical expressions are derived in Busemeyer and Townsend (1993) (cf. Bogacz et al. (in press) and Gardiner (1985):

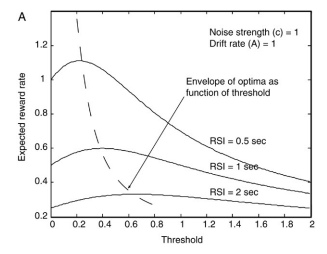
$$ER = \frac{1}{1 + e^{2Az/c^2}},\tag{8}$$

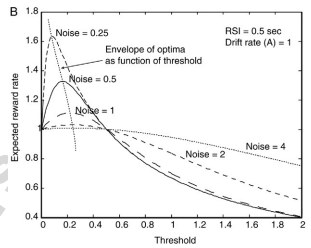
$$DT = \frac{z}{A} \tanh \frac{Az}{c^2},\tag{9}$$

and substituting them into Eq. (7) gives

$$RR\left(\frac{z}{A}, \frac{A}{c}\right) = \frac{1}{\left(T_0 + RSI + \frac{z}{A}\right) + \left(T_0 + RSI - \frac{z}{A}\right) \exp\left(-2\left(\frac{A^2}{c^2}\right) \cdot \frac{z}{A}\right)}.$$
(10)

Fig. 2 shows the expected reward rates given by Eq. (10) as a function of threshold z for various values of RSI (Fig. 2A),





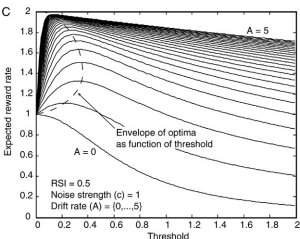


Fig. 2. A: Expected reward rate as a function of threshold for three different RSI conditions, with noise and drift held constant. B: Expected reward rate for several noise values, with RSI and drift held constant. C: Expected reward rate for a range of drift values with RSI and noise held constant.

noise (c; Fig. 2B) and drift rate (A; Fig. 2C) (Bogacz et al., in press). For all values of RSI, noise and drift, this function is smooth and has a unique maximum, indicating that there is a single optimal threshold for maximizing the reward rate, and that a gradient ascent algorithm can be used to find this

Table 1 Hill-climbing algorithm for threshold adaptation that maximizes the reward rate; after Myung and Busemeyer (1989)

Step	Instruction
1	Estimate the reward rate at an initial threshold
2	Randomly take a step upward or downward in threshold value
3	Estimate the reward rate at the new threshold
4	Compute the difference in reward rate estimates
5	Divide the difference by the size (or sign) of the change in threshold value to get an estimate of the gradient of the reward rate curve
6	If the estimated slope is positive, take another step in threshold value in the same direction; else take a step in the opposite direction; go to (3) and repeat.

optimum (although this approach faces some problems that we will consider below).

In each plot, a broken line connects the peaks of the reward rate curves, showing the reward rate and corresponding thresholds associated with optimal performance for different values of the DDM parameters (the 'envelope of the optima' in Fig. 2). Thus, any mechanism that seeks to optimize the reward rate for this decision process must be able to adapt its threshold to the values indicated in the plots in response to changes in task variables.

There is empirical evidence that human participants, performing well-practiced tasks, are capable of such adaptation over relatively short intervals (e.g., in as few as 5–10 trials) following a change in task conditions (Bogacz et al., in press; Ratcliff, Van Zandt, & McKoon, 1999). Several theories have been proposed for how such adaptations may occur (e.g., Busemeyer and Myung (1992), Erev (1998) and Myung and Busemeyer (1989)). However, these have typically been described in terms of discrete updating algorithms (for an example, see Table 1). While such algorithms provide a useful abstract specification of the component processes required for threshold adaptation, several challenges arise when considering how they may be implemented.

First, any reasonable reward rate estimation process takes time, but algorithms like that in Table 1 assume that threshold changes are made at discrete intervals. It is therefore undetermined how long the system should wait at a given threshold in order to develop a reasonable estimate of the associated reward rate before making a modification: with too few trials the estimate will be poor; with too many, convergence will be slow. Second, there is the question of step-size selection: step sizes that are too large cause oscillation of the threshold around the optimum value; step sizes that are too small again cause slow convergence. This problem can be addressed by introducing additional mechanisms that progressively reduce the step size, but this adds complexity to the model. These algorithms also typically require a memory of old reward rate values, and a means to compare new and old values in order to compute gradients. In the sections that follow, we describe a neural network model that addresses these issues. The model operates in continuous time, requires no explicit value comparison mechanisms, and achieves rapid and stable adaptation to threshold values that are nearly optimal.

We begin by describing an implementation of a neural network mechanism for detecting threshold crossing. We then describe a mechanism for reward estimation. Finally, we demonstrate how the latter can be used to adapt the threshold mechanism in order to maximize the reward rate.

4. Thresholds as an affine function of reward rate

The threshold in the DDM is a step function applied to the accumulated evidence (for evidence less than the threshold, the output is 0; for evidence greater than the threshold, the output is 1). In order to implement such a crisp function in a neural network, a McCulloch–Pitts neuron (McCulloch & Pitts, 1943) can be used, or an approximation based on a sigmoid unit with strong gain (see Fig. 3A). Inputs that fall below the inflection point of the curve will fail to activate the unit, while those that fall above it will activate the unit maximally. The effective threshold can be manipulated by providing a constant input (or bias) to the unit, with a positive bias in effect shifting the function to the left (decreasing the threshold), and a negative bias shifting it to the right (increasing the threshold).

