Dynamic decision policy reconfiguration under outcome uncertainty

Krista Bond, Kyle Dunovan, Alexis Porter, Jonathan Rubin, Timothy Verstynen

1Department of Psychology, Carnegie Mellon University, United States; 2Center for the Neural Basis of Cognition, United States; 3Carnegie Mellon Neuroscience Institute, United States; 4Department of Psychology, Northwestern University, United States; 5Department of Mathematics, University of Pittsburgh, United States; 6Department of Biomedical Engineering, Carnegie Mellon University, United States

Abstract  Humans and other mammals flexibly select actions under noisy and unstable conditions. To shed light on the mechanism driving this flexibility, we evaluated how the underlying decision policy evolves when humans change their minds about the most rewarding action. Participants performed a dynamic variant of the two-armed bandit task that manipulated the certainty in relative reward probabilities (conflict) and the reliability of action-outcome contingencies (volatility). We found that conflict and volatility contributed to shifts in exploratory states by changing both the rate of evidence accumulation (drift rate) and the amount of evidence needed to make a decision (boundary height). Following a switch in the optimal choice, the drift rate and the boundary height reduce, allowing variability in the accumulation process to predominate action selection, leading to a fast exploratory state. These changes facilitate the discovery of the new optimal choice, with a quick recovery of the boundary height to baseline. In parallel, the drift rate gradually returns to its asymptotic value as the belief in the value of the optimal choice stabilizes. Together, these decision dynamics suggest that, in the context of volatile two-choice decisions, humans adopt a combined information-threshold and drift rate mechanism in response to environmental changes. Unlike previous observations, we found no evidence that fluctuations in norepinephrine, as measured by pupillometry, associated with this adaptive shift toward an exploratory policy. We conclude that the multifaceted processes underlying a decision can rapidly reconfigure to adapt action selection policy under multiple forms of environmental uncertainty.

Introduction  Humans are faced with uncertain choices on a daily basis. Should someone buy coffee from the cafe that serves their favorite roast or try the new cafe that opened down the street? If their favorite drink is bitter one day, is that a sign to switch to a new blend or is one subpar experience inadequate to prompt a switch? Ultimately, these decisions converge to a single dilemma: whether we choose an action that we believe is likely to produce desirable results (i.e., exploit) or risk choosing another action that is less certain, on the chance that it will produce a more positive outcome (i.e., explore) (O’Reilly, 2013).

Navigating the exploration-exploitation dilemma depends on the environmental context. To
illustrate this, Figure 1A shows what happens when a simple reinforcement learning (RL) agent tries
to maximize reward in a dynamic variant of the two-armed bandit task (Sutton and Barto (1998); see Methods). Here, the relative difference in reward probability for the two actions (conflict) and the frequency of a change in the optimal action (volatility) was independently manipulated. For each level of conflict and volatility, a set of tabular Q-learning (Sutton and Barto, 1998) agents played the task with learning rate held constant while the degree of randomness of the selection policy ($\rho$ in a Softmax function) varied. The agent that returned the most reward was identified as the agent with the best exploration-exploitation balance. Increasing either form of uncertainty led to selecting agents with more random or exploratory selection policies (Fig. 1A).

As the value of the optimal choice decreases relative to the value of a suboptimal choice (conflict increases), the learner tends to exploit what she already knows. But as action values grow unstable (volatility increases) when the clarity of the optimal choice is constant (constant conflict), the learner is biased toward exploration (Bland et al., 2012). As these two forms of uncertainty change together, the gradient of action selection strategy also changes. These simulations highlight the multivariate contributions to optimal action selection policy.

Less certain environments prompt a more frequent change of mind regarding the best option, and thus they are conducive to more exploratory decision policies. How do underlying decision processes change to reflect this shift toward exploration after detecting a possible change in the optimal choice? One popular set of algorithms for describing the dynamics of decision making are accumulation-to-bound processes like the drift-diffusion model (DDM; Ratcliff (1978)). The normative form of the DDM proposes that a decision between two choices is described by a noisy accumulation process that drifts towards one of two decision boundaries at a specific rate (Fig. 1B). Two parameters of this model are critical in determining the degree of randomness of a selection policy: the rate of evidence accumulation (drift rate; $v$) and the amount of information required to make a decision (boundary height; $a$). Decreasing both drift rate and boundary height leads to more random decisions (Fig. 1C), however, the speed of these decisions depends on the ratio of the two parameters (Fig. 1D). Thus exploratory policies can result in either fast or slow decisions, depending on the relative configuration of drift rate and boundary height.

Previous modeling work has shown that these properties of a DDM process can be modulated by feedback signals and choice history (Pedersen et al., 2017; rat, ???; Dunovan and Verstynen, 2019; Dunovan et al., 2019; Mendonça et al., 2020; Urai et al., 2018) with different mechanisms for adapting the drift rate and the boundary height. When the statistics of sensory signals are equivalent for all actions, drift rate fluctuations appear to track the relative value of an action or the value difference between actions (Dunovan et al., 2019; Mikhael and Bogacz, 2016; Bariselli et al., 2018; Rubin et al., 2020). In contrast to value estimation, selection errors have been linked to changes in the boundary height (Forstmann et al., 2008, 2010; Bogacz et al., 2010; Herz et al., 2016a, 2017; Dunovan et al., 2019; Dunovan and Verstynen, 2019) and internal estimates of environmental change (Nassar et al., 2010; Wilson and Niv, 2012; Nassar et al., 2012; Behrens et al., 2007). Given the adaptive sensitivity of the drift rate and the boundary height to value estimation and selection, respectively, these decision parameters may define unique states on a manifold of fast or slow and exploratory or exploitative decision policies. These policies, in turn, adaptively reconfigure based on current environmental feedback signals by modulating value estimation and the rate of selection errors (Fig. 1E). Agents can move along the decision manifold from exploitative states (bright colors, Fig. 1E) to different types of exploratory states (darker colors, Fig. 1E) as they commit a greater number of selection errors prompted by change in action-outcome contingencies. As the system relearns properties of the environment, the decision policy migrates along the manifold to return to an exploitative state until a change occurs again.

One plausible mechanism for this migration along the manifold of selection policies is the locus coeruleus norepinephrine (LC-NE) system, which has been linked to adaptive behavioral variability in response to uncertainty (Urai et al., 2017; Dayan and Yu, 2006; Bouret and Sara, 2005). The LC-NE system has two distinct modes (Aston-Jones and Bloom, 1981) that map onto distinct decision
A) The degree of conflict and volatility shifts the optimal balance between exploration and exploitation. B) The drift diffusion model. C) Accuracy (probability that left choice selected is selected; P(L)) as a function of coordinated changes in the rate of evidence accumulation (v) and the amount of information needed to make a decision, or the boundary height (a). D) Reaction time as a function of changes in the rate of evidence accumulation and the boundary height. E) Decision policy reconfiguration.

Figure 1. A) The degree of conflict and volatility shifts the optimal balance between exploration and exploitation. B) The drift diffusion model. C) Accuracy (probability that left choice selected is selected; P(L)) as a function of coordinated changes in the rate of evidence accumulation (v) and the amount of information needed to make a decision, or the boundary height (a). D) Reaction time as a function of changes in the rate of evidence accumulation and the boundary height. E) Decision policy reconfiguration.

states (Aston-Jones and Cohen, 2005). In the phasic mode, a burst of LC activity results in a global, temporally precise release of NE. This increases the gain on cortical processing and encourages exploitation. In the tonic mode, NE is released without the temporal precision of the phasic mode, increasing baseline NE (Aston-Jones and Bloom, 1981). This encourages disengagement from the current task and facilitates exploration. The dynamic fluctuation of these two modes is thought to optimize the trade-off between the exploitation of stable sources of reward and the exploration of potentially better options (Aston-Jones and Cohen, 2005). Thus the LC-NE system, which can be indirectly measured by fluctuations in pupil diameter (Aston-Jones and Cohen, 2005; Jepma and Nieuwenhuis, 2011), may be a central mechanism for modulating selection policies.

Here we investigated the dynamics of decision policy changes in response to different types of outcome uncertainty in a series of experiments. To shift the greediness of the selection policy, we varied the ambiguity in optimal choice (conflict) and the instability of action outcomes (volatility) as human participants performed a dynamic variant (Daw et al., 2006) of the two-armed bandit task (Sutton and Barto, 2018). We predicted that, in response to suspected changes in action-outcome relationships, humans would exhibit a stereotyped adjustment in the drift rate and boundary height, that changes in response to different types of uncertainty. In addition, using pupillary data, we explored whether the LC-NE system covaried with the decision dynamics that we previously observed. We predicted that fluctuations in norepinephrine would shift the boundary height in response to a change in action outcomes to facilitate exploration.
Results

Across two experiments we used a two-armed bandit task with equivalent sensory uncertainty between targets to independently manipulate the reward conflict and the volatility of action outcomes in order to measure how underlying decision processes responded to environmental perturbations (see Stimuli and Procedure). Both of these experiments shared a common feedback structure. Participants were asked to select either the left or right target presented on the screen using the corresponding key on a response box. Rewards were probabilistically determined for each target and, if a reward was delivered, it was sampled from a Gaussian distribution. The optimally rewarding target delivered reward with a predetermined probability (P(optimal)) and the suboptimal target gave reward with the inverse probability (1 − P(optimal)). After a delay determined by the rate parameter of a Poisson distribution (λ), the reward probabilities for the optimal and suboptimal targets would switch.

