

## Should I stay or should I go? How the human brain manages the trade-off between exploitation and exploration

Jonathan D Cohen, Samuel M McClure and Angela J Yu

*Phil. Trans. R. Soc. B* 2007 **362**, 933-942

doi: 10.1098/rstb.2007.2098

---

### References

[This article cites 41 articles, 9 of which can be accessed free](#)

<http://rstb.royalsocietypublishing.org/content/362/1481/933.full.html#ref-list-1>

[Article cited in:](#)

<http://rstb.royalsocietypublishing.org/content/362/1481/933.full.html#related-urls>

### Rapid response

<http://rstb.royalsocietypublishing.org/content/362/1481/933/reply>

[Respond to this article](#)

<http://rstb.royalsocietypublishing.org/letters/submit/royptb;362/1481/933>

### Email alerting service

Receive free email alerts when new articles cite this article - sign up in the box at the top right-hand corner of the article or click [here](#)

# Should I stay or should I go? How the human brain manages the trade-off between exploitation and exploration

Jonathan D. Cohen<sup>1,2,\*</sup>, Samuel M. McClure<sup>1</sup> and Angela J. Yu<sup>1</sup>

<sup>1</sup>Department of Psychology and Center for the Study of Brain, Mind and Behaviour, Princeton University, Princeton, NJ 08540, USA

<sup>2</sup>Department of Psychiatry, University of Pittsburgh, Pittsburgh, PA 15213, USA

Many large and small decisions we make in our daily lives—which ice cream to choose, what research projects to pursue, which partner to marry—require an exploration of alternatives before committing to and exploiting the benefits of a particular choice. Furthermore, many decisions require re-evaluation, and further exploration of alternatives, in the face of changing needs or circumstances. That is, often our decisions depend on a higher level choice: whether to exploit well known but possibly suboptimal alternatives or to explore risky but potentially more profitable ones. How adaptive agents choose between exploitation and exploration remains an important and open question that has received relatively limited attention in the behavioural and brain sciences. The choice could depend on a number of factors, including the familiarity of the environment, how quickly the environment is likely to change and the relative value of exploiting known sources of reward versus the cost of reducing uncertainty through exploration. There is no known generally optimal solution to the exploration versus exploitation problem, and a solution to the general case may indeed not be possible. However, there have been formal analyses of the optimal policy under constrained circumstances. There have also been specific suggestions of how humans and animals may respond to this problem under particular experimental conditions as well as proposals about the brain mechanisms involved. Here, we provide a brief review of this work, discuss how exploration and exploitation may be mediated in the brain and highlight some promising future directions for research.

**Keywords:** exploration; uncertainty; learning; neurotransmitters; prefrontal cortex; decision making

## 1. INTRODUCTION

Should I stay or should I go now?  
If I go there will be trouble  
And if I stay it may be double  
So come on and let me know  
Should I stay or should I go?

(The Clash)

Every researcher has personal experience with the exploration–exploitation dilemma. At some point in the conduct of a study, when the data are still inconclusive, it may become necessary to decide how to proceed. On the one hand, there is the option to continue with the experiment, in the hope that with more effort and data, the results will look more promising. Alternatively, the experiment can be scrapped in favour of a modified experimental design, a new approach to the problem, or an entirely new research topic. That is, the experimenter faces a trade-off between the value of exploitation versus exploration. This example highlights the importance of this problem in decision making, one that has typically been ignored in psychological research on cognitive control and executive function.

The need to balance exploitation with exploration is confronted at all levels of behaviour and time-scales of decision making from deciding what to do next in the day to planning a career path. It is confronted by individuals in love (as captured by the lyrics above) and by entire armies at war (should a campaign focus intensively on one battle or seek to identify new opportunities to surmount the enemy). Nor is it limited to human behaviour. It is confronted by fungi deciding whether to concentrate growth at a local site or send out hyphae to sample more distant resources (Watkinson *et al.* 2005); by ant colonies exploring options for a new nest before settling on and exploiting a particular site (Pratt & Sumpter 2006); by engineers generating algorithms to deploy a fleet of automata to map the expanses of a new environment (Leonard *et al.* in press) and by machine learning theorists—who coined the phrase ‘exploration versus exploitation’—in their efforts to improve the ability of reinforcement learning (RL) algorithms to function adaptively in changing environments (e.g. Kaelbling *et al.* 1996).

In general, how agents should and do respond to the trade-off between exploration and exploitation is poorly understood. In part, this reflects the difficulty of the problem: there is no known optimal policy for trading off exploration and exploitation in general, even when the objectives are well specified. Gittins & Jones (1974) and Gittins (1979) presented a strategy and proved its

\* Author for correspondence (jdc@princeton.edu).

One contribution of 14 to a Discussion Meeting Issue ‘Mental processes in the human brain’.

optimality for a limited class of problems in which the decisions are made from a finite number of stationary bandit processes (e.g. options for which the reward is delivered with *unknown* but *fixed* probabilities), and when the agent discounts their value exponentially over time. Gittins proved that if being optimal consists of maximizing the cumulative reward over an infinite horizon when the value of each reward is discounted exponentially as a function of when it is acquired, then the optimal policy is to calculate the expected total future rewards associated with each option at a particular time—a value known as the Gittins index—and to select that bandit with the greatest Gittins index (Gittins & Jones 1974; Gittins 1979). The significance of Gittins' contribution is that it reduced the decision problem to computing and comparing these scalar indices. In practice, computing the Gittins index is not tractable for many problems for which it is known to be optimal. However, for some limited problems, explicit solutions have been found. For instance, the Gittins index has been computed for certain two-armed bandit problems (in which the agent chooses between two options with independent probabilities of generating a reward), and compared to the foraging behaviour of birds under comparable circumstances; the birds were found to behave approximately optimally (Krebs *et al.* 1978).

While the Gittins index lends formal rigour to the problem of exploration versus exploitation, proof of its optimality requires strong assumptions about the environment and the agent. The properties of the individual bandits must be frozen unless acted upon (i.e. the pay-off structure of the environment must be stationary), all options must be available at all decision points (i.e. there cannot be any 'side paths') and agents must discount the value of rewards exponentially into the future (Gittins 1979; Berry & Fristedt 1985; Banks & Sundaram 1994). Real-world problems typically violate one or more of these assumptions.