In our model, subthreshold levels of excitation provided to threshold units from non-evidence-accumulating units will act to reduce the effective threshold with respect to the accumulators. An input x > 0, scaled by a positive synaptic weight w, reduces the effective threshold by wx. Thus, if a level z_{max} of evidence is required for threshold crossing in the absence of additional excitation, the level drops to $z_{\text{max}} - wx$. This defines an *affine function* of x (a linear transformation plus a constant): precisely the transformation required by the abstract threshold adaptation algorithm described in Section 6. Accordingly, our algorithm may be implemented by connecting one or more additional units to the threshold detectors with appropriate synaptic weights w and by setting the bias (γ in Eq. (5)) of the threshold detectors to z_{max} (see Fig. 9). We shall further discuss implementation, and possible neural substrates, in Sections 6 and 7.

5. Estimating reward rate

The threshold adaptation algorithm to be described in Section 6 seeks to optimize performance in the TAFC task by using a running estimate of the current reward rate to modulate behavior. Here we show how that estimate of reward rate can be computed by a linear filter or 'leaky integrator' that is incremented in response to reward impulses s(t) while decaying continuously over time (as in, e.g., Sugrue, Corrado, and Newsome (2004a)). The estimate, r(t), evolves according to

$$\dot{r}(t) = \frac{1}{k} \cdot (s(t) - r(t)). \tag{11}$$

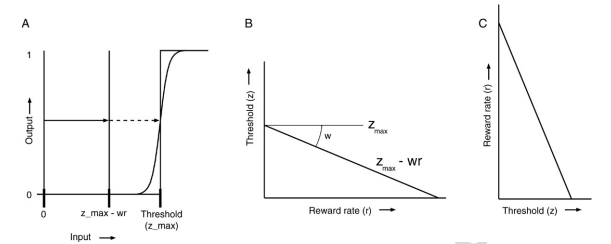


Fig. 3. A: Thresholds can be implemented by a simple step function, or McCulloch–Pitts neuron (McCulloch & Pitts, 1943), which in turn can be approximated by a sigmoid with strong gain. A threshold can be reduced from z_{max} to $z_{\text{max}} - wr$ by giving the unit additional excitation wr, representing weighted reward rate, so that evidence need only be accumulated to this level to produce a response. B: Threshold represented as a linear function of reward rate. C: By exchanging the vertical and horizontal axes, this prescription for threshold setting based on reward rate can be compared to the predicted reward rate as a function of threshold.

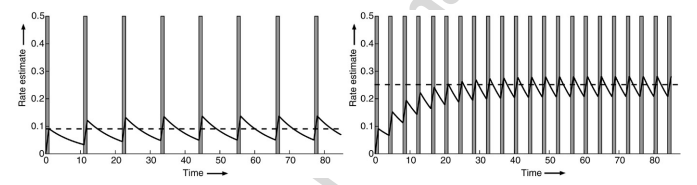


Fig. 4. Reward rate estimate r(t) (sawtooth curve) in response to a sequence of short reward pulses s(t) (gray rectangles). Here the rewards are pulses of height 1 and width 1. In both cases the estimated reward rate oscillates around the true frequency of reward pulses, shown by the dotted line. Left panel: slow pulse rate; Right panel: fast pulse rate.

Thus, r(t) at a given time is an exponentially-weighted time-average of the instantaneous reward signal s(t) in which the time constant k determines the speed of adaptation to changes in s(t). Following a step change in s(t), r(t) approaches s(t) exponentially at rate 1/k. More generally, large values of k attenuate high frequency fluctuations in s(t) (Oppenheim & Willsky, 1996). Discrete rewards can be modeled in continuous time as a sequence of narrow pulses or Dirac-delta impulses (see Fig. 4), in which case r(t) will approach the steady state mean of the reward rate.

Eq. (11) computes a continuous version of the time-discounted averaging usually seen in discrete-time reinforcement learning algorithms (cf. Doya (2000)). The averaging of Eq. (11) may be computed exactly by a linearized connectionist unit (Eq. (5)) in which recurrent self-excitation of strength k-1 is balanced against an activation function of slope 1/k, as can be seen by simple algebra:

$$\dot{y}(t) = -y(t) + \frac{1}{k} \cdot [s(t) + (k-1) \cdot y(t)]$$

$$= \left(\frac{k-1}{k} - 1\right) \cdot y(t) + \frac{1}{k} \cdot s(t)$$

$$= \frac{1}{k} \cdot (s(t) - y(t)). \tag{12}$$

A single unit can therefore compute the reward rate estimate required by the threshold adaptation algorithm. The use of a sigmoid rather than a linear activation function would make the relationship of Eq. (12) to Eq. (11) approximate rather than exact.

6. Threshold adaptation algorithm

We are now in a position to describe an algorithm that builds on the DDM of Eq. (1) and the reward rate estimator of Eq. (11). The algorithm models the DDM threshold as an affine function of a continuously evolving reward rate estimate, as suggested in Section 4. This is implemented in a neural network by exciting the threshold detector of Fig. 3A in proportion to the reward rate estimate. The resulting system has an attractor near the optimal

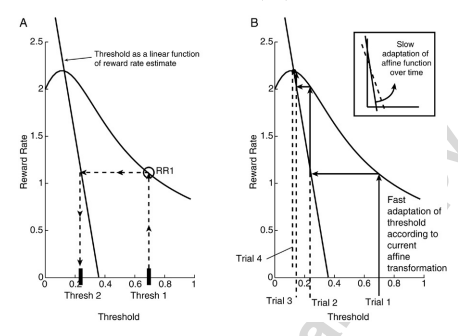


Fig. 5. A: Threshold selection after a single trial of the discrete-time description of the algorithm. B: Convergence to nearly optimal threshold after four trials.

threshold across a range of RSI conditions, and its large domain of attraction makes it robust to noise in reward rate estimates and in the activation of threshold and accumulator units.

