Mechanisms of adjustments to different types of uncertainty in the reward environment across mice and monkeys

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Abstract

Despite being unpredictable and uncertain, reward environments often exhibit certain regularities, and animals navigating these environments try to detect and utilize such regularities to adapt their behavior. However, successful learning requires that animals also adjust to uncertainty associated with those regularities. Here, we analyzed choice data from two comparable dynamic foraging tasks in mice and monkeys to investigate mechanisms underlying adjustments to different types of uncertainty. In these tasks, animals selected between two choice options that delivered reward probabilistically, while baseline reward probabilities changed after a variable number (block) of trials without any cues to the animals. To measure adjustments in behavior, we applied a set of metrics based on information theory that quantify consistency in behavior, and fit choice data using reinforcement learning models. We found that in both species, learning and choice were affected by uncertainty about reward outcomes (in terms of determining the better option) and by expectation about when the environment may change. However, these effects were mediated through different mechanisms. First, more uncertainty about the better option resulted in slower learning and forgetting in mice, whereas it had no significant effect in monkeys. Second, expectation of block switches accompanied slower learning, faster forgetting, and increased stochasticity in choice in mice, whereas it only reduced learning rates in monkeys. Overall, while demonstrating the usefulness of entropy-based metrics in studying adaptive behavior, our study provides evidence for multiple types of adjustments in learning and choice behavior according to uncertainty in the reward environment.

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Introduction

Learning and choice in dynamic environments are shaped by a tradeoff between adaptability and precision (Farashahi et al., 2017a; Soltani, Murray, et al., 2021). In stable environments, learning can be slow to allow more precise estimates of reward contingencies. In contrast, in unstable and thus less predictable environments, learning can become faster, slower, or show no change depending on where the decision maker falls in terms of the adaptability and precision tradeoff. This is because faster learning enables faster updates but also comes at the cost of less precision, which may not be desirable in certain situations. Consistent with this, Behrens and colleagues (Behrens et al., 2007) have shown that more uncertainty in terms of volatility can result in higher learning rates, whereas others have found no evidence for such adjustments (Donahue & Lee, 2015, Farashahi et al., 2019). Instead, it was shown that volatility can decrease the weight of more uncertain information (estimated reward probability) relative to that of more certain information (reward magnitude) and improve encoding of task-relevant signals (Donahue & Lee, 2015; Massi et al., 2018; Farashahi et al, 2019).

Estimation and adjustments to uncertainty are tightly linked to detecting and learning regularities in the reward environment. Such regularities can modulate learning and decision making to enable the decision maker to anticipate and prepare for changes (Atilgan et al., 2022). Nevertheless, any regularity is accompanied by an inevitable uncertainty due to the stochastic nature of the reward environment. It is thus not surprising that the influences of different types of uncertainty on learning and the neural mechanisms by which these effects are mediated have been topics of much interest (Yu & Dayan, 2005; Behrens et al., 2007; Winstanley & Floresco, 2016; Faraji et al., 2018; Soltani & Izquierdo, 2019; Piray & Daw, 2021). For example, it has been shown that during a dynamic foraging task in mice, serotonin neurons in mice represent expected uncertainty and track unexpected uncertainty for updates (Grossman et al., 2022). In monkeys, neurons in the dorsolateral prefrontal cortex track the statistical variance of reward outcomes ('risk') resulting from choices for specific options (Grabenhorst et al., 2019). Nonetheless, similarities and differences between adjustments to different types of uncertainty are not fully understood.

Understanding adjustments to uncertainty requires studying behavior in dynamic learning tasks that involve different types of uncertainty, for example the kind resulting from changing reward probabilities for different choice alternatives between blocks of trials. To reveal mechanisms underlying such adjustments, often choice data are fit using different reinforcement learning (RL) models that have been augmented with additional components to achieve better goodness-of-fit (Farashahi et al., 2017b, Wittmann et al., 2020; Womelsdorf et al., 2021; Grossman et al., 2022; Collins & Shenhav, 2022). Although such RL models can reveal possible mechanisms underlying

flexible behavior, fit of choice data requires many trials, which makes the application of this method to study continuous adjustments in behavior very challenging.

To address this issue, we recently have developed a set of new behavioral metrics based on information theory that can quantify consistency in trial-by-trial response to reward feedback using a relatively small number of trials (Trepka et al., 2021). These metrics provide a convenient, model-free tool to measure randomness in adaptive choice behavior and guide further model development. More importantly, these metrics can be computed from any segment of the task, enabling semi-continuous quantification of behavioral adjustments over time.

Here, we used these metrics to study adjustments in learning and choice behavior to different types of uncertainty in the reward environment. To that end, we re-analyzed data from mice and monkeys that performed similar dynamic foraging in which selection of two choice alternatives was rewarded probabilistically, whereas baseline reward probabilities switched after a certain number of trials (referred to as the block length). We investigated the influences of uncertainty in determining the better option as well as uncertainty about the time of block switches on learning and choice using entropy-based metrics. These include overall entropy of choice strategy in terms of stay or switch, entropy of reward-dependent strategy, entropy of option-dependent strategy, and mutual information between reward outcome and choice strategy. While incorporating probability of win-stay and lose-switch strategies, these metrics can better summarize local response to reward feedback, which then can be used to predict global choice behavior or its adjustments (Trepka et al. 2021).

With regard to uncertainty in determining the better option based on reward feedback, we hypothesized that increased uncertainty should result in slowing down of learning to allow more accurate estimation of reward probabilities and/or in more stochastic behavior to enable more exploration. This could happen via different non-exclusive mechanisms such as a decrease in the learning rates and an overall increase in the stochasticity of choice. In addition, we hypothesized that when blocks are longer than expected, choice or adopted choice strategy would be more random as animals expect changes in reward probabilities more strongly and in the absence of any changes in the environment. Finally, expectation of block switches could interact and/or depend on the level of uncertainty in determining the better option.

Briefly, we found that stochasticity in choice behavior and response to reward feedback measured by entropy-based metrics were systematically affected by uncertainty about the better option and expectation of block switches in both mice and monkeys. Further, by fitting RL models to different groups of choice data identified by the application of entropy-based metrics, we identified possible mechanisms by which different types of uncertainty influence behavior across the two species.

Methods

Experimental paradigms

To examine whether and how animals use regularities and accompanied uncertainty of the reward environment to guide their behavior, we reanalyzed data from mice and monkeys performing two dynamic foraging tasks with variable block lengths. There were key similarities in these experiments that allowed us to compare performance directly across species. First, while baseline reward probabilities remained fixed within a block of trials, probability of reward assignment on the unchosen option increased over time in a similar fashion in both tasks. Second, the block length changed from block to block but the range of block lengths were similar across the two species: with 40-100 trials in mice and 50-150 trials in monkeys.

Mouse experiment. In the mouse experiment (Bari et al., 2019), water-restricted mice were habituated for 1-2 day and head-restrained for the task. 18 male C57BL/6J (The Jackson Laboratory, 000664) mice aged 6-20 weeks were used in the experiment. A "go-cue" was signaled by an odor, after which they made a leftward or rightward lick toward one of the two water ports for a probabilistic reward (**Figure 1A**). Reward was a drop of water (2-4 μ L). In 5% of trials, a "no-go cue" was signaled by another odor, after which licks were neither rewarded nor punished. Each trial was followed by an inter-trial interval (ITI), of which the duration was drawn from an exponential distribution with a rate parameter of 0.3 with a maximum of 30s.

Reward probabilities were assigned to the left and right water ports, with one port yielding a higher reward probability ('better' option) than the other ('worse' option). Rewards were baited, such that if a reward was assigned to a particular side and was unchosen, the reward would remain there until that side was selected. Reward probabilities were chosen from 16 different sets of reward schedules (**Figure S1C**). The majority of sessions (88.8%) used two sets of reward probabilities equal to 0.4 and 0.1 (40/10), and 0.4 and 0.05 (40/5). The two reward probabilities were held constant for a fixed number of trials (a block) and reversed (block switches or reversal) without any cues to the animals. The lengths of each block were drawn from a uniform distribution spanning 40-100 trials. On rare occasions when mice demonstrated strong side biases, block lengths were shortened or lengthened.

For a given day of experiment, mice performed one of two versions of the task, a two-probability task (523 sessions) and a multiple-probability task (only 5 sessions). In the two-probability task, only two reward probabilities were assigned from the pool of possible probabilities and reversed for a given session. In the multiple-probability task, the better and worse options were still reversed after block switches, but more than two

reward schedules were used for a given session. These probabilities were chosen such that the ratios would equal {8:1, 6:1, 3:1, 1:1}, to match parameters from the previous experiment by Sugrue et al., 2004. Majority of the mice (16 out of 18) performed the two-probability task only. For the other two mice, multiple-probability sessions were randomly interleaved with two-probability sessions. In total, 18 mice performed 3768 blocks during 528 sessions (mean 7.13 block switches per session). Here, we focus our analyses on the two-probability task only because most of the mouse experiment was done using this task and there was a significant difference between the performance in two-probability and multiple-probability tasks. All behavioral tasks were performed in a dark sound-attenuated chamber, with white noise delivered between 2-60 kHz. All surgical and experimental procedures were in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and approved by the Johns Hopkins University Animal Care and Use Committee. This experimental setup and some analyses of the data for electrophysiological recordings and matching behavior have been reported previously (Bari et al., 2019; Trepka et al., 2021; Bari & Gershman, 2022). All analyses reported in this study are novel.

Monkey experiment. In the monkey experiment (Tsutsui et al., 2016), two adult male macaque monkeys (*Macaca mulatta*) weighing 5.5 - 6.5 kg were trained in a reward-based foraging task. To initiate a trial, the monkeys fixated on the red fixation spot in the center of a computer screen and contacted a touch-sensitive, immobile resting key at the elbow height. After fixation, two stimuli (two different visual fractal objects, A and B) appeared simultaneously on the screen to the left and right of the fixation point (5° visual angle). Two stimuli were assigned pseudorandomly to left and right positions and were not related to reward. To make a choice, monkeys made a saccadic eye movement to the target of its choice within a time window of 0.25–0.75s (**Figure 1B**). Reward was a drop of juice (0.7 ml) delivered via a spout in front of the animal's mouth.