In Experiment 1, we sought to establish the effect of conflict and volatility on latent decision processes. Twenty-four participants completed four sessions (high and low conflict; high and low volatility) composed of 600 trials each. During each session, they were asked to select one of two coin boxes (Exp. 1: Fig. 2A). The levels of conflict and volatility for all four conditions in Experiment 1 are shown as gray dots in Fig. 2C. Experiment 2 was an out-of-set validation of the effects that we observed in Experiment 1. In Experiment 2, participants were asked to choose between one of two Greebles (one male, one female). Each Greeble probabilistically delivered a monetary reward (Exp. 2: Fig. 2B). Participants were trained to discriminate between male and female Greebles prior to testing to prevent errors in perceptual discrimination from interfering with selection on the basis of value estimation. Four participants completed nine sessions composed of 400 trials each, generating 3600 trials in total per subject. The levels of conflict and volatility for all nine conditions in Experiment 2 are shown as black dots in Fig. 2C. Importantly, Experiment 2 manipulated the same forms of uncertainty as Experiment 1, but had different perceptual features and more expansively sampled the space of conflict and volatility. Given the similarity in design, the results for both of these experiments are presented together below.

The influence of ambiguity and instability on speed and accuracy

We first considered the effect of our conditional manipulations on the speed and accuracy of choices for both Experiments 1 and 2. In Experiment 1, accuracy (i.e., optimal choice selection) suffered as the optimal choice grew more ambiguous, with accuracy in the low conflict condition being 1.2 times higher than what is observed in the high conflict condition (Fig. 3A; β = 1.213, 95% CI: 1.192, 1.235, z=21.36, p<2e-16). In contrast, increasing conflict had no observable impact on reaction times (Fig. 3A; β = −6.902×5, 95% CI: -0.002, 0.002, t=-0.06, p=0.951). As expected, participants also became less accurate as the instability of action outcomes (i.e. volatility) grew (Fig. 3B; β = 0.092, 95% CI: 0.077, 0.111, z=10.36, p<2e-16). Under volatile conditions, participants also took slightly longer to make a decision (β = −0.012, 95% CI: -0.015, -0.010, t=−10.80, p<2e-16); however, while this effect on reaction times was statistically reliable, the impact of volatility on reaction times was weak (increasing volatility increased reaction time by ~13 ms on average; Fig. 3B).

Experiment 2 served as a high powered test of whether the effects we observed in Experiment 1 were replicable. In this experiment, we found similar effects of conflict and volatility on accuracy as we observed in Experiment 1. Accuracy increased as conflict decreased (i.e. as the probability of reward increased; β=1.312, 95% CI=1.259,1.368, z=12.752, p<2e-16). As the environment grew less volatile, accuracy increased (β=1.131, 95% CI=1.085,1.179, z=5.798, p=6.72e-09). Unlike in Experiment 1, reaction times in Experiment 2 reliably increased with the probability of reward (β=0.022, 95% CI=0.021,0.024, t=23.430, p<2e-16). Reaction time also decreased as volatility increased (i.e., as epoch length decreased; Fig. 3D; β=0.006, 95% CI=0.004,0.008, t=5.966, p=2.49e-09).

In light of the data from both of these experiments, we conclude that increasing the ambiguity of the optimal choice (conflict) and increasing the instability of action outcomes (volatility) decreases the probability of selecting the optimal choice. We observed inconsistent effects of conflict and
Figure 2. Task and uncertainty manipulation. A) In Experiment 1, participants were asked to choose between one of two “mystery boxes”. The point value associated with a selection was displayed above the chosen mystery box. The sum of points earned across trials was shown to the left of a treasure box on the upper right portion of the screen. B) In Experiment 2, participants were asked to choose between one of two Greebles (one male, one female). The total number of points earned was displayed at the center of the screen. C) The manipulation of con/uniFB02ict and volatility for Experiments 1 (gray) and 2 (black). Each point represents the combination of degrees of conflict and volatility. Under high conflict, the probability of reward for the optimal and suboptimal target is relatively close. Under high volatility, a switch in the identity of the optimal target selection is relatively frequent.

Tracking estimates of action value and environmental volatility

We calculated trial-by-trial estimates of two ideal observer parameters of environmental states that varied as a function of our manipulations of conflict and volatility (see Cognitive model for calculation details; Nassar et al. (2010); Vaghi et al. (2017)). Belief in the value difference ($\Delta B$) reflects the difference between the learned values of the optimal and suboptimal targets. For ease of interpretation, we refer to the complement of belief as doubt, such that when belief decreases doubt increases. We predicted that $\Delta B$ would capture the degree of conflict that participants estimated, with the difference in belief between these choices decreasing as conflict increases and increasing as conflict decreases. To capture the estimated probability of fundamental shifts in action values, we calculated how often the same action gave a different reward (change point probability; $\Omega$). We predicted that change point probability would capture the degree of volatility that participants estimated, increasing with the frequency of a switch in the identity of the optimal choice. We used the data from Experiment 1 to assess how well these learning estimates captured our imposed manipulations, and observed similar results in Experiment 2 (Supp. Fig. 1).

Consistent with our assumption that increasing conflict for the optimal choice should increase the doubt in the value of that choice, in Experiment 1 we observed a sharp decrease in $\Delta B$ after a switch in action outcomes and a gradual return to asymptotic values (Fig. 4A) with a decreased difference in reward probability resulting in increased doubt ($\Delta B$; $\hat{\beta} = 0.216,95\%\ CI:0.206, 0.224, t=46.24, p<2e^{-16}$). We found that both high and low volatility conditions resulted in a similar time course of belief updating relative to a switch in action outcomes (Fig. 4A). As expected, stable conditions allowed the learner to more fully update her belief in the value of the optimal choice over all trials ($\hat{\beta} = 0.058,95\%\ CI:0.050, 0.068, t=12.32, p<2e^{-16}$), though to a smaller degree than...
Figure 3. Behavior. A) Mean accuracy and reaction time for the manipulation of conflict in Experiment 1. B) Mean accuracy and reaction time for the manipulation of volatility in Experiment 1. Each point represents the average for a single subject. The distribution to the right represents the bootstrapped uncertainty in the mean difference between conditions (high conflict or high volatility subtracted from low conflict or low volatility). Distributions with 95% CIs that do not encompass 0 are marked with an asterisk. C) Mean accuracy for Experiment 2. Each purple line represents a subject. The black line represents the mean accuracy calculated across subjects. D) Reaction time distributions for each subject for Experiment 2. The black line represents the mean reaction time calculated over subjects. Error bars indicate a bootstrapped 95% confidence interval. For panels C and D, λ values shown above each plot specify the average period of optimal choice stability and the probability of reward shown on the x-axis specifies the degree of conflict.

low conflict conditions allowed (see Fig. 4B). Increasing volatility resulted in a sharp increase in the estimate of $\Omega$ at the onset of a change point with a quick return to a baseline estimate of change (Fig. 4C). Notably, this estimate of $\Omega$ was more sensitive to change points when conditions were relatively volatile, with a more pronounced peak in response to a change under high volatility conditions than under low volatility conditions (Fig. 4C). Correspondingly, over all trials, $\Omega$ was higher under more volatile conditions (Fig. 4D; $\beta = -0.022$, 95% CI: -0.023 to -0.020, $t = -30.74$, $p < 2e-16$) indicating sensitivity to the increased frequency of action outcome switches in the reward schedule.

When the identity of the optimal choice was clear (i.e. when conflict was low), the estimate of $\Omega$ was more sensitive to the presence of a true change point than when the optimal choice was ambiguous (i.e. when conflict was high) (Fig. 4C, D). This observation is consistent with the idea that increasing the difficulty of value estimation and, thereby, the assignment of value to a given choice also impairs change point sensitivity. Interestingly, increasing conflict nevertheless resulted in a net increase in $\Omega$ (Fig. 4D; $\beta = -0.006$, 95% CI: -0.007 to -0.004, $t = -8.64$, $p < 2e-16$), likely because higher conflict conditions increased the baseline estimate of change instead of enhancing sensitivity to true change points (see change point response and relative baseline values for the high conflict condition in Fig. 4C). Here, the system conservatively over-estimates the volatility of action outcomes, assuming a slightly greater frequency of changes in the probability of reward for the optimal choice than we imposed (actual proportion of change points for high conflict condition: $0.041 \pm 0.004$; estimated $\Omega$: $0.105 \pm 0.014$).

Reassuringly, net change point probability was much greater when change points were indeed
Figure 4. Changes in ideal observer estimates as a function of condition for Experiment 1. A) Changes in the belief in the value of the optimal target ($\Delta B$) as a function of conflict and volatility over time. B) Mean belief in the value of the optimal choice across conditions. C) Changes in change point probability ($\Omega$) as a function of conflict and volatility over time. D) Mean change point probability across conditions. Error bars represent 95% CIs.

more frequent (see increased $\Omega$ estimates for high volatility conditions over high conflict conditions in Figure 4D). These results suggest that our formulation of these ideal observer estimates adequately captures our manipulation of volatility and conflict at a continuous level.

We now consider the link between these internal estimates of uncertainty, $\Delta B$ (or doubt, the inverse of $\Delta B$) and $\Omega$ and the behavioral effects we observed in Experiments 1 and 2. In Experiment 1, we found no substantial association between doubt and reaction time ($\beta = -0.003$, 95% CI: -0.005, 0.000, $t=-2.45$, $p=0.015$). Similarly, a one standard deviation increase in $\Omega$ did not reliably relate to reaction time to a meaningful degree ($\beta = 0.002$, 95% CI: 0.000, 0.005, $t=2.15$, $p=0.032$). Further, $\Omega$ and $\Delta B$ did not interact to affect reaction time ($\beta = -0.001$, 95% CI: -0.003, 0.001, $t=-1.24$, $p=0.215$). These results are consistent with the null effects of our conditional manipulations on reaction time (see The influence of ambiguity and instability on speed and accuracy). However, $\Omega$ and $\Delta B$ did interact to affect accuracy ($\beta = 0.916$, 95% CI: 0.905, 0.927, $z=-14.51$, $p<2e-16$), with the effect of belief on accuracy decreasing as the probability of a change point increased, consistent with the conditional effects of conflict and volatility decreasing the accuracy of selection.