6.1. Discrete time description

The algorithm proposed below can be understood most easily by first considering the discrete time version illustrated in Fig. 5. Suppose one starts with an arbitrary threshold value, Thresh1, as in Fig. 5A. The appropriate curve from Fig. 2 specifies the expected reward rate RR1 at that threshold value. After the first trial, *Thresh1* is updated in response to a new estimate of the reward rate (based on how long the trial took and whether it produced a reward or not) by mapping RR1 to Thresh2 by the affine transformation. The process is repeated to compute RR2, Thresh3, etc., and progress can be traced by a staircase or 'cobweb diagram' (cf. Jordan and Smith (1999)), as indicated in Fig. 5B. Given the unimodal shape of the expected reward rate function and the slope of the transformation line, rapid convergence to their unique intersection occurs, and we note that step sizes adapt automatically, as shown by the decreasing size of the steps in Fig. 5B as the intersection is approached.

To obtain near-optimal performance, the transformation line must intersect the reward rate curve at or near its apex. Further on, we suggest that this line can be chosen to approximate the relationship between threshold and optimal reward rate over a range of task parameters (see Fig. 6), and that this line itself may be subject to adaptation over longer time scales.

6.2. Continuous time system

We define the continuous time system as a set of SDEs augmented by conditions for threshold crossing and decision

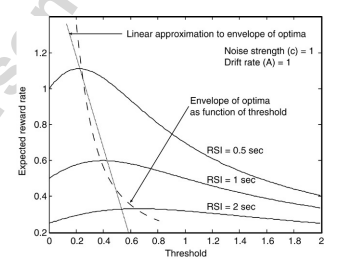


Fig. 6. The affine function that translates reward rate estimates into threshold produces the best performance when it approximates the envelope of optima (dashed curve), because it intersects the expected reward rate curves near their peaks for a range of RSI conditions. Different ranges of RSI conditions require different affine approximations, and the function is assumed to adapt slowly in response to RSI conditions experienced over a longer time scale.

variable resetting:

$$dx = \begin{cases} Adt + c_1 dW & \text{if RSI}(t) = 0\\ 0 & \text{if RSI}(t) = 1 \end{cases}$$
 (13)

$$x = 0 \text{ if RSI}(t) = 1 \tag{14}$$

$$dr = \frac{1}{k} \cdot (-r(t) + R(t))dt + c_2 dW$$
 (15)

$$z(t) = \max(0, z_{\text{max}} - w \cdot r(t)) \tag{16}$$

$$R(t) = \begin{cases} \delta(t - t') & \text{if reward present at time } t' \\ 0 & \text{otherwise} \end{cases}$$
 (17)

$$\tau$$
 = time of last threshold crossing (18)

$$RSI(t) = \begin{cases} 1 & \text{if } |x| \ge z \text{ or } t - \tau < RSI_{\text{max}} \\ 0 & \text{otherwise.} \end{cases}$$
 (19)

Here x (Eqs. (13) and (14)) is the decision variable in the DDM, r in Eq. (15) is the running estimate of reward rate, and z in Eq. (16) is the threshold. Eqs. (13), (14) and (19)model the effects of RSI (which impacts reward rate) as well as the assumption that the decision variable starts at the origin on each new trial.³ Specifically, when a stimulus is present, the RSI variable is set to 0 (Eq. (19)) and first-passage of the decision variable x beyond either threshold $\pm z$ is taken as the time of decision, τ (Eq. (18)). At this point, Eq. (19) specifies the response–stimulus interval RSI_{max} for the next trial, and Eq. (13) resets the decision variable to 0. For correct responses, Eqs. (15) and (17) apply a Dirac-delta impulse to the reward rate estimator, which otherwise decays exponentially (see Fig. 4). The threshold z is determined entirely by the reward rate estimate r via the affine function of Eq. (16). In Section 6.4 we show that the continuous system (Eqs. (13)–(19)) shares the property of rapid convergence to near-optimal thresholds of its discrete time version.

6.3. Robustness, generality and parsimony

Given the affine function $z=z_{\rm max}-wr$ relating threshold to reward rate, the continuous system (Eqs. (13)–(19)) accomplishes gradient ascent without explicit memory, reward-rate comparison or step-size reduction mechanisms. It is also robust to estimation errors and noise, since the intersection point of Fig. 5 is a global attractor. Furthermore, it can be used to adapt the threshold to its optimal value across a range of task parameters. As noted earlier, behavioral evidence suggests that human participants, performing well-practiced tasks, are capable of adaptation over as few as 5–10 trials following a change in task conditions (Bogacz et al., in press; Ratcliff et al., 1999).

The method's performance depends on the slope w and intercept $z_{\rm max}$ of the affine function. This function defines a parsimonious linear approximation of the relationship between the optimal threshold and reward rate across a range of task conditions, as in Fig. 2, provided that the resulting line passes close to the reward rate maxima indicated by the dashed curve of Fig. 6.⁴ We shall assume that these parameters have been learned, for a given task, through practice under different trial conditions (e.g., RSI or noise level). In Section 6.5 we indicate how this can be done by reinforcement learning. First we demonstrate that, like the discrete time version of the algorithm described in Section 6.1, the continuous system (Eqs. (13)–(19)) rapidly converges to near-optimal thresholds.