The reward probabilities of objects A and B were independently calculated in every trial, such that the instantaneous reward probability increased for the unchosen option as a function of the number of consecutive unchosen trials (Eq. 1):

$$P = 1 - (1 - P_0)^{n+1}$$
 (Eq. 1)

where P is the instantaneous reward probability of an object, P_0 is an experimentally imposed, baseline reward probability, and n is the number of trials that the object had been consecutively unchosen. This equation implies that an unchosen option became more rewarding, the longer time it remained unselected. The reward probability fell back to the baseline probability with every choice of that object, regardless of whether that choice was rewarded or not. Functionally, this algorithm implements the same baiting mechanism in the mouse experiment described above. The baseline reward

probabilities (P_0) were chosen from 20 different sets of reward schedules (**Figure S1D**). The two baseline probabilities were held constant for each option for a fixed number of trials ('block') and changed across blocks ('block switches') such that the previously better option (with higher P_0) became the worse option (lower P_0). The lengths and reward schedules of each block changed randomly from one block to the next, typically ranging 50–150 trials without any cues to the animals. In total, the two monkeys performed 251 blocks during 103 session days (mean 2.43 block switches per session). All animal procedures conformed to the US National Institutes of Health Guidelines and were approved by the Home Office of the United Kingdom. This experimental setup and some analyses of the electrophysiological recording data have also been reported previously (Tsutsui et al., 2016). All analyses reported in this study are novel.

Entropy-based metrics

Here, we utilized previously introduced entropy-based metrics to quantify learning and choice behavior (Trepka et al., 2021). This includes entropy of choice strategy (H(str)), conditional entropy of reward-dependent strategy (ERDS), conditional entropy of optiondependent strategy (EODS), mutual information between reward outcome and choice strategy (MIRS), and mutual information between choice and choice strategy (MIOS) (Figure 1). Briefly, H(str) measures consistency or randomness in the adopted choice strategy in terms of stay or switch, whereas ERDS and EODS measure the same consistency but considering reward feedback and/or choice in the previous trial. MIRS, on the other hand, measures the amount of information shared between the adopted strategy (stay or switch) and reward feedback (win or lose). The maximum value for these quantities is equal to 1 corresponding to completely random behavior (equally adopting stay or switch strategy) or no relationship between reward outcome and choice strategy. Conceptually, metrics based on conditional entropy and mutual information aim to capture the degree to which the animal's choice behavior can be explained or predicted by the information prior to that decision, such as by presence of reward (ERDS, MIRS) or previous selection (EODS, MIOS).

More specifically, ERDS is calculated as follows:

$$\begin{split} ERDS &= H(str|rew) = H(str) - I(str;rew) \\ &= -(P(stay,win) \times log_2(\frac{P(stay,win)}{P(win)}) + P(switch,win) \times log_2(\frac{P(switch,win)}{P(win)}) \\ &+ P(stay,lose) \times log_2(\frac{P(stay,lose)}{P(lose)}) + P(switch,lose) \times log_2(\frac{P(switch,lose)}{P(lose)})), \end{split}$$
 (Eq. 2)

where *str* is strategy coded as stay (1) or switch (0), *rew* is the previous reward outcome coded as reward (1) or no reward (0). As seen above, the more traditional measures win-stay and lose-switch appear in this equation as $\frac{P(stay,win)}{P(win)}$ and $\frac{P(stay,lose)}{P(lose)}$, respectively. Additionally, H(str) above is the entropy of strategy (stay or switch), and I(str; rew) is the mutual information of strategy and reward (MIRS):

$$\begin{split} H(str) &= -(P(stay) \times log_2 P(stay) + P(switch) \times log_2 P(switch). \quad \text{(Eq. 3)} \\ MIRS &= I(str; rew) = -(P(stay, win) \times log_2(\frac{P(stay, win)}{P(stay) \times P(win)}) + P(switch, win) \times log_2(\frac{P(switch, win)}{P(switch, win)}) + P(stay, lose) \times log_2(\frac{P(stay, win)}{P(switch, win)}) + P(switch, lose) \times log_2(\frac{P(switch, win)}{P(switch, win)}) + P(switch, lose) \end{split}$$

$$log_2(\frac{P(switch, win)}{P(switch, win)}). \quad \text{(Eq. 4)} \end{split}$$

As these terms suggest, ERDS measures the dependence of adopting a response strategy on reward feedback (*win* or *lose*). Lower ERDS values correspond to decreased randomness in the variable and thus more consistency in the utilized strategy.

Similarly, we also used the entropy of option-dependent strategies (EODS), equal to a conditional entropy of stay/switch strategy (*str*) given the choice of better or worse option from the previous trial (*opt*):

$$\begin{split} EODS &= H(str|opt) = H(str) - I(str;opt) \\ &= -(P(stay,better) \times log_2(\frac{P(stay,better)}{P(better)}) + P(switch,better) \times log_2(\frac{P(switch,better)}{P(better)}) \\ &+ P(stay,worse) \times log_2(\frac{P(stay,worse)}{P(worse)}) + P(switch,worse) \times log_2(\frac{P(switch,worse)}{P(worse)})), \text{ (Eq. 5)} \end{split}$$

where *opt* is the previous choice (1 = better option and 0 = worse option), and I(str; opt) is the mutual information of strategy and option (MIOS):

$$\begin{split} \mathit{MIOS} &= \mathit{I(str;opt)} = -(\mathit{P(stay,better)} \times log_2(\frac{\mathit{P(stay,better)}}{\mathit{P(stay)} \times \mathit{P(better)}}) + \mathit{P(switch,better)} \times \\ log_2(\frac{\mathit{P(switch,better)}}{\mathit{P(switch)} \times \mathit{P(better)}}) + \mathit{P(stay,worse)} \times log_2(\frac{\mathit{P(stay,worse)}}{\mathit{P(stay)} \times \mathit{P(worse)}}) + \mathit{P(switch,worse)} \times \\ log_2(\frac{\mathit{P(switch,worse)}}{\mathit{P(switch,worse)}})). \end{split}$$
 (Eq. 6)

Note that from the above equations, ERDS = H(str) - MIRS and EODS = H(str) - MIOS. This implies that, for example, increased ERDS in one setting could be due to increased H(str) or decreased MIRS, or both.

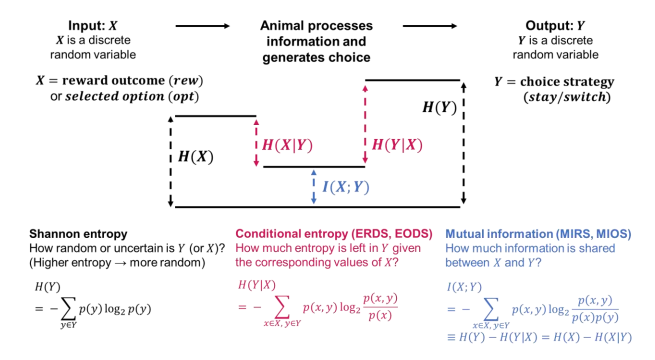


Figure 1. Application of information theory to studying choice behavior under uncertainty. The stochastic nature of the reward environment results in uncertainty in reward outcome that can be measured using Shannon entropy (H(X)). The animal used information in the environment to generate choice, that can be quantified as repeating the previous choice (stay) or switching to the alternative choice option (for two-alternative choice tasks). Randomness in choice can also be measured using Shannon entropy (H(Y)). The link between previous reward outcome and animals' strategy can be quantified using conditional entropy (e.g., H(Y|X)) measuring uncertainty in strategy given reward outcome on the previous trial, or mutual information (I(X; Y)) measuring shared information.

Quantifying expected uncertainty and expectation of block switches in the reward environment

In both mouse and monkey experiments, baseline reward probabilities associated with two choice options were fixed within a block of trials, but block lengths changed randomly from one block to the next. This means that within a given block, animals could develop an estimation for uncertainty in reward feedback, often referred to as expected uncertainty corresponding to uncertainty attributable to the stochastic or probabilistic nature in reward outcome (Soltani & Izquierdo, 2019). Additionally, the animals could form an expectation about block lengths or how long reward probabilities remain unchanged. Such expectation could have significant effects on behavior because the animals could erroneously perceive that a block switch happens after a

certain number of trials have passed. Both types of expectation and their effects on animals' behavior might be read from the animal's responses, such as from the changes in the adopted strategies captured by ERDS or EODS. Therefore, we defined different measures to quantify the different types of expectation as explained below.

Expected uncertainty. As the animals performed a wide range of different reward schedules (**Figure S1**), we defined expected uncertainty as the difference between baseline reward probabilities for better (P_B) and worse option (P_W) for each block: $\Delta P = P_B - P_W$. For the monkey experiment, base reward probabilities (P_0) for better and worse options were used to compute this metric. The larger the quantity ΔP , the easier it is for the animal to discern which is the better option, and hence, experienced less uncertainty in determining the better option. This uncertainty was deemed 'expected' in the sense that P_B and P_W were stable for the whole block and reflected stochasticity in the environment. We note quantifying expected uncertainty with ΔP is different from the expected uncertainty defined as the overall variance in reward outcome, which can be computed using the average absolute value of reward prediction error (Soltani & Izquierdo, 2019).

Using ΔP , we further categorized the blocks into *more uncertain* and *less uncertain* blocks based on the median split of ΔP within each subject. Blocks that had larger ΔP than the median were categorized as less uncertain, and smaller ΔP as more uncertain blocks. We hypothesized that increased uncertainty should result in more stochastic behavior, and therefore *more uncertain* blocks with ΔP smaller than median would be characterized by higher entropies or lower mutual information. Overall, the mouse experiment had a median ΔP of 0.35 (M = 0.330, SD = 0.042), and the monkey experiment had median ΔP of 0.25 (M = 0.281, SD = 0.122).

Expected block length. As the animals experienced a range of different block lengths, we hypothesized that the animals keep an expectation of how long the current block is, and that it is updated at every block switch. More specifically, considering L_i (length of block i), we calculated the expected block length $E[L_i]$ as follows:

$$E[L_{i+1}] = E[L_i] + \gamma_E(L_i - E[L_i]), \quad (i > 2)$$
 (Eq. 7)

where γ_E is an update rate (for computing block length expectation) controlling how much to weigh the recent evidence, analogous to the learning rate in the reinforcement learning models. The amount of update depends on the *block length prediction error* (*BLPE*) equal to the difference between actual and expected block length at block i:

$$BLPE_i = L_i - E[L_i]. mtext{(Eq. 8)}$$

Negative BLPE indicates that a given block was shorter than expected, and positive BLPE suggests that a block was longer than the expected length.

Equation 7 implies that, the larger the γ_E is, the animal weighs the recent evidence more so to update the estimated block length for the next block. For our analysis, we set γ_E = 0.1. For the very first block, $E[L_1]$ was initialized with a median block length of each dataset: $E[L_1]$ = 54 and $E[L_1]$ = 51 for mice and monkeys, respectively. Any sessions which did not have any block switches were omitted from this update and $E[L_i]$ was held constant for that block, given that it did not contribute to forming an expectation of block switches. Mouse data did not have such single-block sessions, whereas monkey data had 40 blocks that terminated without block switches (15.9% of total blocks).

Alternatively, the expected block length can be defined from a window of size *N*, as the average block length for the past *N* blocks:

$$E[L_i]_N = \frac{1}{N} \sum_{j=i-N}^{i-1} L_j \quad (i > 2).$$
 (Eq. 9)

Here, the larger window size N corresponds to smaller update rate γ_E , meaning that the animals estimate expected block length from a longer timescale.