Perhaps, the most important exception to Gittins' assumptions is that real-world environments are typically non-stationary; i.e. they change with time. To understand how organisms manage the balance between exploration and exploitation in non-stationary environments, investigators have begun to study how organisms adapt their behaviour in response to the experimentally induced changes in reward contingencies. Several studies have now shown that both humans and other animals dynamically update their estimates of rewards associated with specific courses of action, and abandon actions that are deemed to be diminishing in value in search of others that may be more rewarding (e.g. Sugrue *et al.* 2004; Daw *et al.* 2006; Gilzenrat & Cohen in preparation). At the same time, there is also longstanding evidence that humans sometimes exhibit an opposing tendency. When reward diminishes (e.g. following an error in performance), subjects often try harder at what they have been doing rather than less (e.g. Rabbitt 1966; Laming 1979; Gratton *et al.* 1992). The balance between exploration and exploitation also seems to be sensitive to time horizons. Humans show a greater tendency to explore when there is more time left in a task, presumably because this allows

them sufficient time later to enjoy the fruits of those explorations (Carstensen *et al.* 1999). A full account of how people regulate the balance between exploration and exploitation must account for these diverse, and in some cases seemingly discrepant, patterns of behaviour.

Recent findings are also beginning to shed light on the neural mechanisms that underlie exploratory and exploitative behaviours. These findings consistently implicate the involvement of neuromodulatory systems thought to be involved in assessing reward and uncertainty. The midbrain dopamine system has been implicated in the signalling of reward prediction errors critical for learning the value of specific actions (Montague *et al.* 1996; Schultz *et al.* 1997) and for decision-making based on those values (McClure *et al.* 2003). The locus coeruleus (LC) noradrenergic system has been proposed to govern the balance between exploration and exploitation in response to reward history (Aston-Jones & Cohen 2005). And the basal forebrain cholinergic system together with the adrenergic system have been proposed to monitor uncertainty, signalling both expected and unexpected forms, respectively, which in turn might be used to promote exploitation or exploration (Yu & Dayan 2005).

Regulating the balance between exploitation and exploration is a fundamental need for adaptive behaviour in a complex and changing world. In the rest of this article, we consider the progress outlined above that has been made in understanding this problem in formal terms and in identifying the mechanisms that have evolved in natural organisms for meeting this challenge. While there has been recent progress in identifying relevant empirical phenomena and candidate neural mechanisms, such work is still in the earliest stages. Accordingly, the connection between theory and data remains largely speculative. Our primary purpose here is to call attention to the problem and point to relevant lines of research that show promise in addressing it.

## 2. OPTIMAL PERFORMANCE IN STATIONARY ENVIRONMENTS: THE GITTINS INDEX

In a landmark paper, Gittins & Jones (1974) developed a straightforward means for calculating the optimal strategy for decision making in multi-armed bandit problems. Bandit problems are well suited for studying the tension between exploitation and exploration since they offer a direct trade-off between exploiting a known source of reward (continuing to play one arm of the bandit) and exploring the environment (trying other arms) to acquire information about other sources of reward (Kaelbling 1996).

For an  $n$ -armed bandit problem, an agent is required to choose between  $n$  options, each of which delivers reward with a probability  $p_i$ . The probability of obtaining reward from a bandit,  $p_i$ , may change through time but only when a choice is made for that bandit. The goal for the agent is to maximize expected rewards,  $V_i$ , where rewards earned in the future are discounted by an exponential discount factor  $\delta \in (0, 1)$ .

Gittins & Jones proved that optimal performance can be obtained by tracking a single index  $v_i$  of the form

$$v_i = \sup_{T>0} \frac{\langle \sum_{t=0}^T \delta^t R_i(t) \rangle}{\langle \sum_{t=0}^T \delta^t \rangle}, \quad (2.1)$$

for each of the bandits, which is a normalized sum of future rewards discounted by the delay until they are accrued. The sum is taken until a time  $T$ , which is defined as the stopping time, or the point at which selecting from bandit  $i$  will be terminated. Gittins & Jones proved that optimal behaviour is assured as long as that action is always taken which has the greatest index value. Critically, the Gittins index for any given bandit is independent of the expected outcomes of all other bandits. This implies that once the bandit with greatest index is known, behaviour should continue on this bandit until its index value falls below its original value. This is true because the index values for all other bandits do not change as long as these bandits are not selected. Computationally, calculating the Gittins index (equation (2.1)) is demanding and may not reasonably be expected to be calculated in the brain.

The Gittins index provides a normative account of how agents should act when faced with a particular form of the exploration–exploitation dilemma. Krebs *et al.* (1978) tested whether the foraging behaviour of birds is optimal when confronting a two-armed bandit problem similar to that solved by the Gittins index. In the experiment, the birds were presented with two feeding posts that gave food reward with fixed probability. The problem was a simplification of the general problem solved by the Gittins index, since the probability of obtaining reward from a feeding post was not allowed to change when selected and since the experiment was of finite length. The investigators found that the time at which birds stopped exploring (operationalized as the point at which they stayed at one feeding post) closely approximated that predicted by the optimal solution. Despite their findings, Krebs *et al.* (1978) recognized that it was highly unlikely that their birds were carrying out the complex calculations required by the Gittins index. Rather, they suggested that the birds were using simple behavioural heuristics that produces exploration times that qualitatively approximate the optimal solution. However, there are more fundamental problems with the Gittins index, beyond complexity of calculation.

As noted earlier, Gittins' proof requires that rewards should be discounted exponentially for delay (Berry & Fristedt 1985) whereas it is generally accepted that most animals (including humans) show hyperbolic discounting (e.g. Ainslie 1975). Additionally, if there is a cost associated with switching from one behaviour to another, then not only is the Gittins index no longer optimal, but also there is *no* optimal index that may be calculated independently for each bandit (Banks & Sundaram 1994). It is well recognized that, under many conditions, humans exhibit costs when switching from one task to another (e.g. Allport *et al.* 1994; Rogers & Monsell 1995). Most importantly, the Gittins index assumes that, although the pay-offs for each bandit are probabilistic and each must be sampled sufficiently to determine its expected value, the *actual*

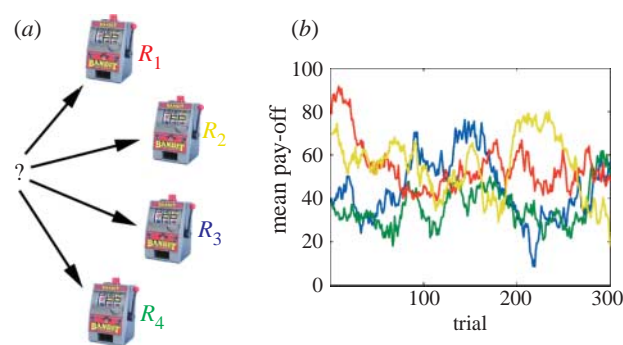


Figure 1. Daw *et al.* (2006) examined how subjects handle the exploration–exploitation problem in a four-armed bandit problem. (a) In each trial of their task, subjects selected one of the four bandits and received a reward based on its current mean pay-off perturbed by noise. (b) The expected value of each bandit changed continuously over time.

expected value of each remains fixed except when acted upon. That is, if nothing is done to a bandit, then its true value remains stable across time. However, both the needs of most organisms and the environments in which they live are not stable in this way. Things change over time, even when they are not acted upon, and often in unpredictable ways. To date, no universally optimal algorithm has been described that prescribes how to trade-off between exploration and exploitation in non-stationary environments, and it is not clear that doing so is possible. Thus, understanding how animals respond to this problem must also be guided by empirical investigation, both of behaviour and underlying neural mechanisms.