We found similar results in Experiment 2. Accuracy increased as doubt decreased, or as the belief in the value of the optimal choice increased (Fig. 3C; $\beta=1.156$, 95% CI=1.126,1.186, $z=10.884$, $p<2e-16$). As the estimate of $\Omega$ increased, accuracy decreased (Fig. 3C; $\beta=0.899$, 95% CI=0.869,0.930, $z=6.220$, $p=4.97e-10$). As observed in our analysis of the behavioral data in terms of our conditional manipulations, reaction times in Experiment 2 reliably increased to a small degree with $\Delta B$ (Fig. 3D; $\beta=0.004$, mean 95% CI=0.002,0.005, $t=6.028$, $p=1.7e-09$), unlike in Experiment 1. Consistent with our previously observed effect of volatility on reaction time in Experiment 2, reaction time decreased as $\Omega$ increased ($\beta=-0.004$, 95% CI=-0.005,-0.002, $t=-4.716$, $p=2.43e-06$), although this decrease in reaction time was only on the order of 4 ms for a one standard deviation increase in $\Omega$, on average.
Across experiments, we find that the relationship between ideal observer estimates and behavior mirrors the relationship between our conditional manipulations and behavior. Specifically, selection accuracy increased with the belief in the value of the optimal choice in both experiments, with an increase in the estimate of change point probability decreasing with the probability of selecting the optimal choice. As before, we observed inconsistent effects of the belief in the value of the optimal choice and change point probability on reaction times across experiments. While neither of these internal estimates of uncertainty had a reliable effect on reaction times in Experiment 1, Experiment 2 suggests a slight but statistically reliable increase in reaction time with the belief in the value of the optimal choice and a decrease in reaction time as the probability of a change increased.

**Different forms of uncertainty impact distinct decision processes**

Having evaluated how optimal choice ambiguity and unstable action outcomes affected gross behavior, we next moved on to how the underlying decision policy responded to these sources of uncertainty. To this end, we fit the behavioral data from Experiment 1 to versions of the adaptive DDM (Table 1, left panel) in which the ideal observer parameters explained trial-wise variability in either the rate of information accumulation (\( \omega \)), the amount of information needed to select an action (\( \alpha \)), or both.

Recall that we used the ideal observer parameters \( \Delta B \) and \( \Omega \) as proxies for internal estimates of belief in the difference in learned target values and change point probability, respectively. This provided a continuous quantification of our manipulation of conflict and volatility (see Tracking estimates of action value and environmental volatility). Experiment 2 provided a high powered, within-subject test of whether the mapping between decision processes and ideal observer estimates that we observed in Experiment 1 would replicate. To select the model that best accounted for the data, we compared the deviance information criterion (DIC) scores (Spiegelhalter et al., 2002) for various models where the ideal observer parameters for environmental state (\( \Delta B \), \( \Omega \)) covary with different parameters in the DDM process (Table 1). DIC scores provide a measure of model fit adjusted for model complexity and quantify information loss. A lower DIC score indicates a model that loses less information. Here, a difference of \( \leq 2 \) points from the lowest-scoring model cannot rule out the higher scoring model; a difference of 3 to 7 points suggests that the higher scoring model has considerably less support; and a difference of 10 points suggests essentially no support for the higher scoring model (Spiegelhalter et al., 2002; Burnham and Anderson, 1998).

Using the data from Experiment 1, we found that the model mapping \( \Delta B \) to \( \omega \) and \( \Omega \) to \( \alpha \) provided the best model fit to the data given the models considered (minimum \( \Delta_{\text{DIC}} = 13.1 \)) and fit the observed data substantially better than the null, intercept-only model (\( \Delta_{\text{DIC}} = -2698 \), see left panel of Table 1), which predicts the decision parameters (\( \alpha \) and \( \omega \)) as a function of the mean values of those parameters. This suggests that as the belief in the value of the optimal choice approaches the reward value for the optimal choice, the rate of information accumulation increases. An internal estimate of change point probability also updates the amount of information required to make a decision. To test whether the effects we observed in Experiment 1 were replicated in an independent data set, we performed a similar model comparison procedure in Experiment 2. Because Experiment 2 followed a replication-based design, we fit a separate model to each subject to assess the replicability of the best fitting model from Experiment 1. For the sake of simplicity, we took the best fitting model from Experiment 1 (Model I) and compared this to the alternative model where the mapping between the ideal observer parameters and the DDM parameters is switched (Model II). This holds the complexity of our hypothesized and alternative model constant.

In three out of four participants in Experiment 2, we replicated the finding that the best model for explaining trialwise variability in DDM parameters was Model I (Table 1, right panel), with the DIC value for this model lower than the inverted model by an average of 31.8 points. Considering the model selection results from both experiments, we find robust evidence for an estimate of
We found that the rate of evidence accumulation, \( v \), increases with the belief in the value of the optimal choice, \( \Delta B \), and \( \Omega \), reliably affected the underlying decision processes, \( v \) and \( a \), we examined the posterior distributions of the regression coefficients for the model supported across experiments (see Hierarchical drift diffusion modeling). We considered the relationship to be reliable if the estimated coefficient maintained the same sign over at least 95% of the mass of the posterior distribution. For Experiment 2, we performed the same analyses of the posterior distributions of regression coefficients and computed the mean and standard deviation of both the reliability of the estimates and the magnitude of the effect over subjects. For Experiment 1, both the relationship between \( a \) and \( \Omega \) and the relationship between \( v \) and \( \Delta B \) were statistically reliable (observed \( p(\beta_{\Delta B} > 0) = 0.002 \); observed \( p(\beta_{v,\Delta B} < 0) < 0.001 \); see Supp. Fig. 2 for raw posterior distributions for Experiment 1).

Next, we asked how these internal estimates of uncertainty updated the decision processes. We found that the rate of evidence accumulation, \( v \), increased with the belief in the value of the optimal choice, \( \Delta B \) (\( \beta_{v,\Delta B} = 0.404 \), empirical 95% CI: 0.390, 0.418). Interestingly, we also observed that the amount of evidence needed to make a decision decreased as the probability of a change point increased (\( \beta_{v,\Delta B} = -0.035 \), 95% CI: -0.057, -0.014).

Experiment 2 replicated the positive relationship between \( \Delta B \) and \( v \) (mean observed \( p(\beta_{v,\Delta B} < 0) = 0.003 \pm 0.005 \)), with the drift rate increasing as the belief in the value of the optimal choice increased (mean \( \beta_{v,\Delta B} = 0.112 \pm 0.061 \)), though to a lesser degree than we observed originally (see Supp. Fig. 3 for raw posterior distributions for Experiment 2). The effect of \( \Omega \) on \( a \) was less reliable than we observed in Experiment 1 (mean observed \( p(\beta_{a,\Omega} < 0) = 0.727 \pm 0.108 \)) though the magnitude of the effect was on par with our previous observations (mean \( \beta_{a,\Omega} = -0.037 \pm 0.017 \), with the boundary height decreasing as the probability of a change point increased.

Across experiments, we find a consistent relationship between internal estimates of environmental uncertainty and the dynamics of the underlying decision processes. Altogether, these results suggest a combined information-threshold and drift rate mechanism for adaptation to change (Fig. 5A). When a change point is detected, the threshold for committing to a choice (\( a \)) decreases, potentiating a quick response. At the same time, the rate of evidence accumulation (\( v \)) approaches zero, allowing the variability in the decision process to dominate, encouraging a random selection. As the learner accrues information about the new optimal choice, the rate of information accumulation slowly recovers to asymptotic levels, with the decision process assuming a more directed path toward the choice that has accrued evidence for reward. Together, these independent, but cooperative, changes in the underlying decision processes point to a mechanism.

### Table 1. Model comparison for Experiments 1 and 2. Roman numerals refer to a given model, as defined by the mapping between the ideal observer estimates and decision parameters in the first two columns. The left panel shows the deviance information criterion (DIC) scores for the set of models considered during the model selection procedure for Experiment 1. The right panel shows the DIC scores for the hypothesized mapping and the inverse mapping, with each set of models representing the fit for each of four subjects. The column labeled DIC gives the raw DIC score, \( \Delta \text{DIC}_{\text{null}} \) lists the change in model fit from an intercept-only model (the null-adjusted fit), and \( \Delta \text{DIC}_{\text{null}} \) provides the change in null-adjusted model fit from the best-fitting model. The best performing model is denoted by an asterisk.

| Experiment 1 | Experiment 2 |
|--------------|--------------|
| \( \Delta B \) | \( \Delta B \) |
| \( \Omega \) | \( \Omega \) |
| DIC | DIC |
| \( \Delta \text{DIC}_{\text{null}} \) | \( \Delta \text{DIC}_{\text{null}} \) |
| \( \Delta \text{DIC}_{\text{null}} \) |

| Experiment 1 | Experiment 2 |
|--------------|--------------|
| \( \Delta B \) | \( \Delta B \) |
| \( \Omega \) | \( \Omega \) |
| DIC | DIC |
| \( \Delta \text{DIC}_{\text{null}} \) | \( \Delta \text{DIC}_{\text{null}} \) |
| \( \Delta \text{DIC}_{\text{null}} \) |

| Model | Experiment 1 | Experiment 2 |
|-------|--------------|--------------|
| I     | \( a \) v   | \( a \) v   |
| II    | \( a \) v   | \( a \) v   |
| III   | \( a \) v   | \( a \) v   |
| IV    | \( a \) v   | \( a \) v   |
| V     | \( a \) v   | \( a \) v   |
| VI    | \( a \) v   | \( a \) v   |
| VII   | \( a \) v   | \( a \) v   |
| \( B \) | \( B \) | \( B \) |
for quickly gathering information in an unbiased manner when the learner needs to update her
evaluation. We now explore these dynamics in more detail.