Finally, we used BLPE (Eq. 8) to categorize blocks as shorter than expected (SE) or longer than expected (LE) blocks and compared behavior across these two types of blocks. We hypothesized that animals could develop an expectation of block switches through experience and use this to modulate their behavior. More specifically, we assumed that if the actual block length (L_i) is past the expected length ($E[L_i]$), the animal would be more likely to incorporate reward feedback as if they signal changes in the environment. In terms of entropy measures, this could be reflected in higher entropies of reward-dependent (ERDS) or option-dependent strategy (EODS), and/or lower mutual information between strategy and reward (MIRS) or previous option (MIOS).

Reinforcement learning (RL) models

RL models. We fit the choice behavior of mice and monkeys using four standard reinforcement learning models. The first model, which we refer to as RL1, has two parameters: a single learning rate α and inverse temperature β representing choice consistency. The value of the chosen option (Q_C) is updated as follows:

$$Q_C(t+1) = Q_C(t+1) + \alpha(R - Q_C(t))$$
 (Eq. 10)

where $C \in \{Left, Right\}$ for mice and $C \in \{Object A, Object B\}$ for monkeys, and R indicates presence (1) or absence (0) of reward on the given trial t.

The second model, referred to as RL1_{decay}, has an additional parameter decay rate γ_{decay} for the value of the unchosen option. In this model, the value of the unchosen option (Q_U) decayed passively, using the following equation:

$$Q_U(t+1) = Q_U(t+1) - \gamma_{decay}(Q_C(t)).$$
 (Eq. 11)

The value estimates for the chosen option were updated as in Eq. 10.

The third and fourth models, RL2 and RL2_{decay}, are identical to RL1 and RL1_{decay} except that there are now two separate learning rates for rewarded and unrewarded trials (α + and α -, respectively). For this set of models, the value of the chosen option (Q_c) was updated as follows:

$$Q_C(t+1) = Q_C(t+1) + \alpha_+(R-Q_C(t))$$
 if $R=1$, or
$$= Q_C(t+1) + \alpha_-(R-Q_C(t))$$
 if $R=0$. (Eq. 12)

RL2 did not include a decay term, whereas RL2_{decay} followed a decay equation for an unchosen option as in Eq.11.

For all models above, the probability of choosing an outcome was then computed using a softmax function:

$$P_1(t) = (1 + e^{\beta(Q_1(t) - Q_2(t))})^{-1},$$
 (Eq. 13)

where $1 = \{Left, Object A\}$ and $2 = \{Right, Object B\}$ for mice and monkeys. The minimum and maximum values of β were set to 1 and 100, respectively.

Model fitting process. We used the standard maximum likelihood estimation method to fit and estimate the best-fit parameters for the described models. One set of model parameters was fit to each block of mice and monkeys choice data. When fitting and simulating RL models with mouse data, we treated infrequent miss trials (average of 3.64 per session) and no-go trials (average of 16.83 per session) as if they had not occurred. For the very first block of the session only, value estimates for the two options (Q) were initialized at 0.5. For the consequent blocks after block switches (within the same session), initial value estimates were set using Q values from the last trial of the previous block, given by its best-fit parameters. Fitting was performed using the MATLAB optimization function *fmincon*, repeating the search for 10 random parameter values.

To quantify goodness-of-fit, we computed the Akaike Information Criterion (AIC) for each block of mouse and monkey dataset:

$$AIC = -2 * logLikelihood + 2 * p$$
, (Eq. 14)

where *p* is the number of free parameters in a given model. Since each block had different number of trials, we further normalized the AIC by the number of trials in each block, i.e., computing AIC per each trial:

$$AIC_{norm} = AIC/L,$$
 (Eq. 15)

where L is the number of trials or length of each block. We determined the winning model as the one with the lowest mean AIC_{norm} across all blocks.

We further computed the probability that a given model is the best model for the dataset given the set of candidate models ($Akaike\ weights$). Specifically, $Akaike\ weights$ (w_i) for the i-th model (M_i) in a set of n models were computed as follows:

$$\Delta \overline{AIC}_{norm}(M_i) = \overline{AIC}_{norm}(M_i) - min\{\overline{AIC}_{norm}(M_1), \overline{AIC}_{norm}(M_2), \dots, \overline{AIC}_{norm}(M_n)\},$$

$$w_i = exp\left(-\frac{1}{2}\Delta \overline{AIC}_{norm}(M_i)\right)/\sum_{j=1}^n exp\left(-\frac{1}{2}\Delta \overline{AIC}_{norm}(M_j)\right), \quad \text{(Eq. 16)}$$

where $\overline{AIC}_{norm}(M_i)$ indicates the mean AIC_{norm} of M_i across all blocks, $\Delta \overline{AIC}_{norm}(M_i)$ is the difference between the mean AIC_{norm} of M_i and the minimum mean AIC_{norm} among all the candidate models.

Statistical analysis and visualization

Due to log-transformation in the entropy equations, the distributions of most of the entropy-based metrics are highly skewed (**Figure S2**). Accordingly, we primarily employed a nonparametric test, a two-sample Kolmogorov-Smirnov test with a null hypothesis that the two samples are from the same continuous distribution. The test statistic *D* for this test measures the difference between the CDFs of the two distributions.

To avoid confounds with the data size and the computed entropy values, we kept the same bin size (number of trials) from each block for computing the metrics. Specifically, we selected the last N trials of each block while ensuring that 10 trials have passed after the start of each block. Blocks that had too few trials to compute reliable entropy values were removed from the analysis. To minimize the number of excluded blocks while keeping its proportion consistent across datasets, we set N = 30 for the mouse and N = 30

20 for the monkey dataset. This removed 511 blocks out of 3768 (13.56%) for mice, and 36 blocks out of 251 (14.34%) for monkeys.

Finally, as an auxiliary to Kolmogorov-Smirnov test, we also used a permutation test to compare the entropy-based metrics between two given conditions. More specifically, for each permutation, we randomly assigned blocks (without replacement) into two groups with sample size n_1 or n_2 , corresponding to the total number of blocks in the two conditions that are being compared. A single entropy-based metric was computed for each type of blocks by concatenating all trial vectors from the sampled permutation. For consistency, we excluded the same 10 trials after the start of each assigned block. Test statistics for each sample were then computed by taking the difference in metrics between the first and the second condition. We computed the p-value by computing the proportion of sampled test statistics that are as extreme as the observed value.

All data analysis was performed in MATLAB 2020b (Mathworks). All violin plots in the figures were created using an adapted version of a custom toolbox *al_goodplot* in MATLAB (Legouhy, 2022). In the violin plots, the rectangles within the kernel distributions indicate the 25th, 50th (black horizontal line), and 75th percentiles, and circles indicate the mean of the data. For the visualization of the entropy time course, we used confidence intervals from the basic bootstrapping method which computes the difference between observed and sampled test statistics (David & Hinkley, 1997). As mentioned above, for determining statistical significance, a non-parametric test (Kolmogorov-Smirnov test) was primarily used. A parametric t-test was used where appropriate, after verifying that the sample distribution was symmetric and approximately normal. All tests were two sided and alpha was set at 0.05, except otherwise mentioned.

Results

To study how uncertainty related to different types of regularities in the reward environment affects learning and choice behavior, we used multiple metrics based on information theory (**Figure 1**) to reanalyze data from two dynamic foraging tasks in mice and monkeys. In both tasks, animals were trained to choose between two alternative choice options (left and right port for mice, and two abstract pictures for monkeys; **Figure 2A-B**) in order to obtain reward. Reward, a drop of water for mice or a drop of juice for monkeys, was delivered probabilistically and independently of the unchosen option (see below for more details). After a variable number of trials (a block) the reward probability changed for the two options without any cue to the animals, often resulting in a switch between the better and worse options (sometimes referred to as a reversal). The length of each block changed randomly from one block to the next.

In the mouse experiment, rewards were baited such that if reward was assigned on a given side and that side was not selected, reward would remain on that side until the next time that side was selected. The monkey experiment did not have a baiting mechanism, but instead, instantaneous reward probability increased for the unchosen option such that selection of the unchosen option became more rewarding as time since an option was chosen increased. The reward probability for a given option fell back to the assigned baseline probability with every choice of that option, irrespective of whether that choice was rewarded or not.

Overall, both mice and monkeys learned reward associations and their choice closely tracked the reward schedules, reflected in the close match between the probability of choosing the better option, P(Better), and the relative probability of reward for that option (**Figure 2C-D**). As expected, when animals chose the better option more consistently, conditional entropy of reward-dependent strategy (ERDS) was lower (**Figure 2E-F**). This was also reflected as a negative correlation between P(Better) and ERDS (Spearman's correlation, r = -0.647, p = 0), option-dependent strategy (EODS; r = -0.836, p = 0), and baseline entropy of strategy (H(str); r = -0.675, p = 0). In monkeys, however, the relationship between performance and ERDS was not significantly correlated (r = -0.0185, p = .788), whereas higher performance was associated with lower EODS only (r = -0.400, $p = 1.20 \times 10^{-9}$). Interestingly, H(str) was positively correlated with performance (r = 0.315, $p = 2.50 \times 10^{-6}$) indicating that an overall random strategy facilitated the performance in monkeys.

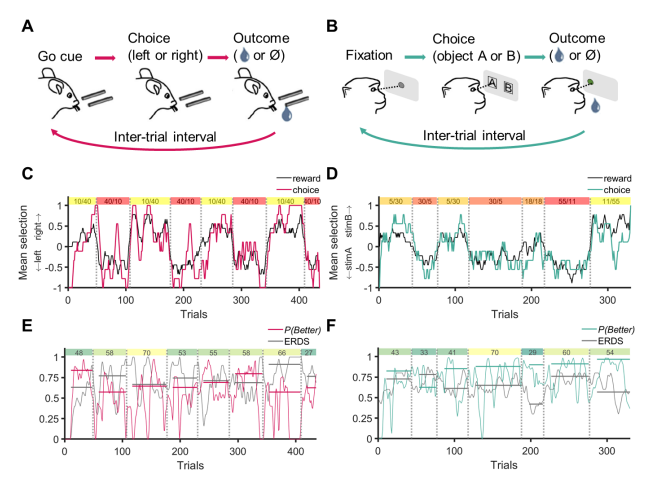


Figure 2. Experimental paradigms and example behavioral results in mice and monkeys. (A, B) Timeline of a single trial during experiments in mice (A) and monkeys (B). To initiate a trial, mice received an olfactory go cue (or no-go cue in 5% of trials), and monkeys fixated on a central point. Animals indicated a choice by licking (mice) or making a saccadic eye movement (monkeys) toward their choice among the two options (left/right port for mice and two abstract pictures for monkeys). Reward (a drop of water for mice and a drop of juice for monkeys) was delivered stochastically based on dynamically assigned reward probability on the chosen option. (C, D) Average choice and reward outcomes using a moving window of 10 trials for a representative session in mice (C) and monkeys (D). Numbers in the colored boxes indicate reward probabilities (in %) of two options within each block. (E, F) Time course of performance and consistency in reward-dependence strategy in example sessions in mice (E) and monkeys (F). Performance (probability of choosing the better option, P(Better)) and conditional entropy of reward-dependent strategy (ERDS) are computed from a moving window of 10 trials from the same example sessions. Numbers in the colored boxes show the length L (number of trials) of each block. Horizontal lines show the overall metrics during each block, computed from all trials in the block (excluding ten trials after block switch).