6.4. Simulations

Numerical simulations of the continuous-time algorithm demonstrate its effect on the behavioral variables of interest: response time and accuracy. According to an optimal analysis of the DDM in this task, thresholds should be set lower for faster RSI conditions, producing faster mean RT and higher error rates. Fig. 7 shows representative timecourses of the decision process variables for one parameterization of the abstract diffusion model in Eqs. (13)–(19), along with the optimal thresholds and associated reward rates before and after a task condition switch. Note how the system updates from arbitrary initial conditions and again following the switch.

The right panel of Fig. 8 shows RT distributions and error rates for the model of Eqs. (13)–(19), illustrating a speed–accuracy tradeoff similar to tradeoffs seen in human behavioral data (left panel, from a pilot study we conducted). In each panel, median and interquartile RTs are shown on the left and error rates on the right. These demonstrate significantly faster RTs for shorter RSIs (Wilcoxon rank-sum test, p < 0.01, each pairwise comparison), and also significantly higher error rates for shorter RSIs (pairwise t-test, p < 0.001).

Finally, Fig. 9 shows the neural implementation of the abstract model (in which a single decision variable and threshold are decomposed into two accumulators and two threshold units), and Fig. 10 shows activation timecourses of its units in response to changing RSI conditions. As pointed out in Section 4, the slope w of the affine function (Fig. 3B) corresponds to the connection weight from the reward rate estimator to the thresholding units, and the intercept $z_{\rm max}$ can be interpreted as the bias (γ) of the sigmoid (Eq. (5)), or as an additional input.

6.5. Learning the critical parameters

The model we have described thus far assumes a linear approximation of the relationship between optimal threshold and reward rate. However, the best approximation differs considerably across different ranges of task parameters (see Fig. 6). Here we consider the possibility that this approximation – the affine function used to adapt the threshold – can itself be adapted through learning. To achieve this, reward rates must be experienced for different values of task parameters, such as RSI and stimulus discriminability.

Noting that the reward rate of Eq. (10) depends on the signal-to-noise ratio A/c, threshold-to-signal ratio z/A, and RSI in a complex, nonlinear manner, it is clear that the slope and intercept parameters w and $z_{\rm max}$ represent a substantial compression of information. However, this is useful only insofar as the affine function $z=z_{\rm max}-wr$ (shown as the straight lines in Figs. 5 and 6) reasonably approximates the envelope of reward rate optima (shown as the dashed curves in Figs. 2 and 6) over a given range of task conditions. For example, the best linear approximation for 500 ms-1 s RSIs would significantly differ from the one for 1-2 s values (see Fig. 6). Since linearizations may provide good approximations only over limited ranges of task parameters, it seems reasonable

³ This last, rather unrealistic assumption implies a discontinuous decision variable trajectory over time, but it can be relaxed without loss of generality by introducing a refractory period and modeling the system as a stable, rapidly decaying Ornstein–Uhlenbeck process during the RSI, as done in Fig. 7; we use the simpler system here for ease of discussion.

⁴ Ideally one would use the envelope of optima – the dashed curve – to transform reward rate estimates into thresholds, but we are constrained to linearity by the plausible neural mechanism assumed in Section 4: superposition of synaptic inputs to the threshold units.

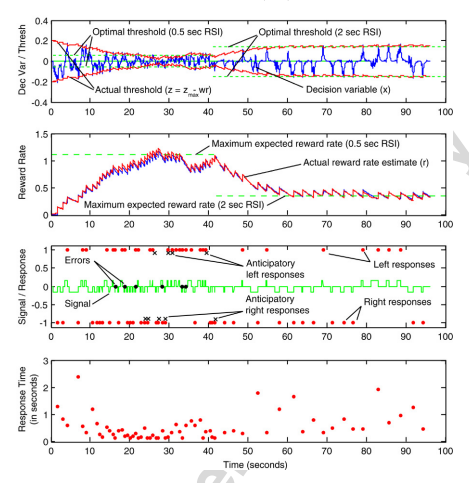


Fig. 7. Timecourses of variables in a sample run of the algorithm. The DDM (Eq. (1)) is simulated directly in the top timecourse, where the decision variable traces out a trajectory within bounds formed by the dynamic threshold (each point of contact between these represents a decision). Optimum thresholds for the two RSI conditions simulated (0.5 s RSI, until about $t \approx 42$ s; 2 s RSI, from 42 s to end) are shown as dashed lines. The second timecourse shows a leaky integrator estimate of the reward rate (Eq. (11)), superimposed on an estimate with added Gaussian noise, plus dashed lines indicating the expected reward rate for the optimum threshold in each RSI condition. The third plot shows the signal direction (upward square pulse = left; zero = RSI; downward square pulse = right); responses are dots plotted at height 1 for left and -1 for right; dots at height 0 are premature responses occurring due to low threshold, and \times 's denote errors. The fourth plot shows RTs.

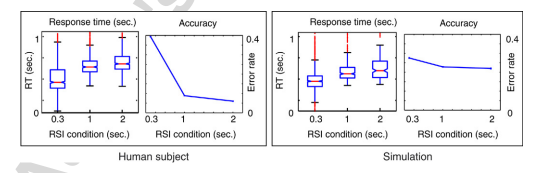


Fig. 8. Speed–accuracy tradeoffs produced by a human subject (left panel) and by the algorithm (right panel). The horizontal axis in all plots represents the RSI condition for blocks of trials (0.3, 1 or 2 s). The left plot in each panel shows a boxplot of RT, with median RT represented by the middle notch, interquartile RT range (25th–75th percentile) denoted by height of box, and outliers denoted by whiskers. The right plots show error rates.

to assume that the approximation given by the affine function may be tuned to accommodate different task environments. This can be accomplished by generic reinforcement learning.