### 3. MODELLING EXPLOITATION VERSUS EXPLORATION IN NON-STATIONARY ENVIRONMENTS

Daw *et al.* (2006) recently addressed this problem in a study that used a variant of the  $n$ -armed bandit problem in which the pay-offs of each bandit changed slowly over time (figure 1). In this setting, therefore, the cost of persisting with one behaviour (i.e. playing only one bandit) was not only the opportunity cost of failing to learn more about the value of the others, but also the possibility that what has already been learned about them will fall out of date. Daw *et al.* (2006) proposed three possible models for how subjects might guide their choices in this situation.

The first model used a simple decision rule, in which the subject maintains a record of the expected value for each option, based on past experience, and usually chooses the option with the greatest value (exploitation) though sometimes, with a fixed probability, picks randomly among the other alternatives (exploration). This is often referred to as the ‘epsilon-greedy’ algorithm (Sutton & Barto 1998). According to a second model, options are chosen by probability matching, i.e. with a probability weighted by their estimated values. This is often referred to as the ‘soft max’ decision rule (e.g. Thrun 1992), as it favours choosing the option with the maximum value (this option will have the highest probability), though this tendency is ‘softened’ by both the value of the competing options as well as randomness (noise)

added to the decision rule. Thus, in this model, the balance between exploitation and exploration is governed by both the relative value of the alternatives as well as a parameter (referred to as gain or, inversely, temperature) that determines how tightly decisions are constrained by the contrast of value among the alternatives: with higher gain, decisions are determined more by relative value (exploitation); with lower gain, decisions are more evenly distributed at random (exploration).

Finally, they entertained a third model, according to which choices are made using the soft max decision rule, but with a critical added factor: options that have not been selected receive an ‘uncertainty bonus’ that augments their probability of being chosen (i.e. promotes exploration). This captures the opportunity cost that is formalized by the Gittins index for stationary environments, and that is particularly important in non-stationary environments: the more time allocated to one option the less one knows about the others, which may be (or have become) more valuable.

Daw *et al.* (2006) compared the behaviour of subjects playing their *n*-arm bandit task to predictions from each of the three models. The model that provided the best fit was the soft max decision rule. Importantly, although subjects did periodically explore options other than the one currently deemed to be most valuable, they did not find evidence that this was driven by an uncertainty bonus (i.e. growing uncertainty about the competing alternatives). However, there are several caveats that must be kept in mind. First, it is possible that the specifics of the environment did not adequately favour the use of an uncertainty bonus. For example, the pay-offs of each bandit changed continuously and relatively slowly over time in their experiment. In the real world—to which real-world organisms are presumably adapted—the dynamics of environmental change may be very different, and therefore call for a different policy of exploration (and computation of uncertainty bonus) than was assumed by Daw *et al.* (2006). Another important factor may be social context—people may be enticed to explore the environment when they have information about the behaviour of others, and they may also place a greater premium on exploration when they face competition from others for resources.

These are questions that beg more detailed formal analysis. Nevertheless, to our knowledge, the Daw *et al.* (2006) study was the first to address formally the question of how subjects weigh exploration against exploitation in a non-stationary, but experimentally controlled environment. It also produced some interesting neurobiological findings. Their subjects performed the *n*-armed bandit task while being scanned using functional magnetic resonance imaging (fMRI). Among the observations reported was task-related activity in two sets of regions of prefrontal cortex (PFC). One set of regions was in ventromedial PFC and was associated with both the magnitude of reward associated with a choice, and that predicted by their computational model of the task (using the soft max decision rule). This area has been consistently associated with the encoding of reward value across a variety of task domains (O’Doherty *et al.* 2001; Knutson *et al.* 2003; McClure *et al.* 2004; Padoa-Schioppa & Assad

2006). A second set of areas observed bilaterally in frontopolar PFC was significantly more active when subjects chose to explore (i.e. chose an option other than the one estimated by their model to be the most rewarding) rather than exploit. This finding is consistent with the hypothesis that more anterior and dorsal regions of PFC are responsible for top-down control, biasing processes responsible for behaviour in favour of higher level goals, especially when these must compete with otherwise prepotent behaviours (e.g. Miller & Cohen 2001). Such top-down control may be important for exploration, insofar as this involves selecting an action that has been less recently associated with reward. That is, when a decision is made to pursue an exploratory behaviour, this may rely on support from higher level control processes. However, this begs the question of how the system decides when it is appropriate to explore. That is, what mechanisms are responsible for assessing the reliability and value of current rewards, and using this information to determine when to continue to pursue current sources of reward (exploit) or take a chance in pursuing new behaviours (explore). Several lines of investigation have begun to address this question.

#### 4. UNCERTAINTY AND EXPLOITATION VERSUS EXPLORATION

One line of work that has direct relevance addresses the question of how the brain encodes different forms of uncertainty. Yu & Dayan (2005) proposed that a critical function of two important neuromodulators—acetylcholine (ACh) and norepinephrine (NE)—may be to signal expected and unexpected sources of uncertainty. While the model they developed for this was not intended to address the trade-off between exploitation and exploration, the distinction between expected and unexpected uncertainty is likely to be an important factor in regulating this trade-off. For example, the detection of unexpected uncertainty can be an important signal of the need to promote exploration. To see this, consider the following scenario.

You are asked to observe a series of coin tosses, told that the coin is biased, and your job is to determine whether it is biased towards heads or tails. The first several tosses produce the following sequence: heads, heads, tails, heads, heads, heads, heads. If you are forced to choose at this point, like most observers, you would probably say that the coin is biased towards heads. If the next flip comes up tails, that is OK. You know that the outcome of any particular toss is uncertain. This represents an *expected* form of uncertainty. However, consider what happens if the subsequent set of tosses is: heads, tails, tails, tails, tails, tails, tails... At some point, you will revise your determination and say that the coin is biased towards tails. Perhaps, the coin was surreptitiously switched (i.e. the world has changed) or your determination was wrong in the first place. In either case, having come to assume that the coin is biased towards heads, you have now been confronted with an *unexpected* form of uncertainty and must revise your model of the world accordingly, along with the choice of any actions that depend on it.