Entropy-based metrics capture changes in behavior due to block switches

To examine animals' response to changes in reward schedule caused by block switches, we next computed different behavioral metrics before and after those events. In order to determine whether the metrics changed significantly after block switches, we used a paired-sample t-test for each pair of metrics around block switches. As expected, the overall performance decreased after block switches for both mice (**Figure 3A**; t(2992) = 27.6, $p = 1.41 \times 10^{-149}$; Cohen's d = 0.52) and monkeys (**Figure 3D**; t(139) = 6.13, $p = 8.48 \times 10^{-9}$; Cohen's d = 0.499). The decrease was more pronounced for mice because before block switches mice exhibited stronger selection of the better side (M = 0.744, SD = 0.204) than monkey selection of the better stimulus (M = 0.617, SD = 0.11).

Comparisons of entropy-based metrics in mice revealed that block switches increased inconsistency in reward-dependent choice strategy (ERDS) (**Figure 3B**; t(2992) = -10.12, $p = 1.07 \times 10^{-23}$; Cohen's d = -0.165) mainly by increasing the overall entropy in choice strategy (stay vs. switch) measured by H(str) (**Figure 3A**; t(2992) = -10.01, $p = 3.33 \times 10^{-23}$; Cohen's d = -0.165), as there was no evidence for a change in the link between reward outcome and adopted choice strategy measured by mutual information of reward-dependent choice strategy (MIRS) (**Figure 3B**; t(2992) = -0.32, p = 0.750; Cohen's d = -0.01). Similarly, mice became less consistent in how they stayed on or switched from the better option measured by EODS (**Figure 3C**; t(2992) = -11.8, $p = 1.95 \times 10^{-31}$; Cohen's d = -0.197), whereas the link between selected option and adopted choice strategy (measured by MIOS) became weaker after block switches (**Figure 3C**; t(2992) = 5.23, $p = 1.80 \times 10^{-7}$; Cohen's d = 0.093).

In contrast, comparisons of entropy-based metrics in monkeys showed that block switches only made the link between reward outcome and choice strategy (measured by MIRS) weaker (**Figure 3E**; t(139) = 2.41, p = .0171; Cohen's d = 0.174) without any changes in reward-dependent choice strategy (**Figure 3E**; t(139) = -0.82, p = .411; Cohen's d = -0.049) or the overall choice strategy, H(str) (**Figure 3D**; t(139) = 1.73, p = .0862; Cohen's d = 0.139). Similarly, the link between the selected option and adopted choice strategy (measured by MIOS) became weaker right after block switches (**Figure 3F**; t(139) = 5.49, $p = 1.85 \times 10^{-7}$; Cohen's d = 0.435) with a noticeable but not statistically significant change in option-dependent choice strategy, EODS (**Figure 3F**; t(139) = -2.04, p = .0429; Cohen's d = -0.16).

The similar and contrasting effects of block switches in the two species could also be seen from the time course of behavioral metrics over time (**Figure 3G-J**). Comparing the consistency of choice behavior in mice and monkeys around block switches, we observed that monkeys' behavior was more random and less predictable by previous

reward outcome or choice. Kolmogorov-Smirnov tests between mouse and monkey data revealed significantly higher H(str) (**Figure S2D**; D = 0.441, $p = 5.15 \times 10^{-35}$), ERDS (**Figure S2B**; D = 0.276, $p = 5.85 \times 10^{-1014}$), and EODS (**Figure S2C**; D = 0.361, $p = 1.19 \times 10^{-23}$) in monkeys. While MIOS did not significantly differ between two species (**Figure S2F**; D = 0.0944, p = 0.0517), MIRS was slightly higher in monkeys (**Figure S2E**; D = 0.231, $p = 5.93 \times 10^{-10}$). This opposing patterns of MIRS and ERDS across species can be explained by noticing that the larger shared information between reward outcome and subsequent choice strategy for monkeys than mice was not strong enough to result in more consistent reward-dependent strategy (measured by ERDS) in monkeys perhaps due to an overall more random choice strategy (H(str)) in monkeys.

Comparing the entropy metrics within each species using paired t-test, we found that mice exhibited an option-dependent strategy more so than a reward-dependent one: EODS tended to be lower than ERDS (t(3226) = 11.49, $p = 5.48 \times 10^{-30}$; Cohen's d = 0.202) and MIOS tended to be higher than MIRS (t(3226) = -11.49, $p = 5.48 \times 10^{-30}$; Cohen's d = -0.202). In contrast, monkeys showed more of reward-dependent strategy compared to an option-dependent one, as indicated by lower ERDS than EODS (t(214) = -2.36, p = 0.0193; Cohen's d = -0.161) and higher MIRS than MIOS (t(214) = 2.36, p = 0.0193; Cohen's d = 0.161).

Effects of environmental regularities and uncertainty on behavior

One of the main aims of this study was to investigate how animals' choice behavior was affected by regularities and accompanied uncertainty in the reward environment. To answer this question, we first ran exploratory regression analyses to predict behavioral metrics based on a few experimental parameters (**Figures S3, S4**). This included the difference in reward probabilities for the better and worse choice options (P_B and P_W) reflecting uncertainty in determining the better option, the sum of reward probabilities measuring richness of the reward environment, and the block length in the previous and current blocks, aiming to measure whether animals developed an expectation about block lengths and the stability of reward environment. We note, however, that regression may not be the most suitable analysis because of the non-linear nature of entropy-based metrics (**Figure S2**). Nevertheless, these regression analyses pointed to the effects of a difference in reward probabilities and block length on multiple entropy-based metrics, indicating that certain regularities in the environment affected learning and choice behavior.

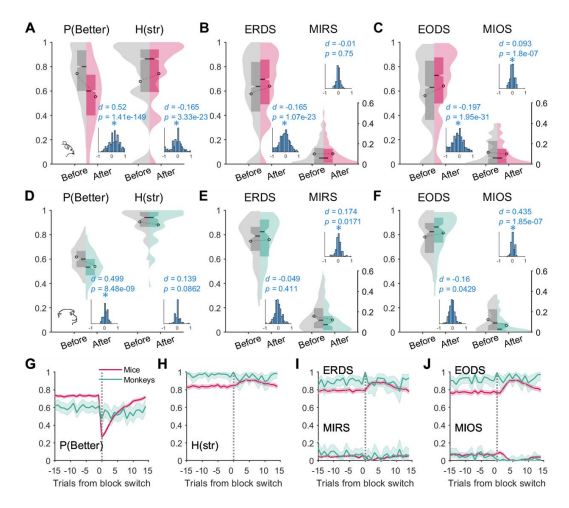


Figure 3. Changes in performance and behavioral metrics around block switches. (A-F) Violin plots show distributions of performance and entropy-based metrics computed from 15 trials before and after block switches in mice (A-C) and monkeys (D-F). In the violin plots, the rectangles within the kernel distributions indicate 25th, 50th (median, black horizontal line), and 75th percentiles. Circles indicate the mean of the data. Insets show histograms of paired difference between metrics before and after each block switch. Reported values are Cohen's d and p-values based on paired-sample t-test comparing before and after block switches (two-sided test, corrected for multiple comparison of two). (G-J) Time course of different metrics around block switches. Plots show running averages of metrics computed from the same trials in time relative to block switches (vertical dotted line) across all blocks of the experiment. Shaded error bars indicate 95% confidence intervals from bootstrapping method using N = 10,000 samples for each trial point. Overall, mice had significantly lower H(str), ERDS, EODS, and MIRS than monkeys.

To better understand these effects, we next grouped blocks based on the block length and the difference in reward probabilities, and computed different metrics for each group of blocks. First, we divided blocks based on the difference in reward probabilities for the better and worse options and categorized blocks into more uncertain and less *uncertain* blocks based on the median split of $P_B - P_W$ in each subject (see **Methods** for more details). The difference in reward probabilities measures uncertainty in determining the better choice option. Blocks that had larger $P_B - P_W$ than the median were categorized as less uncertain and those with smaller $P_B - P_W$ as more uncertain blocks. Second, we also divided blocks depending on whether the current block length was longer than expected or not. More specifically, we assumed that animals developed an expectation of block switches by estimating a value for the length L of a block at each block i, denoted $E[L_i]$, updated at every block switch. This update was done using the block length prediction error (BLPE) equal to the difference between the actual and expected block lengths (see **Methods** for more details). Negative BLPE corresponds to a block that was shorter than expected (SE blocks) based on the animal's experience whereas positive BLPE indicates that a block was *longer than expected* (*LE* blocks).

With regard to uncertainty in determining the better option, our hypothesis was that increased uncertainty should result in slower learning and/or more stochastic behavior. In addition, we hypothesized that when blocks are longer than expected (LE blocks), choice or adopted choice strategy would be more random as animals expect changes in reward probabilities more strongly and in the absence of any changes in the environment. Finally, expectation of block switches could interact or depend on the level of uncertainty in the environment.

Supporting these hypotheses, we found that entropy-based metrics were systematically

affected by both uncertainty and expectation of block switches, but differently for mice and monkeys. In mice, entropy of choice (H(str); **Figure 4A**), reward-dependent choice strategies (ERDS; **Figure 4B**), and option-dependent choice strategies (EODS; **Figure 4C**) were significantly larger in the more uncertain environments, with no significant difference between mutual information (MIRS or MIOS; **Figure 4B, C**). Considering the link between entropies, conditional entropies and mutual information (ERDS = H(str) $^-$ MIRS and EODS = H(str) $^-$ MIOS), these results suggest that the observed increase in H(str) could be due to changes in the effects of reward feedback on choice. In monkeys, however, there was a trend toward a decrease (instead of increase) in the entropy of choice (H(str)), suggesting that animals' strategy became more consistent when reward probabilities were closer to each other; i.e., when the environment was more uncertain (**Figure 5A**; Kolmogorov-Smirnov test; D = 0.277, p = .0250). Post-hoc analysis revealed that this decrease in H(str) in monkeys was due to higher probability of

switching in the more uncertain environment (M = 0.668, SD = 0.11) compared to the less uncertain one (M = 0.508, SD = 0.142). Therefore, monkeys increased their exploration in more uncertain environments.