Environmental instability prompts a stereotyped decision trajectory
So far we have established that both the drift rate and the boundary height are independently
manipulated by two different estimates of environmental uncertainty with different temporal dy-
namics. This suggests that a change in action-outcome contingencies prompts a unique trajectory
through the space of possible decision policies (Fig. 1E). To visualize this trajectory, we plot the
temporal relationship between drift rate and boundary height beginning with the trial prior to the
change point and ending three trials after the change point (Fig. 5B). Following a shift in response
contingencies, the navigation of this decision manifold follows a stereotyped pattern. The bound-
ary height ($a$) and rate of evidence accumulation ($v$) initially decrease rapidly (see the trial prior to
the change point in Fig. 5B) and then gradually recover and stabilize over time. This decision tra-
jectory is consistent across subjects (Fig. 5B, right panel; see Supp. Fig. 4 for decision trajectories
from Experiment 1).

To capture the consistency of this change-point driven response in the relationship between $a$
and $v$ over time, we also represented the trialwise shift in these two decision variables as vectors.
The trial-by-trial estimates of drift rate and boundary height were calculated from the best-fit model
and z-scored (see Different forms of uncertainty impact distinct decision processes for model se-
lection). Then the difference between each sequential set of $(a, v)$ coordinates was calculated to
produce a vector length. The arc tangent between these differenced values was computed to yield
an angle in radians between sequential decision vectors, concisely representing the overall deci-
sion dynamics ($\theta$, Fig. 5B, see Decision vector representation for methodological details). We find
that the distribution of $\theta$ prior to a change point averages to $\sim 135$ deg, sharply changes in response
to the observation of a change point ($\sim 270$ deg) and steadily returns to values prior to the onset
of a change (Fig. 5C; see Supp. Fig. 5 for vector distributions). One trial after the change point, $v$
sharply decreases along with $a$, after which $a$ quickly recovers and $v$ steadily progresses toward its
baseline value. Altogether, we find that the decision trajectory quickly responds to a shift in action
outcomes and also quickly recovers and stabilizes.

Having characterized the change-point-evoked trajectory through the manifold of decision poli-
cies, we next asked whether conditions of increased volatility and increased conflict might modify
its path. We conducted a comparison of null models and models specifying an influence of conflict
and volatility using circular regression (Mulder and Klugkist, 2017). We began our model compar-
ison with two null hypotheses. The first null model estimated decision dynamics (the relationship
between $a$ and $v$; $\theta$) as a function of the intercept, or the average of the decision dynamics $\theta$:

$$\theta = \beta_0$$

We call this the absolute null model.

Our second null model estimated decision dynamics ($\theta$) solely as a function of time, with the
time scale of consideration determined by the results of a stability analysis from Experiment 1 (see
Model proposals and evaluation; Supp. Fig. 6):

$$\theta = \beta_0 + \beta_{t,1.5}$$

We call this the time-null model.

Our model comparison logic was as follows. We first evaluated whether the posterior probabil-
ity of the time-null model was greater than for the absolute null model. This would suggest that
time was a better predictor of decision dynamics than the average response. If the 95% CI of the
posterior probability for the time-null model exceeded the 95% CI of the absolute null model, we
quantified the evidence for alternative models relative to the time-null model, with the sole effect
of time as the new null hypothesis.
Figure 5. The decision manifold. A) Changes in ideal observer estimates of uncertainty over time and their effect on the boundary height and the drift rate. Directly after a change point, the boundary height decreases and the drift rate slows. Over time, the boundary height returns to its baseline value and the drift rate increases. B) Representing decision space in vector form. An angle was calculated between sequential values of ($\omega$, $v$) coordinates, beginning with the trial prior to the change point. The right-hand side shows the decision trajectories for all participants. Note that these trajectories are z-scored. C) Distributions depicting the angle between drift rate and boundary height. The area of the shaded region is proportional to the density and the arrow represents the circular mean.

We used Bayes Factors to quantify the ratio of evidence for two competing hypotheses. If the ratio is close to 1, then the evidence is equivocal. As the ratio grows more positive, there is greater evidence for the model specified in the numerator, and if the ratio is less than 1, then there is evidence for the model specified in the denominator (Jeffreys, 1998). Evidence for the null hypothesis is denoted $B_{F_{01}}$ and evidence for the alternative hypothesis is denoted $B_{F_{10}}$. Because Experiment 2 took a replication-based approach, a separate model was fit for each participant for all proposed models.

To determine whether perturbations of volatility and conflict affected decision dynamics, we modeled decision dynamics ($\theta$) as a function of $\lambda$ and $p$, where $\lambda$ corresponds to the average period of stability and $p$ corresponds to the mean probability of reward for the optimal choice (see Fig. 6 for the full set of models considered). We explored the potential influence of volatility and conflict on the relationship between drift rate and boundary height ($\theta$) by examining the posterior probability for each hypothesized model given the set of alternative hypotheses (Model proposals and evaluation; Fig. 6A). We found that the model describing the relationship between $\theta$ and time relative to a change point was more probable than the absolute null model (Fig. 6A).

We also present the evidence for the absolute null model against each alternative model as a Bayes Factor ($B_{F_{01}}$) (Fig. 6B). The 95% confidence interval for the $B_{F_{01}}$ comparing the ratio of evidence for the absolute null model and the model specifying time-dependent effects of volatility included 1, suggesting inconclusive evidence for either of these models. Likewise, the 95% confidence interval for the $B_{F_{10}}$ comparing the evidence for the absolute null model against the model specifying time-dependent effects of conflict included 1, suggesting no substantive difference between them. Given the equivocal evidence for the two models specifying time-dependent effects of conflict and volatility relative to the absolute null model, we excluded them from further comparison with the time-null model.
Figure 6. Model comparisons for the effect of volatility and conflict on the relationship between drift rate and boundary height (θ). A) The posterior probability for models testing for an effect of volatility and conflict on θ. B) The Bayes Factor for the intercept null model relative to the alternative models specifying either an effect of time alone or a conditional effect on θ. C) The Bayes Factor for the time null model relative to the surviving alternative models specifying a conditional effect on θ. Note that time refers to time relative to the onset of a change point. All models specifying an interaction also include main effects. Dotted horizontal lines refer to grades of evidence (Wagenmakers, 2007).

The remainder of the models had substantially negative $BF_{01}$s (Fig. 6B), suggesting that they better fit the data than the absolute null model. To evaluate our second null hypothesis that time alone best accounted for the data, we computed the $BF_{01}$ for the time-null model against the surviving models from the absolute null analysis. We find that, for all of the remaining models, the $BF_{01}$ is substantially positive (Fig. 6C), indicating that the time-null model best accounted for the data (posterior probability of time-null model given the set of models considered: $0.76 \pm 0.473$; posterior prob. for 3/4 participants > 0.99).

These analyses suggest that the relationship between the rate of evidence accumulation and the boundary height is only related to time relative to a change point. We find no evidence to suggest that changing the degree of volatility or changing the degree of conflict changes the path of this decision trajectory. Thus, the trajectory through the manifold of decision policies is solely sensitive to the presence of a change point rather than either the history of change point frequency or the history of optimal choice ambiguity. This suggests that adaptation to environmental changes in action-outcome contingencies involves a rapid, coordinated decrease in both the amount of information needed to make a decision and the rate of information accumulation, with a stereotyped return to a stable baseline soon thereafter until another change occurs.
No evidence for locus-coeruleus norepinephrine (LC-NE) system contribution to the decision trajectory

The locus coeruleus-norepinephrine (LC-NE) system is known to modulate exploration states under uncertainty and pupil diameter shows a tight correspondence with LC neuron firing rate (Aston-Jones and Cohen, 2005), with changes in pupil diameter indexing the explore-exploit decision state (Jepma and Nieuwenhuis, 2011). Similar to the classic Yerkes-Dodson curve relating arousal to performance (Yerkes et al., 1908), performance is optimal when tonic LC activity is moderate and phasic LC activity increases following a goal-related stimulus (Aston-Jones et al., 1999). Because of this link between LC-NE and the regulation of behavioral variability in response to uncertainty, we expected that LC-NE system responses, as recorded by pupil diameter, would associate with environmental uncertainty and the trajectory through decision policy space following a change in action-contingencies. Specifically, if the LC-NE system were sensitive to a change in the optimal choice, then we should observe moderate a spike in phasic activity following a change in action-outcome contingencies.

We characterized the evoked pupillary response on each trial using seven metrics: the mean of the pupil data over each trial interval, the latency to the peak onset and offset, the latency to peak amplitude, the peak amplitude, and the area under the curve of the pupillary response (see Pupil data preprocessing; Fig. 7A). We then submitted these metrics to principal component analysis to reduce their dimensionality while capturing maximum variance. Evoked response characterization and principal component analyses were conducted for each session and for each subject. The 95% CI for the number of principal components needed to explain 95% of the variance in the data was calculated over subjects and sessions to determine the number of principal components to keep for further analysis. To aid in interpreting subsequent analysis using the selected principal components, the feature importance of each pupil metric was calculated for each principal component and aggregated across subjects as a mean and bootstrapped 95% CI (Fig. 7). We found that the first two principal components explained 95% of the variance in the pupillary data. Peak onset, peak offset, and latency to peak amplitude had the greatest feature importance for the first principal component (Fig. 7B, upper panel). Mean pupil diameter and peak amplitude had the greatest feature importance for the second principal component (Fig. 7B, lower panel). Thus, for interpretability, we refer to the first and second principal components as timing and magnitude.