This problem is closely related to the example we gave at the beginning of this article (concerning the

collection of experimental data), and as we have noted elsewhere (Aston-Jones & Cohen 2005), the distinction between expected and unexpected forms of uncertainty may be an important element in choosing between exploitation versus exploration. As long as prediction errors can be accounted for in terms of expected uncertainty—that is the amount that we expect a given outcome to vary—then all other things being equal (e.g. ignoring potential non-stationarities in the environment), we should persist in our current behaviour (exploit). However, if errors in prediction begin to exceed the degree expected—i.e. unexpected uncertainty mounts—then we should revise our strategy and consider alternatives (explore).

Yu & Dayan (2005) proposed that ACh levels are used to signal expected uncertainty, and NE to signal unexpected uncertainty. They describe a computationally tractable algorithm by which these may be estimated that approximates the Bayesian optimal computation of those estimates. Furthermore, they proposed how these estimates, reflected by NE and ACh levels, could be used to determine when to revise expectations

$$NE > \frac{ACh}{(0.5 + ACh)}. \quad (4.1)$$

They showed that this closely approximates the Bayesian optimal solution to, and people's behaviour in, a variant of a commonly used selective attention task (the 'Posner paradigm'; Posner *et al.* 1980).

This work provides another instructive example of the value in conducting a mathematical analysis of optimal performance in a task, and using this to guide the generation of hypotheses about the specific mechanisms—in this case neural—that govern behaviour in that task. Furthermore, it lends precision to hypotheses about the function of neuromodulatory systems. Despite their ubiquity in the brain, theories about these systems have typically been vague, proposing non-specific functions such as the mediation of motivation and arousal. Yu and Dayan's model assigns precise functions to ACh and NE, specified in mathematical form, that can be used to generate specific testable predictions.

As suggested above, it is not hard to imagine how the functions ascribed to ACh and NE in representing estimates of expected and unexpected forms of uncertainty might play an important role in regulating the balance between exploitation and exploration. As estimates of unexpected uncertainty rise, and NE approaches the threshold defined by equation (4.1), the system promotes a revision of current expectations. This could be an important signal to search for a new model of the environment and a corresponding behavioural strategy—i.e. exploration. Sometimes, however, unexpected events are followed by the opposite tendency: an increase in commitment to the current behavioural strategy. For example, following errors in simple reaction time tasks people often become more cautious and improve their performance (i.e. become more accurate; Rabbitt 1966; Laming 1979). Similarly, following interference in selective attention tasks, subjects typically increase the focus of their attention and improve performance (Gratton *et al.*

1992), especially when such interference is relatively rare (Carter *et al.* 2000; Kerns *et al.* 2004).

The Yu & Dayan model also sensibly predicts that performance should be best when expectations are most accurate. However, when outcomes in a task become *too* predictable, people often become bored and look for other things to do (explore). Video game programmers learned this lesson long ago, and routinely include multiple levels in a game, so that when it becomes too predictable, it is made more difficult in order to retain players' interest (i.e. keep them exploiting).

These observations suggest that additional mechanisms may be involved in evaluating expectations and in regulating the trade-off between exploration and exploitation. Another closely related line of investigation has sought to address some of these observations. It too has suggested an important role for NE, building on detailed physiological observations about the dynamics of NE release, and proposing how this may relate to assessments of reward as well as uncertainty.

## 5. UTILITY AND EXPLOITATION VERSUS EXPLORATION

Virtually all of the NE released in the neocortex originates from a small brainstem nucleus called the LC. Aston-Jones *et al.* (1994, 1997) have observed that in the awake behaving monkey the LC shifts between two operating modes that correspond closely with behavioural performance in a simple target detection task. In the 'phasic mode,' when the animal is performing well (no misses and very few false alarms), the LC shows only moderate levels of tonic discharge, but responds phasically with a burst of activity to target stimuli (but not to distractors). In the 'tonic mode,' the baseline level of discharge is higher, but there are diminished or absent phasic responses to target stimuli. In this mode, reaction time to targets is slower and the animal commits a greater number of false alarms to distractors. These two modes most probably represent a continuum of LC function, consistent with the formal theories described below. However, we will continue to refer to two modes for expository purposes, because the distinction between them (or the extremes of function they represent) has been proposed to be an important factor in influencing the balance between exploration and exploitation.

Usher *et al.* (1999) developed a biophysically detailed model of the LC that accounted for the physiological observations outlined above and suggested that these may play a role in regulating the balance between exploitation and exploration. They proposed that the phasic mode favours exploitation by releasing NE specifically when a task-relevant event occurs, thereby facilitating processing of that event. In contrast, in the tonic mode, sustained release of NE indiscriminately facilitates processing of all events irrespective of their relevance to the current task and thereby favours exploration. Note that the latter aligns well with the role of NE proposed by Yu & Dayan (2005), favouring exploration, if it is assumed that NE in their model corresponds to tonic release.

The Usher *et al.* (1999) model describes physiological mechanisms by which the LC may contribute to regulating the balance between exploitation and

exploration. However, it does not specify what drives the LC towards the phasic (exploitation) or tonic (exploration) modes. Recently, [Aston-Jones & Cohen \(2005\)](#) have proposed that this may be governed by ongoing assessments of utility carried out in ventral and medial frontal structures. As noted earlier, there is extensive evidence that ventral regions within PFC form part of a circuit responsible for encoding reward value (e.g. [Knutson \*et al.\* 2003](#); [O'Doherty \*et al.\* 2001](#); [McClure \*et al.\* 2004](#); [Padoa-Schioppa & Assad 2006](#)). There is also now a substantial body of evidence that medial frontal structures, and in particular the anterior cingulate cortex (ACC), encode costs. Regions within the ACC have consistently been observed to respond to pain, negative feedback, errors in performance, conflicts in processing and even mental effort, all of which represent or are indicative of various forms of cost (e.g. [Miltner \*et al.\* 1997](#); [Carter \*et al.\* 1998](#); [Peyron \*et al.\* 2000](#); [Botvinick \*et al.\* 2001](#); [Holroyd & Coles 2002](#); [Yeung \*et al.\* 2004](#)). Furthermore, recent anatomic evidence indicates that these ventral and medial frontal structures provide dense projections to the LC ([Rajkowski \*et al.\* 2000](#); [Aston-Jones \*et al.\* 2002](#)).