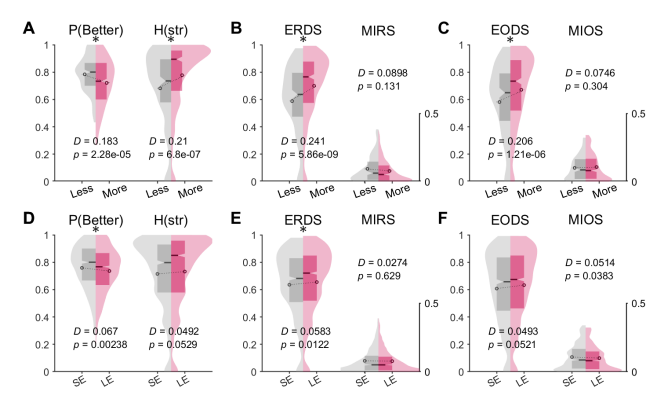


Figure 4. Effects of uncertainty and expectation of block switches on choice behavior in mice. (A-C) Effects on uncertainty on choice behavior. Violin plots show the distributions of performance and entropy-based metrics computed from different types of trials according to uncertainty of the reward schedule. All metrics are computed using the last 30 trials in each block. Asterisks show results of two-sample Kolmogorov-Smirnov test with a null hypothesis that the two samples are from the same continuous distribution (two-sided test, corrected for multiple comparison of two). More uncertainty results in more inconsistent choice driven by more stochasticity. (D-F) Effects of block switch expectation on choice behavior. SE and LE indicate shorter than and longer than expected blocks, respectively, corresponding to trials before and after the animal expected a block switch. Violin plots show the distributions of performance and entropy-based metrics computed from different types of trials according to block length prediction error. Overall, mice adopted less consistent choice strategies when blocks were longer than expected, especially in more uncertain environments.

With regard to expectation of block switches, we found that in mice, ERDS were significantly increased for LE compared to SE blocks (**Figure 4E**; Kolmogorov-Smirnov test; D = 0.0583, p = .0122). H(str) and EODS followed the similar trend, but the

increase was not significant (Kolmogorov-Smirnov test; H(str): D = 0.0492, p = .0529; EODS: D = 0.0493, p = .0521; **Figure 4D**, **F**). The changes in mutual information (MIRS, MIOS) were small or insignificant (**Figure 4E**, **F**). The effect of expectation of block switches on ERDS in the absence of any clear changes in H(str) and MIRS suggest multiple forms of adjustments including changes in learning and stochasticity in choice. Similarly, expectation of block switches increased ERDS in monkeys (Kolmogorov-Smirnov test; D = 0.201, p = .0227; **Figure 5D**) and this was accompanied with a moderate but not significant decrease in MIRS (Kolmogorov-Smirnov test; D = 0.183, p = .0482; **Figure 5D**). Another post-hoc analysis looking at win-stay/lose-switch tendencies of monkeys revealed that this observed change in reward-dependent strategy in monkeys mainly resulted from reduced probability of win-stay from SE (M = 0.627, SD = 0.22) to LE blocks (M = 0.564, SD = 0.196). Lose-switch tendency was overall higher than win-stay but was largely the same between SE (M = 0.738, SD = 0.158) and LE blocks (M = 0.733, SD = 0.156).

Finally, to test possible interactions between uncertainty and reward expectation, we further divided SE and LE blocks according to uncertainty (more or less uncertain) and found that the major effect of block switch expectation was mediated by uncertainty in the reward probability. We found that the differences between SE and LE blocks were mainly significant in more uncertain blocks but not in less uncertain ones (**Figure S5**). This suggests that expectation was more pronounced in more uncertain reward environments because in less uncertain environments, reward feedback could signal changes in the reward schedule more easily.

To compare behavioral changes to actual and expected block switches, we performed further analyses focusing on the LE blocks, which allowed us to examine changes to expected block switches. These blocks are longer than expected (L > E[L]) and therefore have expected block switches situated *prior* to the actual block switches. Accordingly, we computed the changes in the behavioral metrics after expected block switches using paired-sample t-tests, similar to the analyses presented in **Figure 3**.

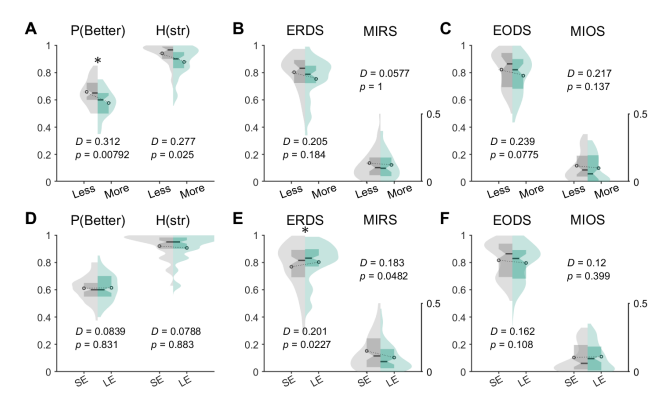


Figure 5. Effects of uncertainty and expectation of block switches on choice behavior in monkeys. (A-C) Effects on uncertainty on choice behavior. Violin plots show the distributions of performance and entropy-based metrics computed from different types of trials according to uncertainty of the reward schedule. All metrics are computed using the last 20 trials in each block. Asterisks show results of two-sample Kolmogorov-Smirnov tests with a null hypothesis that the two samples are from the same continuous distribution. (D-F) Effects of block switch expectation on choice behavior. SE and LE indicate shorter than and longer than expected blocks, respectively, corresponding to trials before and after the animal expected a block switch. Violin plots show the distributions of performance and entropy-based metrics computed from different types of trials according to block length prediction error. Overall, monkeys adopted less consistent reward-dependent choice strategies when blocks are longer than expected, especially in more uncertain environments.

We found that mice reacted to actual and expected block switches similarly; both of these switches were accompanied with increased H(str), ERDS, EODS, and decreased MIOS (**Figure S6**). Among these metrics, EODS showed the largest changes as suggested by the effect sizes (**Figure S6C**; actual block switch: Cohen's d = -0.209, $p = 6.37 \times 10^{-14}$; expected block switch: Cohen's d = -0.225, $p = 7.24 \times 10^{-24}$). Such significant effects on both EODS and MIOS suggest that the observed adjustment in behavior after perceived block switches could be mainly due to changes in the option-dependent strategy, as the animals anticipate a switch between the better and worse choice

options. Monkeys also similarly reacted to actual and expected block switches with decreased MIRS (**Figure S7E**; actual block switch: Cohen's d = 0.322, p = .0131; expected block switch: Cohen's d = 0.346, p = .00896). ERDS increased after expected block switches (**Figure S7B**, right; Cohen's d = -0.411, p = .00275) more so than after actual block switches (**Figure S7B**, left; Cohen's d = -0.183, p = .197).

These results indicate that monkeys adapted to perceived block switches mainly through changes in the reward-dependent strategy, contrary to mice which mostly showed changes in option-dependent strategy. Overall, the means of paired differences did not significantly differ between actual and expected switches for most of the metrics, as determined by independent-sample t-test even when using the most liberal alpha level at 0.05 (insets in **Figure S6, S7**). Therefore, above analyses based on behavioral metrics revealed that both mice and monkeys showed consistent adjustments to expected and actual block switches but monkeys also exhibited some discrepancies in their adjustments (e.g., MIOS in **Figure S7F** inset).

Computation of expected block length (E(L)) and resulting block length prediction error (BLPE) depend on the parameter γ_E that controls update of E(L) (see Eq. 7 in **Methods**). In all of the above analyses related to the effect of block switch expectation, we set $\gamma_E = 0.1$. Nonetheless, we also tested whether and how observed differences in entropy-based metrics between SE and LE blocks depend on the value of γ_E (**Figures S8** and **S9**). Although we found differences for a wide range of γ_E values, they tended to be more pronounced for smaller values of γ_E , corresponding to integration of block length over longer timescales. This observation is in line with the intuition that capturing regularities of the environment requires estimation based on multiple block switches.

Finally, we conducted an additional permutation test to compare entropy-based metrics across conditions (*less* vs. *more uncertain*, or *SE* vs. *LE*; see **Methods** for more details). This was done to avoid a possible confound due to small sample size for computation of entropy-based metrics that causes bias toward smaller values (though this bias should impact all conditions similarly as equal numbers of trials were used for different conditions). Overall, results from the permutation test were consistent with and in some cases more likely to be significant than those based on the Kolmogorov-Smirnov test. As for the effect of reward uncertainty in mice (**Figure 4A-C**), we confirmed the same increase in H(str) (**Figure S10A**), ERDS (**Figure S10C**), and EODS (**Figure S10G**), with additional significant decrease in MIRS in more uncertain blocks (**Figure S10E**). The added significant effect of MIRS supports our above interpretation that the observed increase in H(str) is most likely due to the changes in the effects of reward feedback on choice strategy. For the effect of uncertainty in monkeys, we found that H(str) (**Figure S10B**), ERDS (**Figure S10D**), EODS (**Figure S10H**), and MIOS (**Figure S10J**) were all significantly smaller in more uncertain environments. The results

of the Kolmogorov-Smirnov test were in the same direction but not significant (**Figure 5A-C**). With regard to the effect of block switch expectation, mice had significantly larger EODS (**Figure S11G**) and smaller MIOS (**Figure S11I**) in LE blocks, whereas monkeys showed significant reduction in MIRS only (**Figure S11F**). These results are consistent with the above findings that mice react to perceived block switches through changes in option-dependent strategy (**Figure S6**), whereas monkeys react through changes in reward-dependent strategy (**Figure S7**).

Reinforcement learning models reveal underlying mechanisms in response to different types of uncertainty

To reveal possible mechanisms by which uncertainty and expectation of block switches affect choice behavior and learning, we used simple RL models (RL1, RL1_{decay}, RL2, RL2_{decay}) to fit data in different blocks of the experiments as described above (**Methods**). The first two models had a single learning rate (RL1), while the other two had separate learning rates for rewarded and unrewarded trials (RL2). We also tested how the fit of data can be improved by including forgetting in the estimated value for the unchosen option or action (captured by decay rate γ in RL_{decay} models).

We found the winning model for fitting mouse data to be RL1 $_{decay}$ that has a single learning rate α , inverse temperature β , and decay rate γ (**Table 1**). The normalized AIC by number of trials (AIC_{norm}) for this model was significantly lower than that of the second-best model, RL2 $_{decay}$ (paired-sample t-test, t(3767) = 12.67, $p = 5.21 \times 10^{-36}$; Cohen's d = 0.2063). For the monkey data, the best model was the simplest model, RL1, however, the AIC_{norm} for this model was not significantly lower than that of the second-best model, RL2 (paired-sample t-test, t(250) = 1.41, p = 0.161; Cohen's d = 0.089). Overall, we found that a single learning rate model provided a better fit to both species than separate learning rate models. Mouse data tended to prefer models with decay terms. In contrast, monkeys' choice data were better fit with models without decay, suggesting a different form of reward value estimation from mice.