Our arguments supporting fast threshold convergence assume an affine function that is fixed or that changes

slowly in comparison to reward rate estimates. The affine function effectively exploits knowledge about the shape of the reward rate curves, trading off generality for speed and task-specificity. Learning the transformation requires exploration of a larger space of possibilities. We therefore propose that such

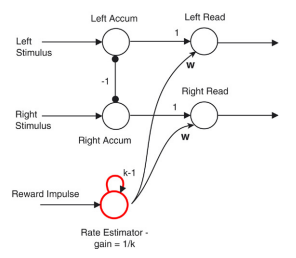


Fig. 9. A five-unit neural network that implements the decision mechanism. Mutually inhibiting accumulator units in the first layer approximate a DDM, units with high gain and bias in the second layer detect threshold crossings, and a reward rate estimator with balanced feedback and gain modulates the detector thresholds according to $z = z_{\text{max}} - wr$.

'environmental models' are learned on a slow time scale, and then used as described above to make rapid improvements to task performance in 'sub-environments'.

As an alternative to adapting the linear approximation, reinforcement learning could be applied directly to threshold adjustment itself. That is, reinforcement learning could be used to produce a full representation of the multi-parameter family of non-linear relationships between thresholds and reward rates. Nevertheless, as with simple hill-climbing, general reinforcement learning methods would still require annealing or step-size reduction schedules tuned to these relationships (cf. Section 3) in order to adapt thresholds quickly to a train of incoming rewards, while simultaneously settling securely on good parameter values in the face of noise (this is another instance of a stability/plasticity tradeoff; Grossberg (1987)). Separating the time scales allows us to obtain the best of both worlds in this task by applying the threshold update algorithm over a time scale of 5–10 trials while simultaneously using reinforcement learning in the background to adjust slope and intercept parameters over a time scale spanning multiple blocks of trials.

In work referred to in Simen et al. (2005) and to be described in a future publication, we have shown that parameters such as slope and intercept can be learned by continuous-time versions of temporal difference algorithms such as the actor–critic method (cf. Doya (2000)). The proposed method draws on the stochastic real-valued unit algorithm of Gullapalli (1990), and is closely related to the Alopex algorithm of Harth and Tzanakou (1974) and a general class of stochastic optimization algorithms (see Kushner and Yin (1997)). Specifically, connection weights from the reward rate estimator to the threshold units of Fig. 9, which correspond to slope, can be updated as in Eq. (15) of Section 6.2, but at a much slower rate (larger k). To obtain the correct hill-climbing behavior, the input (-r(t) + R(t)) in Eq. (15) can be replaced by a product of the *derivatives* of longer-term averages

of reward rate and of the weight itself. Thus, slopes are adjusted only when reward rates are consistently rising or falling, as they would, for example, following a significant change of RSI range. We show that chains of first-order units can approximate derivatives, and also suggest a biologically-plausible neural substrate that employs dopamine modulation of glutamatergic synapses.

7. Discussion

We have demonstrated that very simple neural mechanisms can give rise to speed–accuracy tradeoffs that maximize reward rate and that are similar to the tradeoffs exhibited by human subjects. Our approach exploits attractor dynamics to deal with the challenges faced by straightforward reinforcement learning and gradient ascent approaches to the threshold adaptation problem, as well as more abstract algorithms (e.g., Busemeyer and Myung (1992) and Erev and Barron (2005)). The model we describe operates in continuous time, is able to reliably estimate the reward rate, rapidly and stably converges on a threshold that maximizes the reward rate to a reasonable approximation, and does not require any additional apparatus for explicitly remembering or comparing previous estimates of reward rate with current estimates.

A critical feature of the model is that it relies on a linear approximation of the relationship between optimal threshold and reward rate across a range of task conditions. This serves two purposes. First, it provides a representation of this relationship that is considerably simpler the actual curvilinear relationship (shown in Fig. 2) which is itself the solution to a transcendental equation that can only be approximated numerically (Bogacz et al., in press). Second, and perhaps more importantly, it allows the neural implementation of a threshold adaptation algorithm that uses an affine function to achieve rapid convergence to a threshold in close proximity to the optimal one. However, this use of a linear approximation also represents a potential limitation of the model, insofar as the specific best approximation varies as a function of task parameters (see Fig. 6). We have suggested how the parameters defining the linear approximation could themselves be adapted using simple principles of reinforcement learning, although this adaptation would need to occur on a significantly longer time scale than that of threshold adaptation. The effects of these simultaneous adaptations at different times scales, and their relationship to observations about human performance, remain to be explored in future theoretical and empirical studies.

Previous non-neural modeling work has approached the issue of speed–accuracy tradeoffs from the perspective that subjects try to minimize a cost consisting of a linear combination of speed and accuracy, subject to information processing constraints (e.g., Maddox and Bohil (1998) and Mozer, Colagrosso, and Huber (2002)). This approach seems entirely consistent with an extended version of our model that also maintains an estimate of error rate in the same manner as it estimates reward rate. What is new here is the implementation of these processes using neurally plausible mechanisms and a dynamical systems analysis of how and why they work. Reward