Based on these findings, [Aston-Jones & Cohen \(2005\)](#) have proposed that ongoing assessments of utility carried out in frontal structures are used to govern the mode of LC and thereby regulate the balance between exploitation and exploration. Specifically, they propose that assessments of utility are carried out over both short (e.g. seconds) and long (e.g. minutes) time-scales and that this can reconcile the opposing tendencies (to 'try harder' versus 'give up') following periods of poor performance noted above. For example, consider the following two circumstances. In one, performance in a task has been good and there are still rewards to be accrued from the task, but there are occasional lapses in performance producing transient decreases in utility (e.g. on single trials). In this case, following such a lapse the agent should act to restore performance. That is, exploitation should be promoted when long-term utility has been high, but there has been a momentary decrease. In contrast, consider a second situation in which performance has been poor and utility has progressively declined. At some point, this should encourage disengagement from the current task and exploration of alternative behaviours. That is, how the system responds to a current decrease in utility should depend upon the context of longer term trends in utility, favouring exploitation if long-term utility has been high, and exploration if it has been low. A relatively simple equation can capture these relationships,

Engagement in current task

$$= [1 - \text{logistic}(\text{short-term utility})] \times [\text{logistic}(\text{long-term utility})], \quad (5.1)$$

where *logistic* refers to the sigmoid function  $1/(1 + e^{-\text{utility}})$ . [Aston-Jones & Cohen \(2005\)](#) proposed that high values of this equation favour the LC phasic mode (exploitation), whereas low values favour the tonic mode (exploration; [figure 2](#)). [Usher \*et al.\* \(1999\)](#) and [Brown \*et al.\* \(2005\)](#) both suggest the ways in which this can be accomplished through the regulation of simple

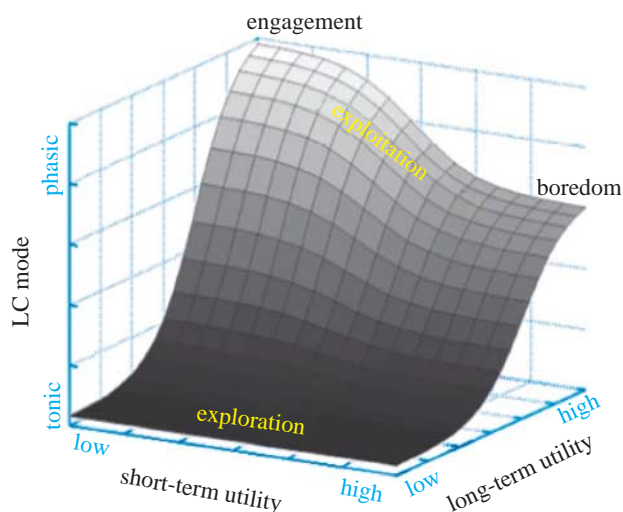


Figure 2. [Aston-Jones & Cohen \(2005\)](#) propose that exploration and exploitation may be mediated by separate short- and long-term measures of utility (cost and reward). Exploration and exploitation, in this model, are mediated by the firing mode of norepinephrine neurons in the locus coeruleus (LC).

physiological parameters (such as electronic coupling and/or baseline afferent drive) within the LC.

This model can also be related to the soft max mechanism that [Daw \*et al.\* \(2006\)](#) found best fits decision-making behaviour in their *n*-armed bandit task. The effect of the LC can be thought of as tuning the softmax function, sharpening it (phasic mode) and biasing decisions towards the most recently rewarded choices (i.e. exploitation) when long-term utility is high, and flattening the function (tonic mode) promoting a more uniform distribution of choices (exploration) when long-term utility is low. Whether such effects are observed in a suitably designed *n*-armed bandit decision-making task remains to be tested. However, recent findings from a simpler, two-armed decision-making task, that used pupilometry to index LC activity ([Aston-Jones & Cohen 2005](#)), have corroborated predictions of the model regarding the relationship of LC activity to decision-making performance ([Gilzenrat & Cohen in preparation](#)). This work has also recently been extended to explore the interaction between these mechanisms and those underlying RL.

## 6. REINFORCEMENT LEARNING AND EXPLOITATION VERSUS EXPLORATION

The trade-off between exploration and exploitation has long been recognized as a central issue in RL ([Kaelbling 1996, 2003](#)). The RL mechanisms act by strengthening associations (e.g. between a stimulus and an action) when these have been associated with a reward (e.g. [Sutton & Barto 1998](#)). There is now strong reason to believe that the dopaminergic (DA) system implements such a mechanism ([Montague \*et al.\* 1996](#); see [Montague \*et al.\* 2004](#) for a recent review). The RL mechanisms function well in stationary environments, in which progressive strengthening of associations makes them robust and efficient, allowing the agent to exploit the current environment. However, this also makes them resistant to change, which is problematic

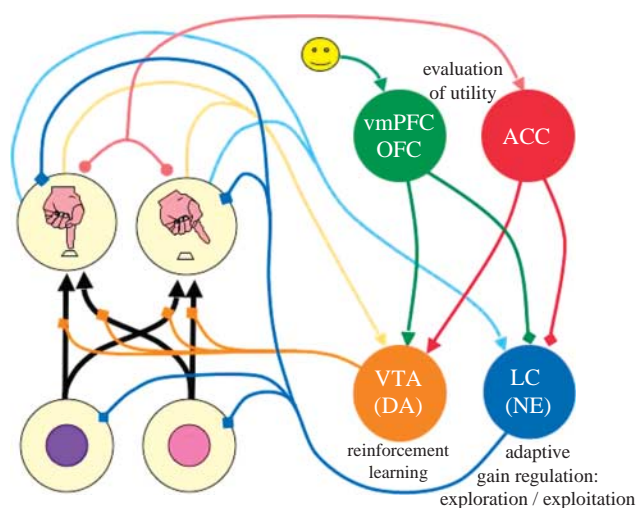


Figure 3. A neural network model of how reward and cost are integrated in the locus coeruleus to adaptively change between exploration and exploitation, as proposed by McClure *et al.* (2006). The left side shows a simple network for decision making in the task. The right side shows evaluative and neuromodulatory mechanisms that regulate the decision-making mechanisms. The model proposes that information about cost (calculated by the anterior cingulate cortex (ACC)) and reward (calculated by the ventromedial prefrontal cortex (vmPFC) and orbitofrontal cortex (OFC)) converge on both the ventral tegmental area (VTA) and the locus coeruleus (LC). This information is used by the VTA to implement a reinforcement learning algorithm that adjusts the weights in the decision network. In the LC, evaluative information sets the mode of responding (phasic or tonic), which, through norepinephrine (NE) release and gain modulation of units in the decision network, regulates the balance between exploration and exploitation (see text for more detailed description).

in non-stationary environments when the system must be able to explore and learn new contingencies.