Accordingly, we analyzed the fitted parameters from the winning model of each species (RL2 $_{decay}$ and RL1 for mice and monkeys, respectively). We hypothesized that a smaller learning rate could be advantageous in more uncertain environments because it allows animals to obtain a more precise estimate of reward value in such environments (cf. Figure 1B in Farashahi et al. 2017a). With regard to the expectation of block switches, we hypothesized that when blocks are longer than expected (LE), the increased randomness in choice behavior would result from an increase in the stochasticity of choice and/or decrease in learning, as suggested by results in the previous section. Higher stochasticity in choice can be reflected in the lower inverse temperature β . In contrast, decreased learning could be mediated by decreased learning rate and/or

faster decay rate for the unchosen option, both of which could reduce the effects of previous reward feedback on choice behavior.

Model	RL1	RL1 _{decay}	RL2	RL2 _{decay}
Parameters -LL AIC norm d (full) d (best) Akaike w.	α, β 0.4825 1.078 (0.0393*) 0.1223* 1000 0.2589* 500 24.6% (4) 0 1 2	α , β , γ 0.4215 1.013 (-0.0259*) -0.2063* 1000 0 25.41% (1) 0 1 2	$\alpha_{+}, \alpha_{-}, \beta$ 0.4417 1.053 (0.0144*) 0.07238* 1000 0.2432* 500 24.91% (3)	$\alpha_{+}, \alpha_{-}, \beta, \gamma$ 0.4061 1.039 (0) 0 0.2063* 25.09% (2)
Model	RL1	RL1 _{decay}	RL2	RL2 _{decay}
Model Parameters -LL AIC norm	RL1 α, β 0.6457 1.378 (-0.0488*)	RL1 _{decay} α , β , γ 0.6436 1.417 (-0.00985)	RL2 α ₊ , α ₋ , β 0.627 1.384 (-0.043*)	RL2 _{decay} $\alpha_{+}, \alpha_{-}, \beta, \gamma$ 0.6269 1.426 (0)

Table 1. Reinforcement learning models used to fit choice data, their parameters, and their goodness-of-fit. Reported are parameters of a given model and goodness-of-fit, given by negative log-likelihood (-LL) and AIC normalized by the number of trials in each block (AIC_{norm}). Values reported in parentheses next to AIC_{norm} are the difference in AIC_{norm} of a given model and the full model (RL2_{decay}). Asterisks indicate the significance of this difference indicated by paired-sample t-test (p < 0.05/6, multiple comparison corrected). d(full) indicates Cohen's d value from paired-sample t-tests between a given model and the full model (RL2_{decay}), with the asterisk indicating significance. d(best) indicates Cohen's d value of paired-sample t-tests between a given model and the best model determined by the lowest AIC_{norm}. Akaike w = Akaike weights, indicating the probability that a given model is the best model. Integers in parentheses are ranks of a given model. Insets show histograms of AIC_{norm} from the mouse (top row, red) and monkey data (bottom, teal).

We found that in mice, learning rate (α) was significantly reduced from less (M = 0.372, SD = 0.252) to more uncertain (M = 0.278, SD = 0.249) blocks, supporting our hypothesis (**Figure 6A**). We also observed a significantly smaller decay rate (γ_{decay}), corresponding to less forgetting of the reward estimate for the unchosen option in more uncertain environments (**Figure 6C**). Overall, these results indicate that more uncertainty resulted in slowing down of value updates in mice. In contrast, the effect of block switch expectation was mediated mostly by changes in inverse temperature parameter (**Figure 6E**). Specifically, when blocks were longer than expected (LE),

inverse temperature β was significantly reduced corresponding to more stochasticity in choice (Kolmogorov-Smirnov test, D = 0.129, $p = 4.85 \times 10^{-14}$). At the same time, longer than expected blocks was accompanied by decreased learning rate and increased decay rate (**Figure 6D-E**). These opposing effects on value updates are interesting as they suggest that when expecting block switches, mice slowed down learning from chosen option while speeding up the forgetting of the unchosen option.

Furthermore, dividing LE and SE blocks into more or less uncertain conditions, we found the same mediating effect of uncertainty as revealed by behavioral metrics. That is, significant changes in the estimated model parameters due to block switch expectation were only observed for more uncertain blocks, consistent with the interpretation that expectation is more pronounced in more uncertain environments. More specifically, the reduction in β for LE blocks was significant only for the more uncertain environment (blue bars in **Figure 6E** inset; Kolmogorov-Smirnov test, D = 0.219, $p = 6.17 \times 10^{-8}$) but not for less uncertain one (gray bars in **Figure 6E** inset; Kolmogorov-Smirnov test, D = 0.105, p = .437). Similarly, an increase in γ_{decay} for LE blocks was only observed for more uncertain blocks (blue bars in Figure 6F inset; Kolmogorov-Smirnov test; D = 0.16, p = .00018) but not for less uncertain blocks (gray bars in **Figure 6F** inset; Kolmogorov-Smirnov test, D = 0.10, p = .503). Interestingly, learning rate did not show this pattern as there was no significant decrease for LE blocks in less uncertain (gray bars in **Figure 6D** inset; Kolmogorov-Smirnov test, D = 0.191, p = .0152) or more uncertain environments (blue bars in **Figure 6D** inset: Kolmogorov-Smirnov test, D = 0.118, p = .0151). This suggests a modulatory effect of local uncertainty about the better option on uncertainty about the expectation of block switches (which could take longer to estimate) via changes in stochasticity in choice and the decay of value estimates for the unchosen option.

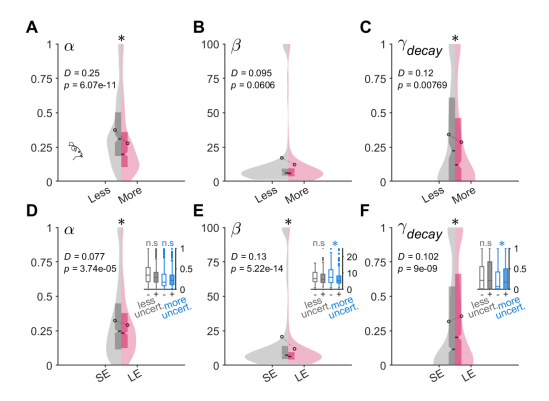


Figure 6. Mechanisms underlying effects of uncertainty and expectation of block switches revealed by fitting of mouse data by reinforcement learning models. Plots show distributions of fitted parameters from RL1_{decay} model to mice choice data. categorized by the level of uncertainty (A-C) or expectation of block length (D-F). α = learning rate for rewarded and unrewarded trials; β = inverse temperature or choice consistency; γ_{decay} = decay rate in value estimate for the unchosen option. SE: shorter than expected, LE = longer than expected blocks. Asterisks show results from a twosample Kolmogorov-Smirnov test. All tests are two-sided, corrected for multiple comparisons of three (number of free parameters). Box plots in the insets show further division of the data into less (gray) or more (blue) uncertain blocks. Asterisks are results of two-sample Kolmogorov-Smirnov tests, corrected for multiple comparisons of six. Interaction effect also exists for the significant RL parameters. Overall, mice responded to the more uncertain reward environment by decreasing the learning rate and slower decay rate for unchosen value. In addition, they responded to trials after an expected block switch by decreasing learning rate, increasing in stochasticity in choice, and having faster decay of unchosen value.

In monkeys, we found evidence that greater uncertainty about the better option resulted in significant changes in the estimated model parameters. More specifically, mean learning rate reduced from less (M = 0.443, SD = 0.35) to more uncertain (M = 0.381, SD = 0.33) environments, similar to what we found in mice (**Figure 7A**; Kolmogorov-Smirnov test, D = 0.166, p = .273). In contrast to mice, however, the effect of block switch expectation was reflected in a significant decrease in the learning rate only

(**Figure 7C**; Kolmogorov-Smirnov test, D = 0.193, p = .0204). This suggests that monkeys slowed down learning, perhaps because around block switches reward feedback was not reliable. Similar to mice, this effects on learning was more pronounced in less uncertain environment (gray bars in **Figure 7C** inset; Kolmogorov-Smirnov test, D = 0.323, p = .0657) compared to more uncertain one (blue bars in **Figure 7C** inset; Kolmogorov-Smirnov test, D = 0.275, p = .123), although not statistically significant. Inverse temperature was not affected by expectation of block length (**Figure 7D**; Kolmogorov-Smirnov test, D = 0.072, p = .908). Overall, these results demonstrate that mice and monkeys respond to regularities and uncertainty in the reward environment via different mechanisms.

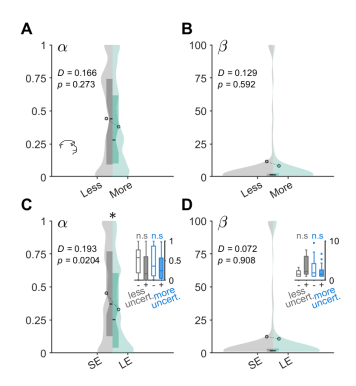


Figure 7. Mechanisms underlying effects of expectation of block switches revealed by fitting of monkey data by reinforcement learning models. Plots show distributions of fitted parameters from RL models to mice choice data, categorized by the effect of expected uncertainty (A-B) or expectation of block length (C-D). α = learning rate for rewarded and unrewarded trials; β = inverse temperature or choice consistency. SE: shorter than expected, LE = longer than expected blocks. Asterisks show results from a two-sample Kolmogorov–Smirnov test. All tests are two-sided, corrected for multiple comparisons of two (number of free parameters). (A-B) No parameters were significantly affected by expected uncertainty of the reward environment, similar to the entropy measures. (C-D) Monkeys responded to trials after an expected block switch by decreasing learning rate. Box plots in the insets show further division of the data into less (gray) or more (blue) uncertain blocks. Asterisks are results of two-sample Kolmogorov-Smirnov tests, corrected for multiple comparisons of four. Interaction effect was not observed for the significant RL parameters.

Discussion

How the brain learns from regularities in the environment and adapts to an uncertainty associated with those regularities are central questions in cognitive and behavioral neuroscience. Here, we used multiple metrics based on information theory to investigate the effects of different types of uncertainty on learning and choice behavior. We found that rodents and primates differed in how they adapted to uncertainty in the environment. Mice employed option-dependent strategies more often, after slowly identifying the better option, and chose to stay on the better rewarding option to exploit rewards. In contrast, monkeys employed reward-dependent strategies more often while switching between choice options more frequently. Fit of choice data using RL models showed that this difference in strategy was mediated via higher learning rates in monkeys, consistent with the interpretation that such strategy requires more weighting of immediate reward outcomes. In addition, inverse temperature representing choice consistency was higher in mice, as they consistently stayed on the better option more than monkeys.