Figure 7. Method for analyzing pupil data. A) The evoked pupillary response was characterized according to seven metrics. B) These pupillary features were submitted to a principal component analysis. The contribution of each feature to the variance explained for the first two components is plotted for each subject.

[Diagram of pupil response with metrics and principal components]
To test for the possibility that fluctuations in norepinephrine covaried with changes in the drift-rate and the boundary height, we evaluated a set of models exploring the relationship between the timing and magnitude components of the pupillary response and shifts in $\theta$. As in our previous model comparison (Fig. 6; see Environmental instability prompts a stereotyped decision trajectory), we found that the model describing the relationship between decision dynamics ($\theta$) and time relative to a change point had the highest posterior probability given the set of models considered (Fig. 8A). To further evaluate the extent of the evidence for the time-null hypothesis, we present the evidence for the time null model against the original model set as $BF_{01}$ (Fig. 8B). We find unambiguous evidence in favor of the time-null model relative to the models specifying the modulation of $\theta$ via the timing and magnitude features of the pupillary response (posterior probability of time-null model given the set of models considered: $0.997\pm0.002$), with substantially positive $BF_{01}$ values. We find no evidence that the pupillary response associates with the dynamics of the decision policy changes in response to a change in action-outcome contingencies.

### Discussion

We investigated how decision policies change when the rules of the environment change. In two separate experiments, we characterized how decision processes adapted in response to a change in action-outcome contingencies as a trajectory through the space of possible types of exploratory and exploitative decision policies. Our findings highlight how, in the context of two choice paradigms, when faced with a possible change in outcomes, humans rapidly shift to a fast exploratory strategy by reducing both the drift rate and the boundary height in a stereotyped manner. Using pupillary data, we were unable to detect a relationship between the LC-NE system and the dynamics of adaptive decision policies in unstable environments. Our findings highlight how the underlying decision algorithm adapts to different forms of uncertainty.

Exploration and exploitation states are not discrete, but exist along a continuum (Addicott et al., 2017). Instead of switching between binary states, humans manage environmental instability by adjusting the degree of exploration and exploitation (Sadeghiyeh et al., 2020; Prat-Carrabin et al., 2020; Feng et al., 2020; Wilson et al., 2014; Payzan-LeNestour and Bossaerts, 2011, 2012; Wilson et al., 2015).
Depending on the relative configuration of parameters in the accumulation to bound process, this adjustment can manifest as either speeded or slowed decisions (Fig. 1E) (Alexandrowicz, 2020; Ratcliff, 1978). Our results suggest that, in the context of volatile two-choice decisions, humans adopt a mechanism that simultaneously changes the information threshold and rate of evidence accumulation to adapt to an environmental change (Fig. 5A). As soon as a shift in action outcomes is suspected, an internal estimate of change point probability increases and an estimate of the belief in the value of the optimal target plummets (Fig. 5A). The rapid increase in change point probability causes a rapid, but slight, contraction of the boundary height on the subsequent trial, thereby lowering the criterion for selecting a new action and allowing variability in the accumulation process to have a greater influence on choice (Fig. 5B-C). These changes lead to fast exploratory decisions that facilitate discovery of the new optimal action and result in a quick recovery of the original threshold value. In parallel, the rate of evidence accumulation for the optimal choice decreases, with an immediate drop that gradually returns to its asymptotic value as the belief in the value of the optimal choice stabilizes. These results show that when a learner confronts a change point, the decision policy becomes more exploratory by simultaneously lowering the amount of evidence needed to make a decision and slowing the integration of evidence over time. Together, these decision dynamics form a mechanism for quickly gathering information in an unbiased manner.

Critically, our finding that underlying decision policies can express exploration as a relative reconfiguration of multiple underlying decision parameters closely parallels recent work in the domain of information-seeking. Information seeking has been decomposed into random and directed components (Wilson et al., 2014). Random exploration refers to inherent behavioral variability that leads us to explore other options, while directed exploration refers to the volitional pursuit of new information. Feng and colleagues recently found that random exploration is driven by changes in the drift rate and the boundary height, with drift rate changes dominating the policy shift (Feng et al., 2020). When environmental conditions encouraged exploration, the drift rate slowed, reducing the signal-to-noise ratio of the reward representation. This finding clearly aligns with our current observations showing that the drift rate sharply decreases in response to a change point and that this change in drift rate dominates the reconfiguration of decision processes, though our experiments were not designed to isolate the directed and random elements of exploration.

Our results are also broadly consistent with a growing body of research converging on the idea that decision policies are not static, but sensitive to changes in environmental dynamics (Dunovan and Verstynen, 2019; Urai et al., 2018). Previous work by our lab (Dunovan and Verstynen, 2019) has shown how, during a modified reactive inhibitory control task, different feedback signals target different parts of the accumulation-to-bound process. Specifically, errors in response timing drove rapid changes in the drift rate on subsequent trials, while selection errors (i.e., making a response on trials where the response should be inhibited) changed the boundary height. Further, there is new evidence that the drift rate adapts on the basis of previous choices, independent of the feedback given for those choices. Urai and colleagues have convincingly demonstrated that choice history signals sculpt the dynamics of the accumulation process by biasing the rate of evidence accumulation (Urai et al., 2018). Our current findings and these previous observations (Pedersen et al., 2017; rat, ????) all highlight how sensitive the parameters of accumulation-to-bound processes are to immediate experience.

At first glance, our results might appear to conflict with prior findings linking relative differences in action value to an increase in boundary height (Cavanagh et al., 2011; Frank, 2006; Frank et al., 2015). However, in contrast with the previous studies, our experiments were designed to not only include an estimate of reward-conflict, but also an estimate of change point probability. This allowed us to directly compare models capturing the effect of conflict and volatility, respectively, on the drift rate and the boundary height (Table: Fig. 1). Including both of these estimates may parse drift rate effects from those on the boundary height, which would otherwise be obscured. Importantly, the model that mapped change point probability to the drift rate and belief to the boundary
height did not explain variability in choice and reaction times as well as the model in which change
point probability drives changes in the boundary height and belief drives changes in the drift rate.
This comparison lends some credence to the possibility that previous observations of a selective
effect of conflict on the boundary height (Cavanagh et al., 2011; Frank, 2006; Frank et al., 2015)
may have been the result of an obscured relationship between conflict and volatility.

We hypothesized that these shifts in decision policies would be linked to changes in phasic
responses of the LC-NE pathways (Aston-Jones and Cohen, 2005). However, our results failed to
find such an association. This suggests that an alternative mechanism drives the adaptive changes
that we observed behaviorally. On possible alternative is the cortico-basal ganglia-thalamic (CBGT)
pathways or "loops". Both recent experimental (Yartsev et al., 2018) and theoretical (Bogacz and
Larsen, 2011; Caballero et al., 2018; Wei et al., 2015) studies have pointed to the CBGT loops as
being a crucial pathway for accumulating evidence during decision making, with the wiring architec-
ture of these pathways being ideal for implementing the sequential probability ratio test (Bogacz
and Gurney, 2007; Bogacz, 2007), the statistically optimal algorithm for evidence accumulation de-
cisions and the basis for the DDM itself (Ratcliff, 1978). Further, multiple lines of theoretical work
have suggested that, within the CBGT pathways, the difference in direct pathway activity between
action channels covaries with the rate of evidence accumulation for individual decisions (Mikhael
and Bogacz, 2016; Bariselli et al., 2018; Dunovan et al., 2019; Rubin et al., 2020), while the indirect
pathways are linked to control of the decision threshold (Wei et al., 2015; Herz et al., 2016b; Bo-
gacz, 2007; rat, ???). This suggests that changes in the direct and indirect pathways, both within
and between representations of different actions, may regulate shifts in decision policies.

Critically, these pathways are also a target of the dopaminergic signaling that drives reinforce-
ment learning (Schultz et al., 1992), suggesting that changes in relative action-value should drive
trial-by-trial changes in the drift rate. Indeed, previous work relating dopaminergic circuitry to deci-
sion policy adaptation suggests that dopamine may play a critical role in modulating decision poli-
cies. Dopamine has substantial links to exploration (Kakade and Dayan, 2002) and recent pharma-
cological evidence suggests a role for dopaminergic regulation of exploration in humans (Chakroun
et al., 2020). More explicitly, both directed and random exploration have been linked to variations
in genes that affect dopamine levels in prefrontal cortex and striatum, respectively (Gershman and
Tzovaras, 2018). Physiologically, previous work has found that a dopamine-controlled spike-timing
dependent plasticity rule alters the ratio of direct to indirect pathway efficacy in a simulated corti-
costral network (Vich et al., 2020), with overall indirect pathway activity linked to the modulation
of the boundary height in a DDM and the difference in direct pathway activation across action chan-
nels associated with changes in the drift rate (Dunovan et al., 2019; Rubin et al., 2020). Moreover,
recent optogenetic work in mice suggests that activating the subthalamic nucleus, a key node in
the indirect pathway, not only halts the motoric response but also interrupts cognitive processes
related to action selection (Heston et al., 2020). Our current observations, combined with this
previous work, suggests that the decision policy reconfiguration that we observe may associate
with similar underlying corticostriatal dynamics, with belief-driven changes to drift rate varying
with the difference in direct pathway firing rates across action channels Dunovan et al. (2019), and
change-point-probability-driven changes to the boundary height varying with overall indirect path-
way activity (Dunovan et al., 2019; Vich et al., 2020). Future physiological studies should focus on
validating this predicted relationship between decision policy reconfiguration and CBGT pathways.