The simplest example of this is a reversal conditioning paradigm, in which the agent learns a set of associations (e.g. that a purple light calls for a response and a pink light does not) and once they are learned the contingencies are reversed. If the RL mechanism ensures rapid and strong learning of the initial association, then it will be difficult to adjust to the change (the purple light will continue to elicit a response). However, if RL operates more weakly, then it will take longer to learn the initial association. A common solution to this problem is to introduce an annealing mechanism. When new learning is required (i.e. there is uncertainty about the environment, and/or utility declines), noise is added to the system, allowing it to randomly explore new associations; noise is then progressively reduced as newly rewarded associations are discovered and these are strengthened. This is similar to the tuning of the softmax decision function described above. Indeed, McClure *et al.* (2006) have described a model showing how the frontal and LC mechanisms described above can function as such an annealing mechanism when integrated with a DA-based RL mechanism (figure 3).

Furthermore, they have shown that the behaviour of the LC in this model closely parallels observations that have been made from LC recordings in a reversal

conditioning paradigm using a target detection task (Aston-Jones *et al.* 1997). Performance of the task following acquisition of the initial target was associated with the LC phasic mode. When the contingencies were reversed, LC tonic activity increased and phasic responses diminished. Then, as the new target was acquired, the LC returned to the phasic mode of responding. These findings provide growing support for the view that the LC noradrenergic system plays an important role in mediating the balance between exploitation and exploration. As the work of Aston-Jones & Cohen (2005) and Yu & Dayan (2005) suggests, ongoing assessments of both uncertainty and utility are likely to be important in regulating this balance.

## 7. OPEN QUESTIONS AND CHALLENGES

In this article, we hope to have drawn attention to the fact that managing the trade-off between exploitation and exploration is a fundamental challenge for the adaptive control of behaviour. While traditionally this has not occupied centre stage in research on executive function and cognitive control, we have reviewed several lines of work that have productively begun to address this issue. Nevertheless, many important questions remain.

First, it should be noted that some of the work we have reviewed addresses the estimation of uncertainty (e.g. Yu & Dayan 2005), while other work focuses more on the computation of utility and action selection (e.g. Usher *et al.* 1999; Aston-Jones & Cohen 2005; Daw *et al.* 2006). All of these are likely to be critical elements in determining the trade-off between exploitation and exploration. However, the specific relationship between these remains to be examined directly. For example, it would be valuable to understand how the mechanisms proposed by Yu & Dayan (2005) to compute assessments of uncertainty (i.e. prediction errors) can be coupled to action selection, and how this relates to the algorithm described by equation (5.1)—proposed by Aston-Jones & Cohen (2005) to relate assessments of utility to LC function and decision-making performance.

It seems inescapable that, in addition to uncertainty and utility, social signals are a critical factor adjudicating the trade-off between exploitation and exploration. Observing others can provide critical counterfactual information about the reward value of behavioural strategies that one has not yet pursued oneself (Montague *et al.* in press). Competition within a social context may also help explain aspects of boredom—i.e. the perplexing tendency to explore alternatives to current behaviour when certainty of outcome (including reward) is at its highest. If it is assumed that more difficult tasks are both more remunerative and less competitive (because fewer agents possess the skills necessary to perform them), then performing a task below one's skill level carries an opportunity cost. That is, it should be possible to find another task for which one is still adequately competent, but that is more difficult and less competitive, and therefore more remunerative. Thus, boredom may in part reflect an adaptive bias towards exploration when performance at ceiling suggests that a more remunerative task can be found (M. Todd 2006, personal communication).



An important super-ordinate question is whether the trade-off between exploitation and exploration should be considered a single problem addressed by a unitary set of mechanisms in the brain, or whether it represents a family of problems spanning different scales, that are addressed by different mechanisms. The time-scale of neuromodulatory function suggests that these mechanisms influence decisions that take place over seconds or minutes. However, faster processes (e.g. saccadic search mechanisms) and longer ones (planning a career) may involve very different mechanisms.

Finally, an equally pressing question is whether it is best to distinguish qualitatively between exploitation and exploration, or whether these represent the extremes of a continuum. For example, the models we have discussed have, for the most part, treated exploration as random search (e.g. increasing noise in an annealing procedure). However, search can often be structured by relatively sophisticated, domain-specific heuristics (for example, in problem solving tasks; Newell & Simon 1972). Such search processes may involve temporally extended, goal-directed behaviours that rely on mechanisms of cognitive control similar to those required for exploitation within the context of simpler tasks. Indeed, the findings of an association between PFC activity and exploration in the Daw *et al.* (2006) study may provide an example of this. These considerations help underscore the need for a precise formulation of the exploitation–exploration trade-off within specific task environments.

More generally, this issue brings into focus an important dimension for considering the trade-off between exploration and exploitation: the extent to which the environment to be explored is well-structured (whether static or changing in predictable ways) versus unknown and unpredictable. To the extent that it is structured, then it should be possible to explore it in a systematic fashion (we might refer to this as ‘controlled exploration’). That is, at least from the theorist’s perspective, it should be possible to identify an optimal strategy for exploration that takes account of knowledge about the various behavioural alternatives (the Gittins index represents a special case of this). Under such conditions, the decision of whether to exploit or explore should weigh both the value of current pursuits as well as informed expectations about the alternatives, and exploration should be deterministic. Indeed, to the extent that an optimal strategy can be found, this might be thought of simply as higher level exploitation. Of course, for realistically complex environments, theoretically optimal strategies are likely to be computationally intractable, at least for biological mechanisms (this is so for the Gittins index, even given its simplifying assumptions). Thus approximations, including stochastic ones (such as some of the mechanisms reviewed in this article) may be more biologically realistic.

At the other end of this dimension are unknown and unpredictable environments. Under such conditions, the decision of whether to exploit or explore may focus more profitably on assessments of performance in the current task rather than on expectations about alternatives. Similarly, strategies for exploration will necessarily rely on cruder assumptions about behavioural alternatives and search among them will be less structured and

presumably more stochastic. Consideration of these factors may be useful in guiding the next generation of hypotheses about the mechanisms governing exploitation and exploration in biological organisms.