Some of the similarities and differences between mice and monkeys can be explained as below. First, mice exhibited an overall more consistent choice strategy, indicated by the lower values of H(str), ERDS, and EODS than those of monkeys. The higher performance (P(Better)) in mice suggests that the lower entropy was mainly due to these animals staying on the better option more consistently than monkeys. We note that this difference could be in part derived from slight differences in tasks rather than reflecting some intrinsic intra-species differences. In the mouse task, the same reward schedule was used throughout each of the whole session, while the reward probabilities reversed between two sides at every block switch (Figure 2C). In contrast, in the monkey task, reward schedules were randomly assigned at every block, making it more challenging to determine the better option (Figure 2D). In other words, the mouse task facilitated the distinction between the better and option by assigning just two reward probabilities for a given session, whereas the reward rate for each option had to be estimated at every block in the monkey task.

Furthermore, the difference in the reward rates between two options (ΔP) was smaller in the monkey experiment, suggesting that the task difficulty was greater for monkeys. In effect, this led to a more switching behavior in the monkeys compared to mice. Specifically, the monkeys showed a short-term tendency to switch between options and a longer-term tendency to repeat choices for the same options—a pattern that can be revealed by inclusion of a choice-history variable in a logistic regression on the monkeys' choices (Grabenhorst et al., 2019) and that has previously been observed in monkeys performing matching tasks (Lau & Glimcher, 2005). The monkeys' higher tendency to switch may have also resulted from the absence of a 'changeover delay', a

manipulation sometimes used to encourage matching behavior, whereby reward would only be delivered on the second trial following a switch between options (Sugrue et al., 2004). Interestingly, H(str) was positively correlated with performance P(Better), indicating that a more equal mix of stay and switch facilitated winning of rewards in monkeys. In contrast, lower H(str) predicted better performance in mice, as they primarily employed a staying strategy to exploit the better rewarding option. Future empirical work should test the influence of task conditions on these observed species differences.

The overall difference between mice's and monkeys' adoption of stay vs. switch strategies could also explain how they adjusted to uncertainty. More specifically, the greater uncertainty in the reward environment (measured by ΔP) had different effects on mice and monkeys. Mice adjusted their strategies by incorporating switching strategies more often, which resulted in increased randomness in strategies reflected by higher entropies (H(str), ERDS, and EODS). Monkeys similarly increased their switching behavior, but even more so than their baseline strategy (which was already very random) such that the overall H(str) was reduced, signaling higher consistency.

The difference in the baseline entropy of strategy (H(str)) between mice and monkeys also partly accounts for the observed difference in their adjustment to expected block switches. The exploitation strategy characterizing the mice behavior – consistently staying on the more rewarding option – is a form of option-dependent strategy which requires a clear sense or confidence about what the better option is. Therefore, in the anticipation or perception of block switches, the link between previous choice option and the subsequent action would be expected to decline as the animal begins to explore the alternative option. This is what MIOS metric in mice revealed (comparison of before/after metrics in **Figure S6** or **Figure S11I**). In contrast, monkeys primarily depended on reward-dependent strategies. Accordingly, in the event of expected block switches, the link between reward feedback and the subsequent action was most significantly reduced (MIRS in **Figure S6E** and **Figure S11F**), as winning a reward did not lead to staying behavior as consistently as before.

Here, we used the difference between the baseline probability of reward for the better and worse options to quantify expected uncertainty in the environment. This was based on the notion that a larger difference in reward probabilities is associated with less uncertainty in identifying the better option using reward feedback. Alternatively, one could use the variance in objective or subjective reward probability to measure risk in economic sense as such "variance risk" can drive choice behavior and neural activity differently than subjective value (Schultz et al., 2008; Grabenhorst, et al., 2019). In this study, we used uncertainty because in both tasks, reward probabilities are unknown and

must be learned from reward feedback (De Groot & Thurik, 2018; Soltani & Izquierdo, 2019) and examined how such uncertainty influences choice behavior and learning.

Comparing the effects of reward uncertainty (ΔP) and block switch expectation, we found that the effects of block switch expectation were more subtle compared to that of ΔP . That is, for all of our analyses comparing the extent of these two effects (**Figure 4**, **5**; **Figure S10**, **S11**), reward uncertainty had a greater impact on behavior than block switch expectation for both species. This is expected because anticipation of block switches requires longer-term statistics of reward outcomes compared to those related to determining the better option (Atilgan et al., 2022). The animals could, for example, return to their previous strategy after a sufficient number of reward feedbacks have signaled that actual block switch has not occurred yet. Still, the entropy-based measures captured such effects from LE blocks, by applying the assumption that expectation of block switches can have more clear effects toward the end of

blocks. This also demonstrates the utility of the entropy metrics, which are capable of detecting nuanced shifts in choice behavior if measured from appropriate binning of trials.

Finally, the observation that expectation of block switches depends on the level of uncertainty suggests interdependence between different types of uncertainty (Soltani & Izquiredo, 2019; Piray & Daw, 2021). In a recent study, Piray & Daw (2021) provided a normative account that uncertainty (stochasticity) reduces learning rate, whereas volatility increases it. Regarding uncertainty, we similarly observed lower mean learning rates for more uncertain conditions. However, on the influence of volatility, previous studies have shown mixed results (Behrens et al., 2007; Donahue and Lee, 2015; Farashahi et al., 2019). Consistent with previous work (Donahue and Lee, 2015; Farashahi et al., 2019), we found decreased rather than increased learning rates in both species. Moreover, we found that the learning rates were not influenced by the mediating effects of greater uncertainty on the block switch expectation.

These results raise a possibility that, while learning rates could be implicated in adjustment to volatile environments, the effect of expectation arising from observation of long-term regularities in the environment might more directly involve other cognitive mechanisms. For example, it has been suggested that active representations of alternative models about the environment ('task set') are stored in long-term memory to be retrieved when the corresponding context reoccurs (Collin & Koechlin, 2012; Soltani & Koechlin, 2022). The neural system for this retrieval likely involves extensive interaction between prefrontal cortex and hippocampus, the latter proposed to be involved in the representation of unexpected uncertainty (Payzan-LeNestour et al., 2013) and one-shot learning (Lee et al., 2015) in humans. Further studies in rodent and

primate could directly probe the causal role for the hippocampus in learning under different forms of uncertainty.

There are limitations and additional considerations related to our findings. Foraging in the wild consists of exposure to time-delays and spatial separation between options, so more naturalistic dynamic foraging tasks could be developed to increase parametric space such that a wider range of values could be explored (Constantino & Daw, 2015; Hall-McMaster et al., 2021). Indeed, the longer time horizon and expansive range size to forage in nonhuman primates compared to rodents influence how the species predict, evaluate, and act on rewards; mediated by a more specialized and expansive prefrontal cortex in the primate brain (Wittmann et al. 2016; Rudebeck & Izquierdo, 2022). Additionally, though we found that stochastic behavior and corresponding entropy metrics were affected by uncertainty of the better option in both mice and monkeys, these effects may be mediated by different neural mechanisms given that mice responded by actions, while monkeys responded to stimuli. Previous work has shown that (freely moving) rodents tend to have greater difficulty learning stimulus-based probabilistic outcomes compared to spatial-based ones (Harris, Aguirre et al., 2021; Chen, Ebitz et al., 2021), whereas monkeys do well in both modalities (Costa, et al., 2016; Rothenhoefer et al., 2017; Taswell et al., 2021). Consistently, different neurons in the dorsolateral prefrontal cortex of monkeys have been shown to track reward uncertainty associated with both stimuli and actions (Grabenhorst et al., 2019). This points to a need to disambiguate the rodents' natural tendency to navigate and learn spatially, versus the influence of task conditions that enhance stable choice such as in this dynamic foraging task.

Conclusion

Together, our results illustrate how higher-order statistics of the environment influence learning and choice strategies. We show that entropy-based metrics can be useful for investigating multiple aspects of adaptive behavior under different forms of uncertainty. By defining the notion of block switch expectation, we showed that shorter than expected blocks and longer than expected blocks were marked by quantitatively different consistencies in behavior as captured by the entropy-based metrics. These findings suggest that mice and monkeys can use reward history to form long-term expectations about the environment and adjust their behavior accordingly. Such expectations of block switches were characterized by increased randomness in choice strategy, similar in its quantity to those induced by the actual block switches. Therefore, our study demonstrates how model-free (entropy-based metrics) and model-dependent (RL models) methods can complement each other to elucidate behavioral adjustments and underlying mechanisms with finer details.

Supplementary Information

The online version contains supplementary material available at XXX.

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Declarations

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Conflicts of interest/Competing interests

The authors declare no conflicts of interests.

Availability of data & materials

The data presented are available upon reasonable request from the authors.

Authors' contribution

Conceptualization of the study: JHW, AS; Mouse experiment conceptualization & data collection: BAB, JYC; Monkey experiment conceptualization & data collection: KT, FG, WS; Analyses of data and illustration: JHW, CGA, AI, AS; Implementation of algorithms, models, analyses, and plots: JHW; Original draft: JHW, CGA, AI, AS; Review & editing: all authors.

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Supplementary Materials

- Figure S1. Distributions of block lengths and reward probabilities in dynamic foraging tasks in mice and monkeys.
- Figure S2. Distributions of main behavioral metrics in mice and monkeys.
- Figure S3. Multiple regression on mice's behavioral metrics.
- Figure S4. Multiple regression on monkeys' behavioral metrics.
- Figure S5. Expectation of block switch depends on the level of uncertainty.
- Figure S6. Mice exhibit similar changes in behavior around actual and expected block switches.
- Figure S7. Monkeys exhibit similar changes in behavior around actual and expected block switches.
- Figure S8. Comparison of different timescales for integration of expectation in mice.
- Figure S9. Comparison of different timescales for integration of expectation in monkeys.
- Figure S10. Effects of reward uncertainty on mouse and monkey behavior shown by permutation test.
- Figure S11. Effects of block switch expectation on mouse and monkey behavior shown by permutation test.