Indeed, the current study raises many more questions about the dynamics of adaptive deci-
sion policies than it answers. For example, we only sparsely sampled the space of possible states
of value conflict and volatility. Future work would benefit from a more complete sampling of the
conflict and volatility space. A psychophysical characterization of how decision states shift in re-
sponse to varying forms of uncertainty will expose potential non-linear relationships between the
decision policy and feedback uncertainty. Moreover, the decisions that we have modeled here
are simple two choice decisions, constrained mostly by the normative form of the traditional DDM
framework (Ratcliff, 1978). Scaling the complexity of the task will allow for a more complete assess-
ment of how these relationships change with more complex decisions that better approximate the choices that we make outside the lab. This could be done by moving the cognitive model to frameworks that can fit processes for decisions involving more than two alternatives (e.g. Tajima et al. (2019)). Finally, because our estimate of the relationship between our ideal observer estimates of uncertainty and human estimates of uncertainty was indirect, this work would benefit from online approximations of ideal observer estimates, as has been done previously (Wilson et al., 2010). Indeed, there can be substantive individual differences in the detection of of change points (Wilson et al., 2010). Thus, an approximation of how well the estimates of change point probability from our ideal observer correspond to estimates that human observers hold is needed. This approximation would validate the fidelity of the relationship between the ideal observer estimates of uncertainty and the decision parameters that we observed.

Together, our results suggest that when humans are forced to change their mind about the best action to take, the underlying decision policy adapts in a specific way. When a change in action-outcome contingency is suspected, the rate of evidence accumulation decreases and less evidence is required to commit to a response, allowing variability inherent to the decision process to play a greater role in response selection and resulting in a fast exploratory state. As the environment becomes stable, the system slowly adapts to an exploitative state. Importantly, we find no evidence that norepinephrine pathways associate with this response. This suggests that other pathways may be engaged in this adaptive reconfiguration of decision policies. These results reveal the multifaceted underlying decision processes that can adapt action selection policy under multiple forms of environmental uncertainty.

Methods and Materials

Participants

Twenty-four participants (19 female, 22 right-handed, 19-31 years old) were recruited for Experiment 1 and paid $20 at the end of four sessions. Four participants (2 female, 4 right-handed, 21-28 years old) were recruited for Experiment 2 and paid $10 for each of nine sessions, in addition to a performance bonus. All participants were sourced from the Carnegie Mellon University community. Both experiments were approved by the Institutional Review Board at Carnegie Mellon University.

Processed data and code are available within a Github repository for this publication. Hypotheses were registered prior to the completion of data collection using the Open Science Framework (Foster and Deardorff, 2017). It should be noted that the direction of the association between change point probability and boundary height was inverted in the preregistration.

Stimuli and Procedure

Experiment 1

To begin the task, each participant read the following instructions:

"You're going on a treasure hunt! You will start with 600 coins in your treasure chest, and you'll be able to pay a coin to open either a purple or an orange box. When you open one of those boxes, you will get a certain number of coins, depending on the color of the box. However, opening the same box will not always give you the same number of coins, and each choice costs one coin. After making your choice, you will receive feedback about how much money you have. Your goal is to make as much money as possible. Press the green button when you're ready to continue. Choose the left box by pressing the left button with your left index finger and choose the right box by pressing the right button with your right index finger. Note that if you choose too slowly or too quickly, you won't earn any coins. Finally, remember to make your choice based on the color of the box. Press the green button when you're ready to begin the hunt!"
On each trial, participants chose between one of two ‘mystery boxes’ presented side-by-side on the computer screen (Fig. 2A). Participants selected one of the two boxes by pressing either a left button (left box selection) or right button (right box selection) on a button box (Black Box ToolKit USB Response Pad, URP48). Reaction time (RT) was defined as the time elapsed from stimulus presentation to stimulus selection. Reaction time was constrained so that participants had to respond within 100 ms to 1000 ms from stimulus presentation. If participants responded too quickly, the trial was followed by a 5 s pause and they were informed that they were too fast and asked to slow down. If participants responded too slowly, they received a message saying that they were too slow, and were asked to choose quickly on the next trial. In both of these cases, participants did not receive any reward feedback or earn any points, and the trial was repeated so that 600 trials met these reaction time constraints. In order to avoid fatigue, a small break was given midway through each session (break time: 0.70±1.42 m). Participants began each condition with 600 points and lost one point for each incorrect decision.

Feedback was given after each rewarded choice in the form of points drawn from the normal distribution $N(\mu = 3, \sigma = 1)$ and converted to an integer. If the choice was unrewarded, then participants received 0 points. These points were displayed above the selected mystery box for 0.9 s. To prevent stereotyped responses, the inter-trial interval was sampled from a uniform distribution with a lower limit of 250 ms and an upper limit of 750 ms ($U(250, 750)$). The relative left-right position of each target was pseudorandomized on each trial to prevent incidental learning based on the spatial position of either the mystery box or the responding hand.

To induce decision-conflict, the probability of reward for the optimal target ($P$) was manipulated across two conditions. We imposed a relatively low probability of reward for the high conflict condition ($P = 0.65$). Conversely, we imposed a relatively high probability of reward for the low conflict condition ($P = 0.85$). For all conditions, the probability of the low-value target was $1 - P$.

Along with these reward manipulations, we also introduced volatility in the action-outcome contingencies. After a prespecified number of trials, the identity of the optimal target switched periodically. The point at which the optimal target switched identities was termed a change point. Each period of mean contingency stability was defined as an epoch. Consequently, each session was composed of multiple change points and multiple epochs. Epoch lengths, in trials, were drawn from a Poisson distribution. The lambda parameter was held constant for both high conflict and low conflict conditions ($\lambda = 25$).

To manipulate volatility, epoch lengths were manipulated across two conditions. The high volatility condition drew epoch lengths from a Poisson distribution where $\lambda = 15$ and the low volatility condition drew epoch lengths from a distribution where $\lambda = 35$. In these conditions manipulating volatility, the probability of reward was held constant ($P = 0.75$).

Each participant was tested under four experimental conditions: high conflict, low conflict, high volatility, and low volatility. Each condition was completed in a unique experimental session and each session consisted of 600 trials. Each participant completed the entire experiment over two testing days. To eliminate the effect of timing and its correlates on reward learning (Byrne et al., 2017; Murray et al., 2009), the order of conditions was counterbalanced across participants.

Experiment 2

Experiment 2 used male and female Greebles (Gauthier and Tarr, 1997) as selection targets (Fig. 2B). Participants were first trained to discriminate between male and female Greebles to prevent errors in perceptual discrimination from interfering with selection on the basis of value. Using a two-alternative forced choice task, participants were presented with a male and female Greeble and asked to select the female, with the male and female Greeble identities resampled on each trial. Participants received binary feedback regarding their selection (correct or incorrect). This criterion task ended after participants reached 95% accuracy (mean number of trials to reach criterion: 31.29, standard deviation over means for subjects: 9.99).

After reaching perceptual discrimination criterion for each session, each participant was tested
under nine reinforcement learning conditions composed of 400 trials each, generating 3600 trials per subject in total. Data were collected from four participants in accordance with a replication-based design, with each participant serving as a replication experiment. Participants completed these sessions across three weeks in randomized order. Each trial presented a male and female Greeble (Gauthier and Tarr, 1997), with the goal of selecting the sex identity of the Greeble that was most profitable (Fig. 2B). Individual Greeble identities were resampled on each trial; thus, the task of the participant was to choose the sex identity rather than the individual identity of the Greeble which was most rewarding. Probabilistic reward feedback was given in the form of points drawn from the normal distribution \(N(\mu = 3, \sigma = 1)\) and converted to an integer, as in Experiment 1. These points were displayed at the center of the screen. Participants began with 200 points and lost one point for each incorrect decision. To promote incentive compatibility (Hurwicz, 1972; Ledyard, 1989), participants earned a cent for every point earned. Reaction time was constrained such that participants were required to respond within 0.1 and 0.75 s from stimulus presentation. If participants responded in \(\leq 0.1\) s, \(\geq 0.75\) s, or failed to respond altogether, the point total turned red and decreased by 5 points. Each trial lasted 1.5 s and reward feedback for a given trial was displayed from the time of the participant’s response to the end of the trial.

To manipulate change point probability, the sex identity of the most rewarding Greeble was switched probabilistically, with a change occurring every 10, 20, or 30 trials, on average. To manipulate the belief in the value of the optimal target, the probability of reward for the optimal target was manipulated, with \(P\) set to 0.65, 0.75, or 0.85. Each session combined one value of \(P\) with one level of change point probability, such that all combinations of change point frequency and reward probability were imposed across the nine sessions (Fig. 2C). As in Experiment 1, the position of the high-value target was pseudo-randomized on each trial to prevent prepotent response selections on the basis of location.

Throughout the task, the head-stabilized diameter and gaze position of the left pupil were measured with an Eyelink 1000 desktop mount at 1000 Hz. Participants viewed stimuli from within a custom-built booth designed to eliminate the influence of ambient sources of luminance. Because the extent of the pupillary response is known to be highly sensitive to a variety of influences (Sirois and Brisson, 2014), we established the dynamic range of the pupillary response for each session by exposing participants to a sinusoidal variation in luminance prior to the reward-learning task. During the reward-learning task, all stimuli were rendered isoluminant with the background of the display to further prevent luminance-related confounds of the task-evoked pupillary response. To obtain as clean a trial-evoked pupillary response as possible and minimize the overlap of the pupillary response between trials, the inter-trial interval was sampled from a truncated exponential distribution with a minimum of 4 s, a maximum of 16 s, and a rate parameter of 2. The eyetracker was calibrated and the calibration was validated at the beginning of each session. See Pupil data preprocessing for pupil data preprocessing steps.