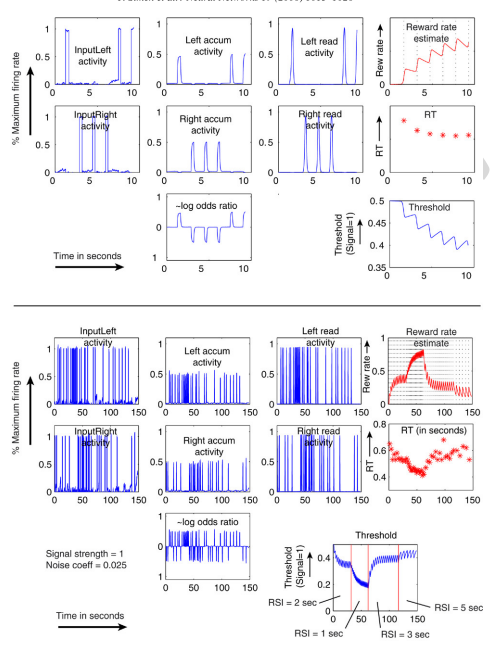


Fig. 10. Activations of all units in the network of Fig. 9 are plotted, the position of each plot corresponding to the position of each unit in the circuit diagram. The top set of panels shows a short timecourse, and the bottom set a longer one that illustrates fast threshold adaptation. The first column shows inputs to left and right channels. The second column plots left and right accumulator activations and the log odds ratio of evidence derived from their difference. The third column shows left and right threshold detectors. The top plot in the fourth column shows the reward rate monitor, with dotted vertical lines marking stimulus onsets, showing that RSI was decreased from 2 to 1 s at about 30 s, and from 1 to 3 s at about 60 s. Response time (RT) is plotted below as asterisks. Note the similarity between the rate monitor activity and the plots of threshold at bottom right.

rate estimators, by directly exciting threshold readout units, can implement a speed–accuracy tradeoff that is nearly optimal across a range of task conditions.

In this respect, our approach is similar to an existing neural model in which continuous feedback in the form of motivational signals influences underlying decision making circuits based on lateral inhibition ('gated dipoles') in order to change their stimulus sensitivity (Grossberg, 1982). To the best of our knowledge, however, that type of model has not been applied specifically to the problem of estimating and responding to the rate of reward in a free response task.

It is also worth noting the similarity of the reward rate estimate discussed in Section 5 to an 'accumulating' eligibility trace as used in reinforcement learning (Sutton & Barto, 1998). The two quantities are computed in the same way, but the purpose of the eligibility trace is typically for changing weights in reinforcement learning, whereas in our case we simply feed the equivalent quantity into the response units as an additional input. In this way, our approach corresponds to an activation-based mechanism, whereas the traditional use of an eligibility trace (as applied to brain modeling) is for governing synaptic plasticity. Combining both approaches should provide

a means for optimizing performance at both short and long time scales.

7.1. Implementation in the brain

Recent findings suggest that the components of our model may reflect the operation of specific neural structures. Previous reports have suggested that areas of LIP as well as the frontal eye field may correspond to the accumulators in the DDM (e.g., Gold and Shadlen (2001), Hanes and Schall (1996) and Shadlen and Newsome (2001)). Other findings have begun to identify neural mechanisms that may be involved in reward rate estimation and the adaptation of decision parameters. For example, given the role of orbitofrontal cortex (OFC) in encoding the reward value of objects or actions (e.g. Rolls (2000)) and evidence that it may also encode the *rate* of reward (Sugrue, Corrado, & Newsome, 2004b), it is possible that the reward rate estimator unit in Fig. 9 may serve as a simple, first-order approximation to OFC function in the type of decision making task discussed here.

Further, activity in areas of monkey parietal cortex thought to be involved in sensorimotor transformation has been shown to be sensitive to expected future reward (Platt & Glimcher, 1999) and the relative values of competing choices (Sugrue et al., 2004a) in oculomotor tasks. Similar effects have been observed in other brain areas thought to be involved in the decision-making process, such as the dorsolateral prefrontal cortex (Barraclough, Conroy, & Lee, 2004; Leon & Shadlen, 1999). Thus it is possible that connections from OFC to threshold detectors implemented in frontal and/or parietal cortex may implement something like our threshold adaptation algorithm.

Given their role in reward processing, the basal ganglia are also promising candidates for involvement in threshold modulation. Along these lines, Frank (2006) discusses a pattern of anatomic connections and a possible role for the subthalamic nucleus (STN) that are consistent with the mechanisms we have described. There, the proposed role of the STN is effectively to increase the threshold when increased conflict is detected between competing responses in a Go/No-Go task. In this way, emphasis shifts toward accuracy and away from speed when response conflict is high. A shift in the opposite direction in response to reward rate might similarly be achieved by connections from OFC to the striatum, given the role that the striatum plays in promoting, rather than inhibiting, the propagation of activity through the basal ganglia (as the STN does).

Our model assumes that speed-accuracy tradeoffs are implemented in the brain at a single stage of processing (the threshold-crossing detection stage) through modulation by an estimate of reward rate. However, neither reward rate estimation nor threshold-crossing detection are functions that are likely to be discretely localized in the brain. As we have noted above, many areas of the brain are sensitive to information about reward. Furthermore, the specific neural mechanisms responsible for information accumulation and threshold adjustment are likely to vary based on the demands of a given

task (e.g., whether it involves visual or auditory information, and a manual or oculomotor response). Presumably these are selected by frontal control mechanisms for task engagement (e.g., Miller and Cohen (2001)). We propose, however, that our model identifies fundamental principles of operation that may be shared in common by the neural mechanisms involved in decision making across different processing domains. Along these lines, it will be important to explore the relationship between these mechanisms and others that have been proposed for the adaptive regulation of performance. This includes the use of processing conflict to adapt threshold as well as attentional variables (e.g. Botvinick et al. (2001)), as well as the role of neuromodulatory mechanisms in adapting processing parameters such as threshold and gain (e.g., Aston-Jones and Cohen (2005)).