## 8. SUMMARY AND CONCLUSIONS

This article began by reviewing efforts to formalize the optimal solution to the trade-off between exploitation and exploration. The Gittins index provides such a solution, but applies to restricted circumstances (e.g. only stationary environments). As yet, no general solution has been found for non-stationary environments and, depending upon the breadth and characteristics of the environment to be considered, this may not be possible. Nonetheless, empirical studies of both behaviour and neural mechanisms have begun to reveal mechanisms that animals may use to adapt to changes in the environment, by regulating the balance between exploitation and exploration. These studies appear to be converging on the view that neuromodulatory systems—in particular, ACh and NE, interacting with DA-mediated RL mechanisms—may play a critical role in regulating this balance within certain domains of behaviour. These systems appear to be responsive to both estimates of uncertainty and utility. However, social signals are also likely to be an important source of information. More generally, the trade-off between exploitation and exploration represents a challenge to behaviour at all levels and over multiple time-scales. It is not yet clear whether neuromodulatory mechanisms serve the same function at all of these levels and time-scales, or whether this relies on other mechanisms that remain to be discovered. Given these considerations, it seems probable that further research will require a mixed (though not yet fully informed) strategy of continuing to exploit promising lines of recent work, while considering new ones to explore.

This work was supported by NIH grants P50 MH062196 (J.D.C.), F32 MH072141 (S.M.M.) and the NIMH Quantitative Neuroscience Training Grant (MH65214). We thank Gary Aston-Jones, Eric Brown, Mark Gilzenrat, Phil Holmes and Leigh Nystrom for their close collaboration in the development of many of the ideas presented in this article. We would also like to thank Nathaniel Daw, Yael Niv and Greg Stephens for their valuable discussions related to this work, as well as Peter Dayan and an anonymous reviewer for their useful suggestions regarding this article. Finally, we would like to offer a profound and heartfelt thanks to Tim Shallice, not only for his comments on this article, but more importantly for the decades of inspiring work and visionary leadership that he has provided to our field.

## REFERENCES

- Ainslie, G. 1975 Specious reward: a behavioral theory of impulsiveness and impulse control. *Psychol. Bull.* **82**, 463–496. (doi:10.1037/h0076860)
- Allport, A., Styles, E. & Hsieh, S. 1994 Shifting intentional set: exploring the dynamic control of task. In *Attention and performance XV* (eds C. Umiltà & M. Moscovitch), pp. 421–452. Cambridge, MA: MIT Press.
- Aston-Jones, G. & Cohen, J. D. 2005 An integrative theory of locus coeruleus–norepinephrine function: adaptive gain and optimal performance. *Annu. Rev. Neurosci.* **28**, 403–450. (doi:10.1146/annurev.neuro.28.061604.135709)

- Aston-Jones, G., Rajkowski, J., Kubiak, P. & Alexinsky, T. 1994 Locus coeruleus neurons in monkey are selectively activated by attended cues in a vigilance task. *J. Neurosci.* **14**, 4467–4480.
- Aston-Jones, G., Rajkowski, J. & Kubiak, P. 1997 Conditioned responses in monkey locus coeruleus neurons anticipate acquisition of discriminative behavior in a vigilance task. *Neuroscience* **80**, 697–715. (doi:10.1016/S0306-4522(97)00060-2)
- Aston-Jones, G., Rajkowski, J., Lu, W., Zhu, Y., Cohen, J. D. & Morecraft, R. J. 2002 Prominent projections from the orbital prefrontal cortex to the locus coeruleus in monkeys. *Soc. Neurosci. Abstr.* **28**, 86–89.
- Banks, J. S. & Sundaram, R. K. 1994 Switching costs and the Gittins index. *Econometrica: J. Econ. Soc.* **62**, 687–694.
- Berry, D. A. & Fristedt, B. 1985 *Bandit problems: sequential allocation of experiments*. London, UK: Chapman and Hall.
- Botvinick, M. M., Braver, T. S., Barch, D. M., Carter, C. S. & Cohen, J. D. 2001 Conflict monitoring and cognitive control. *Psychol. Rev.* **108**, 624–652. (doi:10.1037/0033-295X.108.3.624)
- Brown, E., Gao, J., Bogacz, R., Gilzenrat, M. & Cohen, J. D. 2005 Simple neural networks that optimize decisions. *Int. J. Bifurc. Chaos* **15**, 803–826. (doi:10.1142/S0218127405012478)
- Carstensen, L. L., Isaacowitz, D. & Charles, S. T. 1999 Taking time seriously: a theory of socioemotional selectivity. *Am. Psychol.* **54**, 165–181. (doi:10.1037/0003-066X.54.3.165)
- Carter, C. S., Braver, T. S., Barch, D. M., Botvinick, M. M., Noll, D. C. & Cohen, J. D. 1998 Anterior cingulate cortex, error detection and the on-line monitoring of performance. *Science* **280**, 747–749. (doi:10.1126/science.280.5364.747)
- Carter, C. S., Macdonald, A. M., Botvinick, M., Ross, L. L., Stenger, V. A., Noll, D. & Cohen, J. D. 2000 Parsing executive processes: strategic vs. evaluative functions of the anterior cingulate cortex. *Proc. Natl Acad. Sci. USA* **97**, 1944–1948. (doi:10.1073/pnas.97.4.1944)
- Daw, N. D., O'Doherty, J. P., Seymour, B., Dayan, P. & Dolan, R. J. 2006 Cortical substrates for exploratory decisions in humans. *Nature* **441**, 876–879. (doi:10.1038/nature04766)
- Gilzenrat M. S. & Cohen J. D. In preparation. The role of locus coeruleus in mediating between exploration and exploitation in nonstationary environments: an empirical test in a changing utility task.
- Gittins, J. C. 1979 Bandit processes and dynamic allocation indices. *J. R. Stat. Soc. B* **41**, 148–177.
- Gittins, J. C. & Jones, D. M. 1974 A dynamic allocation index for the sequential design of experiments. In *Progress in statistics* (ed. J. Gans), pp. 241–266. Amsterdam, The Netherlands: North-Holland.
- Gratton, G., Coles, M. G. H. & Donchin, E. 1992 Optimization in the use of information: strategic control of activation and responses. *J. Exp. Psychol. Gen.* **4**, 480–506. (doi:10.1037/0096-3445.121.4.480)
- Holroyd, C. B. & Coles, M. G. H. 2002 The neural basis of human error processing: reinforcement learning, dopamine, and the error-related negativity. *Psychol. Rev.* **109**, 679–709. (doi:10.1037/0033-295X.109.4.679)
- Kaelbling, L. P. 1996 Gittins Allocation Indices. See <http://www.cs.cmu.edu/afs/cs/project/jair/pub/volume4/kaelbling96a-html/node9.html>.
- Kaelbling, L. P. 2003 *Learning in embedded systems*. Cambridge, MA: MIT Press.
- Kaelbling, L. P., Littman, M. L. & Moore, A. W. 1996 Reinforcement learning: a survey. *J. Artif. Intell. Res.* **4**, 237–285.
- Kerns, J. G., Cohen, J. D., MacDonald, A. W., Cho, R. Y., Stenger, V. A. & Carter, C. S. 2004 Anterior cingulate conflict monitoring and adjustments in control. *Science* **303**, 1023–1026. (doi:10.1126/science.1089910)
- Knutson, B., Fong, G. W., Bennett, S. M., Adams, C. M. & Hommer, D. 2003 A region of the mesial prefrontal cortex tracks monetary rewarding outcomes: characterization with rapid event-related fMRI. *Neuroimage* **18**, 263–272. (doi:10.1016/S1053-8119(02)00057-5)
- Krebs, J. R., Kacelnik, A. & Taylor, P. 1978 Tests of optimal sampling by foraging great tits. *Nature* **275**, 27–31. (doi:10.1038/275027a0)
- Laming, D. R. J. 1979 Choice reaction performance following an error. *Acta Psychologica* **43**, 199–224. (doi:10.1016/0001-6918(79)90026-X)
- Leonard, N. E., Paley, D., Lekien, F., Sepulchre, R., Fratantoni, D. M. & Davis, R. E. In press. Collective motion, sensor networks and ocean sampling. *Proc. IEEE*, **95**.
- McClure, S. M., Daw, N. D. & Montague, P. R. 2003 A computational substrate for incentive salience. *Trends Neurosci.* **26**, 423–428. (doi:10.1016/S0166-2236(03)00177-2)
- McClure, S. M., Laibson, D. I., Loewenstein, G. & Cohen, J. D. 2004 Separate neural systems value immediate and delayed monetary reward. *Science* **306**, 503–507. (doi:10.1126/science.1100907)
- McClure, S. M., Gilzenrat, M. S. & Cohen, J. D. 2006 An exploration–exploitation model based on norepinephrine and dopamine activity. In *Advances in neural information processing systems*, vol. 18 (eds Y. Weiss, B. Sholkopf & J. Platt), pp. 867–874. Cambridge, MA: MIT Press.
- Miller, E. K. & Cohen, J. D. 2001 An integrative theory of prefrontal cortex function. *Annu. Rev. Neurosci.* **24**, 167–202. (doi:10.1146/annurev.neuro.24.1.167)
- Miltner, W. H. R., Braun, C. H. & Coles, M. G. H. 1997 Event-related potentials following incorrect feedback in a time-estimation task: evidence for a 'generic' neural system for error detection. *J. Cogn. Neurosci.* **9**, 788–798.
- Montague, P. R., Dayan, P. & Sejnowski, T. J. 1996 A framework for mesencephalic dopamine systems based on predictive Hebbian learning. *J. Neurosci.* **16**, 1936–1947.
- Montague, P. R., Hyman, S. E. & Cohen, J. D. 2004 Computational roles for dopamine in behavioral control. *Nature* **431**, 760–767. (doi:10.1038/nature03015)
- Montague, P. R., King-Casas, B. & Cohen, J. D. In press. Imaging valuation models in human choice. *Annu. Rev. Neurosci.* **29**, 417–448. (doi:10.1146/annurev.neuro.29.051605.112903)
- Newell, A. & Simon, H. A. 1972 *Human problem solving*. Englewood Cliffs, NJ: Prentice-Hall.
- O'Doherty, J., Kringelback, M. L., Rolls, E. T., Hornak, J. & Andrews, C. 2001 Abstract reward and punishment representation in the human orbitofrontal cortex. *Nat. Neurosci.* **4**, 95–102. (doi:10.1038/82959)
- Padoa-Schioppa, C. & Assad, J. A. 2006 Neurons in the orbitofrontal cortex encode economic value. *Nature* **441**, 223–226. (doi:10.1038/nature04676)
- Peyron, R., Laurent, B. & Garcia-Larrea, L. 2000 Functional imaging of brain responses to pain: a review and meta-analysis. *Neurophysiol. Clin.* **30**, 263–288. (doi:10.1016/S0987-7053(00)00227-6)
- Posner, M. I., Snyder, C. R. R. & Davidson, B. J. 1980 Attention and the detection of signals. *J. Exp. Psychol. Gen.* **109**, 160–174. (doi:10.1037/0096-3445.109.2.160)
- Pratt, S. C. & Sumpter, D. J. T. 2006 A tunable algorithm for collective decision-making. *Proc. Natl Acad. Sci. USA* **103**, 15 906–15 910. (doi:10.1073/pnas.0604801103)