**Models and simulations**

**Q-Learning simulations**

A simple, tabular q-learning agent (Sutton and Barto, 1998) was used to simulate action selection in contexts of varying degrees of conflict and volatility. On each trial, \(i\), the agent chooses which of two actions to take according to the policy

\[
p_i = \frac{\exp^{\beta Q_i}}{\sum \exp^{\beta Q_i}}.
\]

Here \(\beta\) is the inverse temperature parameter, \(1/\tau\), reflecting the greediness of the selection policy and \(Q_i\) is the estimated state-action value vector on that trial. Higher values of \(\beta\) reflect more exploitative decision policies.

After selection, a binary reward was returned. This was used to update the \(Q\) table according
We called the belief in the reward attributable to a given action $B$ with $\theta \in [0; 1]$. This reward belief is learned a priori hypothesis was that the drift rate ($a$) and the boundary height ($b$) should change on a trial-by-trial basis according to two estimates of uncertainty from an ideal observer (Bond et al., 2018). We adapted the below ideal observer calculations from a previous study (Vaghi et al., 2017) (for the original formulation of this reduced ideal observer model and its derivation, see Nassar et al., 2010).

First we assumed that reward feedback drove the belief in the reward associated with an action. We called the belief in the reward attributable to a given action $B$. This reward belief is learned separately for each action target. Given the chosen target ($c$) and the unchosen target ($u$), the
belief in the mean reward for the chosen and unchosen targets on the next trial (trial $t+1$) was calculated as:

$$B_{t+1,c} = B_{t,c} + a_t \delta_t,$$

$$B_{t+1,u} = B_{t,u} (1 - \Omega_t) + \Omega_t E(r),$$

where $a_t$ denotes the learning rate, $\delta_t$ the prediction error, and $\Omega_t$ the change point probability on the current trial $t$, as discussed below. $E(r)$ refers to the pooled expected value of both targets:

$$E(r) = \frac{\bar{r}_{t0} + \bar{r}_{t1}}{2},$$

with $\bar{r}_{t0}, \bar{r}_{t1}$ fixed based on the imposed target reward probabilities.

The prediction error, $\delta_t$, was the difference between the reward obtained for the target chosen and the model belief:

$$\delta_t = r_t - B_t,c.$$

The signed belief in the reward difference between optimal and suboptimal targets ($\Delta B_t$) was calculated as the difference in reward value belief between target identities:

$$\Delta B_{t+1} = B_{t,opt} - B_{t,subopt}.$$

Model confidence ($\phi$) was defined as a function of change point probability ($\Omega$) and the variance of the generative distribution of points ($\sigma_n^2$), both of which formed an estimate of relative uncertainty ($RU_t$):

$$RU_t = \frac{\Omega_t \sigma_n^2 + (1 - \Omega_t)(1 - \phi_t)\sigma_n^2 + \Omega_t (1 - \Omega_t)(\Delta B_t)^2}{\Omega_t \sigma_n^2 + (1 - \Omega_t)(1 - \phi_t)\sigma_n^2 + \Omega_t (1 - \Omega_t)(\Delta B_t)^2 + \sigma_n^2}.$$

Thus $\phi$ is calculated as:

$$\phi_{t+1} = 1 - RU_t.$$

An estimate of the variance of the reward distribution, $\sigma_t^2$, was calculated as:

$$\sigma_t^2 = \sigma_n^2 + \frac{(1 - \phi_t)\sigma_n^2}{\phi_t}$$

where $\sigma_n$ is the fixed variance of the generative reward distribution.

The learning rate of the model ($\alpha$) was determined by the change point probability ($\Omega$) and the model confidence ($\phi$). Here, the learning rate was high if either 1) a change in the mean of the distribution of the difference in expected values was likely ($\Omega$ is high) or 2) the estimate of the mean was highly imprecise ($\sigma_t^2$ was high):

$$\alpha_t = \Omega_t + (1 - \Omega_t)(1 - \phi_t).$$

To model how learners update action-values, we calculated an estimate of how often the same action gave a different reward (Vaghi et al., 2017). This estimate gave our representation of change point probability, $\Omega$. The change point probability approached 1 from below as the probability of a sample coming from a uniform distribution, relative to a Gaussian distribution, increased:

$$\Omega_t = \frac{U(r_t)H}{U(r_t)H + N(r_t|B_{t,subopt}, \sigma_t^2)(1 - H)}.$$

In equation (12), $H$ refers to the hazard rate, or the global probability of a change point over trials:

$$H = \frac{n_{c,p}}{n_{trials}}.$$

We expected that the belief in the value of a given action and an estimate of environmental stability would target different parameters of the DDM model. Specifically, we hypothesized that
the belief in the relative reward for the two choices, $\Delta B$, would update the drift rate, $v$, or the rate of evidence accumulation:

$$v_{t+1} = \hat{\beta}_v \cdot \Delta B_t + v_t$$  \hspace{1cm} (14)

while the change point probability, $\Omega$, would decrease the boundary height, $a$, or the amount of evidence needed to make a decision:

$$a_{t+1} = a_t - \hat{\beta}_a \cdot \Omega_t.$$  \hspace{1cm} (15)

Hierarchical drift diffusion modeling

To assess whether and how much the ideal observer estimates of change point probability ($\Omega$) and the belief in the value of the optimal target ($\Delta B$) updated the rate of evidence accumulation ($v$) and the amount of evidence needed to make a decision ($a$), we regressed the ideal observer estimates onto the decision parameters using hierarchical drift diffusion model (HDDM) regression (Wiecki et al., 2013). These ideal observer estimates of environmental uncertainty served as a more direct and continuous measure of the uncertainty we sought to induce with our experimental conditions (see Fig. 4 for how the experimental conditions impacted these estimates). Considering this more direct approach, we pooled change point probability and belief across all conditions and used these values as our predictors of drift rate and boundary height. Responses were accuracy-coded, and the belief in the difference between targets values was transformed to the belief in the value of the optimal target ($\Delta B_{\text{optimal}(t)} = B_{\text{optimal}(t)} - B_{\text{suboptimal}(t)}$). This approach allowed us to estimate trial-by-trial covariation between the ideal observer estimates and the decision parameters.

Markov-chain Monte-Carlo methods were used to sample the posterior distributions of the regression coefficients. Ten thousand samples were drawn from the posterior distributions of the coefficients for each model. We chose this number of samples to optimize the trade-off between computation time and the precision of parameter estimates, and all model parameters converged to stability. This method generates a distributional estimate of the regression coefficients instead of a single best fit.

To identify the regression model that best accounted for the data, we conducted a model selection process using Deviance Information Criterion (DIC) scores. We compared the set of models (Table 1) to an intercept-only regression model ($DIC_i - DIC_{\text{intercept}}$). A lower DIC score indicates a model that loses less information. Here, a difference of $\leq 2$ points from the lowest-scoring model cannot rule out the higher scoring model; a difference of 3 to 7 points suggests that the higher scoring model has considerably less support; and a difference of 10 points suggests essentially no support for the higher scoring model (Spiegelhalter et al., 2002; Burnham and Anderson, 1998).

To test our hypotheses regarding these HDDM regression estimates, we used the posterior distributions of the regression parameters. To quantify the reliability of each regression coefficient, we computed the probability of the regression coefficient being greater than or less than 0 over the posterior distribution. We considered a regression coefficient to be reliable if the estimated coefficient maintained the same sign over at least 95% of the mass of the posterior distribution. Trial-by-trial estimates of boundary height and drift rate were calculated from the regression equation for the winning model.

For Experiment 2, a regression model as described above was fit to each subject.

Analyses

General statistical analysis

Statistical analyses and data visualization were conducted using custom scripts written in R (R Foundation for Statistical Computing, version 3.4.3) and Python (Python Software Foundation, version 3.5.5).

To determine how many trials would be needed to detect proposed condition effects, we conducted a power analysis by way of parameter recovery. For this we simulated accuracy and reaction time data using our hypothesized model (Cognitive model) and calculated the generative or “true”
mean drift rate and boundary height parameters across trials. Then we conducted hierarchical parameter estimation given 200, 400, 600, 800, or 1000 simulated trials. The mean squared error of parameter estimates was stable at 600 trials for all decision parameters. Additionally, as a validation measure, we estimated parameters using component models (drift rate alone, boundary height alone) and a combined model (drift rate and boundary height). We found that the Deviance Information Criterion (DIC) scores among competing models were clearly separable at 600 trials, and in favor of the hypothesized model from which we generated the data, as expected (Acknowledgments). Based on these results, we used 600 trials per condition for each participant for our first experiment. We chose to recruit 24 participants for this experiment to fully counterbalance the four conditions (4! = 24).

Binary accuracy data were submitted to a mixed effects logistic regression analysis with either the degree of conflict (the probability of reward for the optimal target) or the degree of volatility (mean change point frequency) as predictors. The resulting log-likelihood estimates were transformed to likelihood for interpretability. RT data were log-transformed and submitted to a mixed effects linear regression analysis with the same predictors as in the previous analysis. To determine if participants used ideal observer estimates to update their behavior, two more mixed effects regression analyses were performed. Estimates of change point probability and the belief in the value of the optimal target served as predictors of reaction time and accuracy across groups. As before, we used a mixed logistic regression for accuracy data and a mixed linear regression for reaction time data.