7.2. Extensions to a broader range of decision tasks

Our model has focused exclusively on free responding in the TAFC task. However, the basic ideas should extend to address a broader range of more realistic tasks. For example, it should be straightforward to incorporate our reward rate estimation and threshold adaptation mechanisms into models for free responding in multi-alternative decision tasks (e.g., Bogacz and Gurney (in press), McMillen and Holmes (2006) and Usher and McClelland (2001)), as well as Go/No-Go tasks as discussed above. It should also be possible to address other response conditions, such as deadlining, by elaborating the model to maintain an estimate of late-response rate, and to use this in place of reward rate estimates to set an internal deadline for responding. Finally, decision-making tasks that also involve working memory over a delay period can be modeled by implementing thresholds via detector units with strong, selfexcitatory positive feedback - as in the firing rate models of Nakahara and Doya (1998) and Simen (2004) and the spiking models of Seung, Lee, Reis, and Tank (2000) and Wang (2002) - rather than non-self-exciting units with very steep gain in their activation functions (as in Section 4). The analysis presented here in terms of units with activation functions that are step functions carries over without loss of generality to such selfexciting units (Simen, 2004).

We hope that this model helps extend the foundation that has begun to develop for formalizing decision-making tasks, particularly those that integrate continuous time reward monitoring and prediction with sensory mechanisms, and that it may serve as a bridge between psychological function and neural implementation.

Acknowledgments

This work was supported by PHS grants MH58480 and MH62196 (Cognitive and Neural Mechanisms of Conflict and Control, Silvio M. Conte Center) (PS, JDC and PH) and DoE grant DE-FG02-95ER25238 (PH).

References

Aston-Jones, G., & Cohen, J. D. (2005). An integrative theory of locus coeruleus-morepinephrine function: adaptive gain and optimal performance. *Annual Review of Neuroscience*, 28, 403–450.

- Barraclough, D. J., Conroy, M. L., & Lee, D. (2004). Prefrontal cortex and decision making in a mixed-strategy game. *Nature Neuroscience*, 7, 404–410.
- Bogacz, R., Brown, E., Moehlis, E., Holmes, P., & Cohen, J.D. (2006). The physics of optimal decision making: a formal analysis of models of performance in two-alternative forced choice tasks. *Psychological Review* (in press).
- Bogacz, R., & Gurney, K. (2006). The basal ganglia and cortex implement optimal decision making between alternative actions. *Neural Computation* (in press).
- Botvinick, M. M., Braver, T. S., Barch, D. M., Carter, C. C., & Cohen, J. D. (2001). Conflict monitoring and cognitive control. *Psychological Review*, 108(3), 624–652.
- Brody, C. D., Hernandez, A., Zainos, A., & Romo, R. (2003). Timing and neural encoding of somatosensory parametric working memory in macaque prefrontal cortex. *Cerebral Cortex*, 13, 1196–1207.
- Brown, E., Gao, J., Holmes, P., Bogacz, R., Gilzenrat, M., & Cohen, J. D. (2005). Simple neural networks that optimize decisions. *International Journal of Bifurcation and Chaos*, 15(3), 803–826.
- Busemeyer, J. R., & Myung, I. J. (1992). An adaptive approach to human decision making: learning theory and human performance. *Journal of Experimental Psychology: General*, 121, 177–194.
- Busemeyer, J. R., & Townsend, J. T. (1993). Decision field theory: A dynamic-cognitive approach to decision making in uncertain environments. *Psychological Review*, 100, 432–459.
- Cohen, J. D., Dunbar, K., & McClelland, J. L. (1990). On the control of automatic processes: A parallel distributed processing model of the Stroop effect. *Psychological Review*, 97(3), 332–361.
- Cohen, M., & Grossberg, S. (1983). Absolute stability of global pattern formation and parallel memory storage by competitive neural networks. *IEEE Transactions on Systems, Man and Cybernetics*, 13, 815–826.
- Doya, K. (2000). Reinforcement learning in continuous time and space. *Neural Computation*, 12, 219–245.
- Erev, I. (1998). Signal detection by human observers: a cut-off reinforcement learning model of categorization decisions under uncertainty. *Psychological Review*, 105, 280–298.
- Erev, I., & Barron, G. (2005). On adaptation, maximization, and reinforcement learning among cognitive strategies. *Psychological Review*, 112, 912–931.
- Frank, M. (2006). Hold your horses: a dynamic computational role for the subthalamic nucleus in decision making. *Neural Networks* (this issue).
- Freeman, W. J. (1979). Nonlinear gain mediating cortical stimulus-response relations. *Biological Cybernetics*, 33, 237–247.
- Gardiner, C. W. (1985). *Handbook of stochastic methods* (2nd ed.). New York: Springer.
- Gold, J. I., & Shadlen, M. N. (2000). Representation of a perceptual decision in developing oculomotor commands. *Nature*, 404, 390–394.
- Gold, J. I., & Shadlen, M. N. (2001). Neural computations that underlie decisions about sensory stimuli. *Trends in Cognitive Sciences*, 5, 10–16.
- Gold, J. I., & Shadlen, M. N. (2002). Banburismus and the brain: Decoding the relationship between sensory stimuli, decisions and reward. *Neuron*, 36, 299–308
- Grossberg, S. (1982). A psychophysiological theory of reinforcement, drive, motivation and attention. *Journal of Theoretical Neurobiology*, 1, 286–369.
- Grossberg, S. (1987). Competitive learning: from interactive activation to adaptive resonance. *Cognitive Science*, 11, 23–63.
- Grossberg, S., & Gutowski, W. (1987). Neural dynamics of decision making under risk: affective balance and cognitive–emotional interactions. *Psychological Review*, 94(3), 300–318.
- Gullapalli, V. (1990). A stochastic reinforcement learning algorithm for learning real-valued functions. *Neural Networks*, 3, 671–692.
- Hanes, D. P., & Schall, J. D. (1996). Neural control of voluntary movement initiation. Science, 274, 427–430.
- Harth, E., & Tzanakou, E. (1974). Alopex: A stochastic method for determining visual receptive fields. Vision Research, 14, 1475–1482.
- Holmes, P., Brown, E., Moehlis, J., Bogacz, R., Gao, J., Aston-Jones, G., et al. (2005). Optimal decisions: From neural spikes, through stochastic differential equations, to behavior. *IEICE Transactions on Fundamentals on Electronics, Communications and Computer Sciences*, 88(10), 2496–2503.