- Rabbitt, P. M. A. 1966 Errors and error-correction in choice-response tasks. *J. Exp. Psychol.* **71**, 264–272. (doi:10.1037/h0022853)
- Rajkowski, J., Lu, W., Zhu, Y., Cohen, J. D. & Aston-Jones, G. 2000 Prominent projections from the anterior cingulate cortex to the locus coeruleus in Rhesus monkey. *Soc. Neurosci. Abstr.* **26**, 838.15.
- Rogers, R. & Monsell, S. 1995 The costs of a predictable switch between simple cognitive tasks. *J. Exp. Psychol. Gen.* **124**, 207–231. (doi:10.1037/0096-3445.124.2.207)
- Schultz, W., Dayan, P. & Montague, P. R. 1997 A neural substrate of prediction and reward. *Science* **275**, 1593–1599. (doi:10.1126/science.275.5306.1593)
- Sugrue, L. P., Corrado, G. S. & Newsome, W. T. 2004 Matching behavior and the representation of value in the parietal cortex. *Science* **304**, 1782–1787. (doi:10.1126/science.1094765)
- Sutton, R. S. & Barto, A. G. 1998 *Reinforcement learning: an introduction*. Cambridge, MA: MIT Press.
- Thrun, S. B. 1992 The role of exploration in learning control. In *Handbook of intelligent control: neural, fuzzy, and adaptive approaches* (eds D. A. White & D. A. Sofge), pp. 527–559. Florence, KY: Van Nostrand Reinhold.
- Usher, M., Cohen, J. D., Rajkowski, J. & Aston-Jones, G. 1999 The role of the locus coeruleus in the regulation of cognitive performance. *Science* **283**, 549–554. (doi:10.1126/science.283.5401.549)
- Watkinson, S. C., Boddy, L., Burton, K., Darrah, P. R., Eastwood, D., Fricker, M. D. & Tlalka, M. 2005 New approaches to investigating the function of mycelial networks. *Mycologist* **19**, 11–17. (doi:10.1017/S0269915X05001023)
- Yeung, N., Botvinick, M. M. & Cohen, J. D. 2004 The neural basis of error detection: conflict monitoring and the error-related negativity. *Psychol. Rev.* **111**, 931–959. (doi:10.1037/0033-295X.111.4.939)
- Yu, A. & Dayan, P. 2005 Uncertainty, neuromodulation and attention. *Neuron* **46**, 681–692. (doi:10.1016/j.neuron.2005.04.026)