Because we adopted a within-subjects design, all regression analyses of behavior modeled the non-independence of the data as constantly correlated data within participants (random intercepts). Unless otherwise specified, we report bootstrapped 95% confidence intervals for behavioral regression estimates. To prevent any bias in the regression estimates emerging from collinearity between predictors and to aid easy interpretation, all predictors for these regressions were mean-centered and standardized prior to analysis. The Satterthwaite approximation was used to estimate p-values for mixed effects models (Satterthwaite, 1946; Luke, 2017).

Decision vector representation
To concisely capture the change-point-driven response in the relationship between the boundary height and the drift rate over time, we represented the relationship between these two decision variables in vector space. Trial-by-trial estimates of drift rate and boundary height were calculated from the winning HDDM regression equation and z-scored. Then the difference between each sequential set of (a, v) coordinates was calculated to produce a vector length. The arctangent between these subtracted values was computed to yield an angle in radians between sequential decision vectors (Fig. 5B).

For Experiment 1, these computations were performed from the trial prior to the onset of the change point to eight trials after the change point. The initial window of nine trials was selected to maximize the overlap of stable data between high and low volatility conditions (see Supp. Fig. 1). This resulted in a sequence of angles formed between trials -1 and 0 ($\Delta t_1$, yielding $\theta_1$), 0 and 1 ($\Delta t_2$, yielding $\theta_2$), and so on. To observe the timescale of these dynamics, a circular regression (Mulder and Klugkist, 2017) was performed to determine how $\theta$ changed as a function of the number of trials after the change point:

$$\theta = \hat{\beta}_0 + \hat{\beta}_{\Delta t_1} + \ldots \hat{\beta}_{\Delta t_n},$$

To quantitatively assess the number of trials needed for $\theta$ to stabilize, we calculated the probability that the posterior distributions of the regression estimates (Supp. Fig. 6) for sequential pairs of trials had equal means ($\theta_{\Delta t_1} = \theta_{\Delta t_2}$). This result (Supp. Fig. 7) provided an out-of-set constraint on the timescale of the decision response to consider for analogous analyses in Experiment 2.

Experiment 2 used the stability convergence analysis from Experiment 1 to guide the timescale of further circular analyses and, thus, placed a constraint on the complexity of the models pro-
posed (Model proposals and evaluation). Because Experiment 2 took a replication-based approach, a separate model was fit for each participant for all proposed models. We report the mean and 95% CI of the posterior distributions of regression parameter estimates and the mean and standard deviation of estimates across subjects.

The circular regression analyses used Markov-chain Monte-Carlo (MCMC) methods to sample the posterior distributions of the regression coefficients. For both experiments, 10,000 effective samples were drawn from the posterior distributions of the coefficients for each model (Kruschke and Vanpaemel, 2015). Traces were plotted against MCMC iteration for a visual assessment of equilibrium, the autocorrelation function was calculated to verify independence of MCMC steps, trace distributions were visually evaluated for normality, and point estimates of the mean value were verified to be contained within the 95% credible interval of the posterior distribution for the estimated coefficients.

Pupil data preprocessing

Pupil diameter data were segmented to capture the interval from 500 ms prior to trial onset to the end of the 1500 ms trial, for a total of 2000 ms of data per trial. While the latency in the phasic component of the task-evoked pupillary response ranges from 100-200 ms on average (Beatty, 1982), suggesting that our segmentation should end 200 ms after the trial ending, participants tended to blink after the offset of the stimulus and during the intertrial interval (see Supp. Fig. 8 for a representative sample of blink timing). Because of this, we ended the analysis window with the offset of the stimulus. Following segmentation, pupil diameter samples marked as blinks by the Eyelink 1000 default blink detection algorithm and zero- or negative-valued samples were replaced by linearly interpolating between adjacent valid samples. Pupil diameter samples with values exceeding three standard deviations of the mean value for that session were likewise removed and interpolated. Interpolated data were bandpass filtered using a .01 to 5 Hz second-order Butterworth filter. Median pupil diameter calculated over the 500 ms prior to the onset of the stimulus was subtracted from the trial data. Finally, processed data were z-scored by session.

For each trial interval, we characterized the evoked response as the mean of the pupil data over that interval, the latency to peak onset and offset, the latency to peak amplitude, the peak amplitude, and the area under the curve of the phasic pupillary response (Fig. 7A). We then submitted these metrics to principal component analysis to reduce their dimensionality while capturing maximum variance. Evoked response characterization and principal component analysis were conducted for each session and for each subject.

The 95% CI for the number of principal components needed to explain 95% of the variance in the data was calculated over subjects and sessions to determine the number of principal components to keep for further analysis.

To aid in interpreting further analysis using the selected principal components, the feature importance of each pupil metric was calculated for each principal component and aggregated across subjects as a mean and bootstrapped 95% CI (Fig. 7B).

Model proposals and evaluation

To assess the hypothesized influences on $\theta$ in Experiment 2, we began our model set proposal with two null hypotheses. The absolute null model estimates decision dynamics as a function of the intercept, or the average of $\theta$:

$$\theta = \beta_0.$$ 

The second, time-null model estimates decision dynamics solely as a function of time, with the timescale of consideration determined by the results of the stability convergence analysis from Experiment 1:

$$\theta = \beta_0 + \beta_\Delta \Delta.$$
We first evaluated whether the posterior probability of the time-null model given the data was greater than the posterior probability for the absolute null model. If the lower bound of the 95% CI of the posterior probability for the time-null model exceeded the upper bound of the 95% CI for the absolute null model (i.e., the posterior probability was greater for the time-null model and the CIs were non-overlapping), we proceeded to evaluate the evidence for alternative models relative to this time-null model. We evaluated the statistical reliability of the posterior probabilities using a bootstrapped 95% CI computed over subjects.

We considered an explicit set of hypotheses regarding the effect of the pupillary response on boundary height and drift rate dynamics (see Fig. 8 for the full set of models considered). The first two principal component projections of the set of pupil metrics, which we term the timing and magnitude components, respectively, were included in this model set to evaluate the effect of the timing and magnitude of noradrenergic dynamics on the change-point-evoked decision manifold. Under the assumption of a neuromodulatory effect on decision dynamics, these principal components were shifted forward by one trial to match the expected timing of the response to neuromodulation.

To determine whether perturbations of volatility and conflict affected decision dynamics, we estimated decision dynamics as a function of $\lambda$ and $p$, where $\lambda$ corresponds to the average length of an epoch and $p$ corresponds to the mean probability of reward for optimal target selection (see Table 1 for the full set of models considered).

We used Bayes Factors to quantify the ratio of evidence for competing hypotheses (Wagenmakers, 2007). To estimate whether these models accounted for decision dynamics beyond the effect of time alone, we calculate the Bayes Factor for the time-null model relative to each candidate model ($BF_{ij}$). Finally, we calculate the posterior probability of the null model given the full set of alternative models (Wagenmakers, 2007). Note that this approach assumes that each model has equal a priori plausibility.

Bayes Factor visualizations represent the mean and bootstrapped 95% CI with 1000 bootstrap iterations.

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Appendix 1 Figure 1. Initial window selection.

Analysis conducted on data from Experiment 1 to determine the timescale of the response that maximized the intersection between high volatility ($\lambda = 15$) and low volatility ($\lambda = 35$) data. The bolded line represents the mean and the gray lines represent individual subjects. The dotted line indicates the initial window of nine trials used.
Appendix 1 Figure 2. Ideal observer estimates for Experiment 2.

A) The average belief in the value of the optimal target ($\Delta B$) as a function of the probability of reward (conflict) and the average period of stability for the optimal choice ($\lambda$; volatility).

B) Average change point probability ($\Omega$) as a function of conflict and volatility.
Appendix 1 Figure 3. Distributions of HDDM $\beta$ coefficients for Experiment 1.
A) The distribution of the estimate of the relationship between the drift rate ($v$) and the belief in the value of the optimal choice ($\Delta B$). B) The distribution of the estimate of the relationship between the boundary height ($a$) and change point probability ($\Omega$).

Appendix 1 Figure 4. Distributions of HDDM $\beta$ coefficients for Experiment 2.
A) The relationship between the drift rate ($v$) and the belief in the value of the optimal choice ($\Delta B$). B) The relationship between the boundary height ($a$) and change point probability ($\Omega$).
Appendix 1 Figure 5. Decision trajectories for Experiment 1.
The change in boundary height and drift rate over time for Experiment 1. Each line represents the data for one participant.
Appendix 1 Figure 6. Vector distributions.

The distribution of vectors formed between sequential ($a, v$) coordinates, beginning with the trial prior to the change point. The black arrow shown on each panel represents the mean vector.
Appendix 1 Figure 7. Stability analysis.

Analysis conducted on Experiment 1 to determine the timescale of the response to consider for Experiment 2. The estimated angle is plotted as a function of time within an epoch (estimate from circular regression).
Appendix 1 Figure 8. Quantification of stability.

Probability that sequential posterior distributions for $\theta_{\Delta}$ have equal means.
Blink timing for a sample participant. For visibility, thirty trials were selected at random. The onset of the trial is marked as time 0 and the trial ends at 1500 ms. Blinks are marked in black. Blink timing plots are available for all subjects and all conditions in the GitHub repository for this publication.
Appendix 2 Table 1. Power analysis for Experiment 1.

The results of the model comparison analysis using simulated data. Roman numerals refer to a given model, as defined by the mapping between the ideal observer estimates and decision parameters in the first two columns. The column labeled DIC gives the raw DIC score, ΔDICnull lists the change in model fit from an intercept-only model (the null-adjusted fit), and ΔDICbest provides the change in null-adjusted model fit from the best-fitting model. The last row represents the null, intercept-only regression model. The best performing model is denoted by an asterisk.