- Jordan, D. W., & Smith, P. (1999). Nonlinear ordinary differential equations (3rd ed.). New York, NY: Oxford University Press.
- Kushner, H. J., & Yin, G. G. (1997). Stoachastic approximation algorithms and applications. New York: Springer.
- Laming, D. R. J. (1968). Information theory of choice reaction time. New York: Wiley.
- Leon, M. I., & Shadlen, M. N. (1999). Effect of expected reward magnitude on the response of neurons in the dorsolateral prefrontal cortex of the macaque. *Neuron*, 24, 415–425.
- Luce, R. D. (1986). Response times: their role in inferring elementary mental organization. New York: Oxford University Press.
- Maddox, W. T., & Bohil, C. J. (1998). Base-rate and payoff effects in multidimensional perceptual categorization. *Journal of Experimental Psychology: Learning, Memory and Cognition*, 24, 1459–1482.
- McCulloch, W., & Pitts, W. (1943). A logical calculus of the ideas immanent in nervous activity. *Bulletin of Mathematical Biophysics*, 5, 115–133.
- McMillen, T., & Holmes, P. (2006). The dynamics of choice among multiple alternatives. *Journal of Mathematical Psychology*, 50, 30–57.
- Miller, E. K., & Cohen, J. D. (2001). Integrative theory of PFC function. *Annual Review of Neuroscience*, 24, 167–202.
- Mozer, M. C., Colagrosso, M. D., & Huber, D. H. (2002). A rational analysis of cognitive control in a speeded discrimination task. In T. Dietterich, S. Becker, & Z. Ghahramani (Eds.), Advances in neural information processing systems: Vol. 14 (pp. 51–57). Cambridge, MA: MIT Press.
- Myung, I. J., & Busemeyer, J. R. (1989). Criterion learning in a deferred decision making task. American Journal of Psychology, 102, 1–16.
- Nakahara, H., & Doya, K. (1998). Near-saddle-node bifurcation behavior as dynamics in working memory for goal-directed behavior. *Neural Computation*, 10, 113–132.
- Oppenheim, A. V., & Willsky, A. S. (1996). Signals and systems (2nd ed.). New York: Prentice Hall.
- Palmer, J., Huk, A. C., & Shadlen, M. N. (2005). The effect of stimulus strength on the speed and accuracy of a perceptual decision. *Journal of Vision*, 5(5), 376–404.
- Platt, M. L., & Glimcher, P. W. (1999). Neural correlates of decision variables in parietal cortex. *Nature*, 400, 233–238.
- Ratcliff, R. (1978). A theory of memory retrieval. *Psychological Review*, 85, 59–108
- Ratcliff, R., & Rouder, J. N. (1998). Modeling response times for two-choice decisions. *Psychological Science*, 9, 347–356.
- Ratcliff, R., Van Zandt, T., & McKoon, G. (1999). Connectionist and diffusion models of reaction time. *Psychological Review*, 106, 261–300.
- Roitman, J. D., & Shadlen, M. N. (2002). Response of neurons in the lateral intraparietal area during a combined visual discrimination reaction time task. *Journal of Neuroscience*, 22, 9475–9489.
- Rolls, E. T. (2000). The orbitofrontal cortex and reward. *Cerebral Cortex*, 10(3), 284–294.
- Seung, H. S., Lee, D. D., Reis, B. Y., & Tank, D. W. (2000). The autapse: a simple illustration of short-term analog memory storage by tuned synaptic feedback. *Journal of Computational Neuroscience*, 9, 171–185.
- Shadlen, M., & Newsome, W. (2001). Neural basis of a perceptual decision the parietal cortex (area LIP) of the rhesus monkey. *Journal of Neurophysiology*, 86, 1916–1936.
- Simen, P. (2004). Neural mechanims for control in complex cognition. Ph.D. thesis. University of Michigan.
- Simen, P., Holmes, P., & Cohen, J. D. (2005). Threshold adaptation in decision making. Society for Neuroscience Abstracts.
- Smith, P. L., & Ratcliff, R. (2004). Psychology and neurobiology of simple decisions. *Trends in Neuroscience*, 27, 161–168.
- Stone, M. (1960). Models for choice reaction time. *Psychometrika*, 25, 251–260.
- Sugrue, L. P., Corrado, G. S., & Newsome, W. T. (2004a). Matching behavior and the representation of value in the parietal cortex. *Science*, 304, 1782–1787.

- Sugrue, L. P., Corrado, G. S., & Newsome, W. T. (2004b). Neural correlates of value in orbitofrontal cortex of the rhesus monkey. Society for Neuroscience Abstracts
- Sutton, R. S., & Barto, A. G. (1998). *Reinforcement learning*. Cambridge, MA: MIT Press.
- Usher, M., & McClelland, J. (2001). On the time course of perceptual
- choice: the leaky competing accumulator model. *Psychological Review*, 108, 550-592.
- Wald, A., & Wolfowitz, J. (1948). Optimum character of the sequential probability ratio test. *Annals of Mathematical Statistics*, 19, 326–339.
- Wang, X. -J. (2002). Probabilistic decision making by slow reverberation in cortical circuits. *Neuron*, *36*, 955–968.