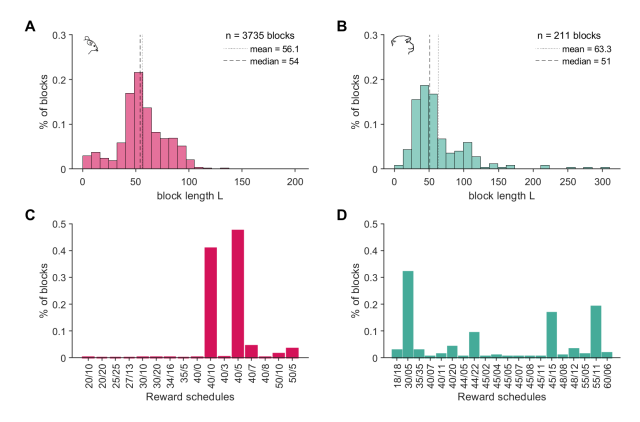


Figure S1. Distributions of block lengths and reward probabilities in dynamic foraging tasks in mice and monkeys. Plots show distributions of block lengths and reward schedules in the mouse (A, C) and monkey (B, D) experiments. The two numbers in the reward schedule correspond to reward probabilities (in percentage) of better and worse options (defined based location for mice and visual stimuli for monkeys) in each block. For some blocks, the two choice options had the same reward probabilities (e.g., 35/35). Total n = 18 mice performed on average 209.3 blocks per each subject. Total n = 2 monkeys performed on average 125.5 blocks (first monkey: 182 blocks, second monkey: 69 blocks). The majority of the blocks (\sim 89%) in mice were either 40/10 or 40/5 schedule. The majority of the blocks (\sim 70%) in monkeys were either 30/5, 45/15, or 55/11 schedule.

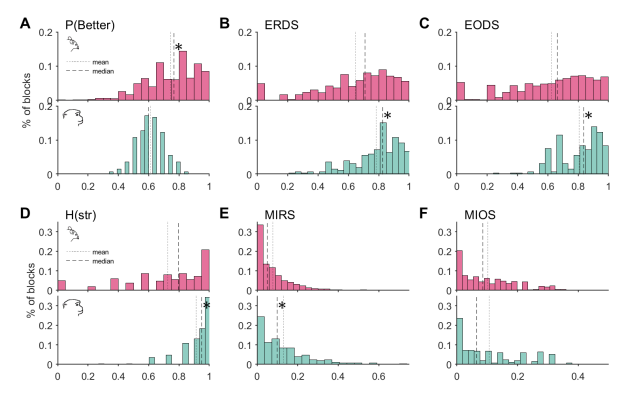


Figure S2. Distributions of main behavioral metrics in mice and monkeys. Plotted are distributions of the behavioral metrics in mouse (top panel, red) and monkey (bottom panel, teal) experiments, including performance or probability of choosing a better option (A), entropy of reward-dependent strategy (B), entropy of option-dependent strategy (C), baseline entropy of strategy (D), mutual information of reward and strategy (E), and mutual information of previous choice option and strategy (F). Asterisk indicates the distribution with significantly larger CDF from intra-species comparison, determined by Kolmogorov-Smirnov test (p < .05). All metrics have minimum values of 0 and maximum of 1. Due to log-transformation in the computation of entropy-based metrics, most of the metrics are highly skewed. To avoid confounds with the data size and the computed entropy values, we kept the same bin size (number of trials) from each block for computing the metrics. For mouse data, metrics are computed from the last N = 30 trials in the block. For monkey data, N = 20. This was chosen to minimize the number of excluded blocks while keeping its proportion consistent across datasets (see **Methods**).

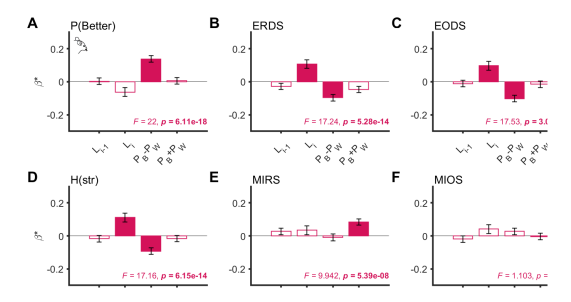


Figure S3. Multiple regression on mice's behavioral metrics. Bar plots show the standardized coefficients of regressors, corresponding to task variables representing different types of uncertainty. The bottom of each panel reports the F-statistic and its pvalue of the regression model. Significant predictors are indicated with a filled color bar. L_{i-1} = previous block length. L_i = length of the current *i*-th block. $P_B - P_W$ = difference between reward probabilities of better (P_B) and worse (P_W) options (see Methods). $P_B + P_W = \text{sum of }$ the two reward probabilities. All behavioral metrics were computed from the last 30 trials of each block, to avoid confounds with data size and computed entropy values by ensuring the same bin size for every block. (A) Response variable in the regression is performance (probability of choosing the better option during each block, P(Better)). Larger $P_B - P_W$ or less expected uncertainty predicted better performance ($\beta^* = 0.156$, $p = 3.32 \times 10^{-15}$). Longer L_i was associated with worse performance ($\beta^* = -0.0539$, p = .0416) but the significance did not survive after correcting for multiple comparisons of four (number of independent variables). (B) Response variable: entropy of reward-dependent strategy (ERDS), equal to H(strlrew). Higher ERDS value indicates a more random response strategy on reward feedback. Longer L_i and smaller $P_B - P_W$ predicted more random reward-dependent strategy (*L*_i: $\beta^* = 0.107$, $p = 5.73 \times 10^{-5}$; $P_B - P_W$: $\beta^* = -0.096$, $p = 1.50 \times 10^{-6}$) (**C**) Response variable: entropy of option-dependent strategy (EODS), equal to H(strlopt). Higher EODS value indicates a more random response strategy on the previous chosen option. Similar to ERDS, Longer L_i and smaller $P_B - P_W$ predicted more random option-dependent strategy (L_i . $\beta^* = 0.097$, $p = 2.54 \times 10^{-4}$; $P_B - P_W$: $\beta^* = -0.103$, $p = 2.90 \times 10^{-7}$). (**D**) Response variable: entropy of strategy. H(str). This quantity represents the baseline entropy of stay or switch strategy, independent of any information from the previous trial. Longer L_i and smaller $P_B - P_W$ predicted more random strategy (*Li.* $\beta^* = 0111$, $p = 2.91 \times 10^{-5}$; $P_B - P_W$: $\beta^* = -0.093$, $p = 3.28 \times 10^{-6}$). (E). Response variable: mutual information of reward and strategy (MIRS), equal to I(rew; str). Note ERDS = H(str) - MIRS. Larger $P_B - P_W$ predicted more shared information between reward and strategy ($\beta^* = 0.084$, $p = 2.09 \times 10^{-5}$). (**F**). Response variable: mutual information of option and strategy (MIOS), equal to I(opt; str). Note EODS = H(str) - MIOS. No predictors were significant.

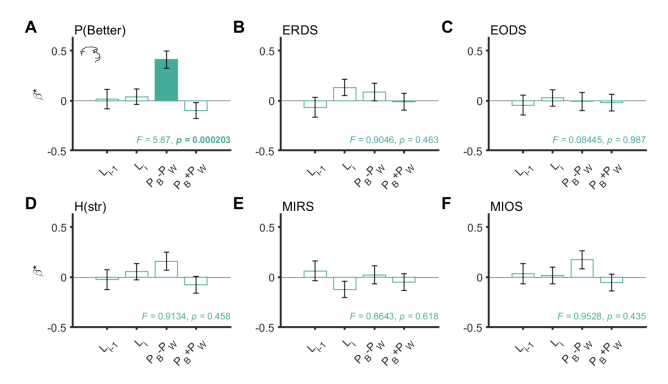


Figure S4. Multiple regression on monkeys' behavioral metrics. Bar plots show the standardized coefficients of regressors, corresponding to task variables representing different types of uncertainty. The bottom of each panel reports the F-statistic and its pvalue of the regression model. Significant predictors are indicated with a filled color bar. L_{i-1} = previous block length. L_i = length of the current block i. $P_B - P_W$ = difference between reward probabilities of better (P_B) and worse (P_W) options (see Methods). $P_B + P_W = \text{sum of }$ the two reward probabilities. All behavioral metrics were computed from the last 20 trials of each block, to avoid confounds with data size and computed entropy values by ensuring the same bin size for every block. (A) Response variable in the regression is performance or probability of choosing the better option during each block, P(Better). Larger $P_B - P_W$ or less uncertainty predicted better performance ($\beta^* = 0.412$, $p = 3.76 \times 10^{-6}$). (B) Response variable: entropy of reward-dependent strategy (ERDS), equal to H(str|rew). Longer L_i was associated with more random reward-dependent strategy but not significantly ($\beta^* = 0.133$, p = .106). (C) Entropy of option-dependent strategy (EODS), equal to H(str|opt). (D) Entropy of strategy, H(str). Larger $P_B - P_W$ or less uncertainty was associated with more random strategy but not significantly ($\beta^* = 0.159$, p = .075). (**E**) Mutual information of reward and strategy (MIRS), equal to I(rew; str). Note ERDS = H(str) - MIRS. Longer L_i was associated with less shared information but not significantly ($\beta^* = -0.122$, p = .140). (F) Mutual information of option and strategy (MIOS), equal to I(opt, str). Note EODS = H(str)- MIOS. Larger $P_B - P_W$ or less uncertainty was associated with more random optiondependent strategy but not significantly ($\beta^* = 0.173$, p = .057).

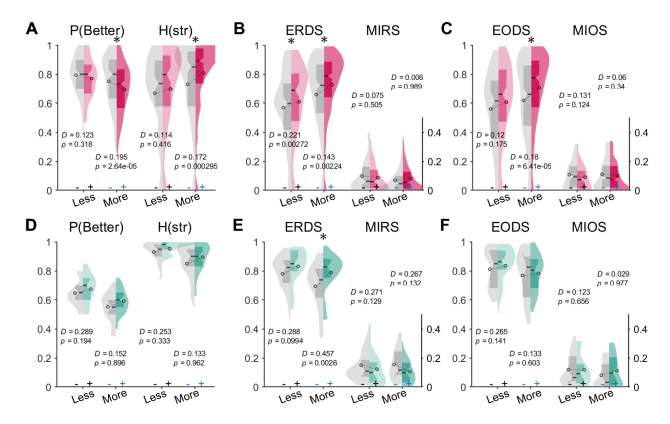


Figure S5. Expectation of block switches depends on the level of uncertainty. Plotted are distributions showing combined effects of uncertainty and block switch expectation on choice behavior for mice (top, A-C) and monkeys (bottom, D-F). For each metric, blocks are divided into four categories: SE and less uncertain (gray, Less-), LE and less uncertain (colored, Less+), SE and more uncertain (gray, More-), and LE and more uncertain (colored, More+). Asterisks show results from a two-sample Kolmogorov-Smirnov test. All tests are two-sided, corrected for multiple comparisons of four. (A) In mice, performance P(Better) was significantly decreased for LE blocks in a more uncertain environment only. Similarly, H(str) or randomness in strategy was significantly increased for LE blocks in a more uncertain environment only. (B) In contrast, ERDS was larger for LE blocks in both less and more uncertain environments. (C) The effect of block switch expectation on EODS was significant in more uncertain environment only. (D-F) For monkeys, the only significant difference between SE and LE was found for ERDS in a more uncertain environment.

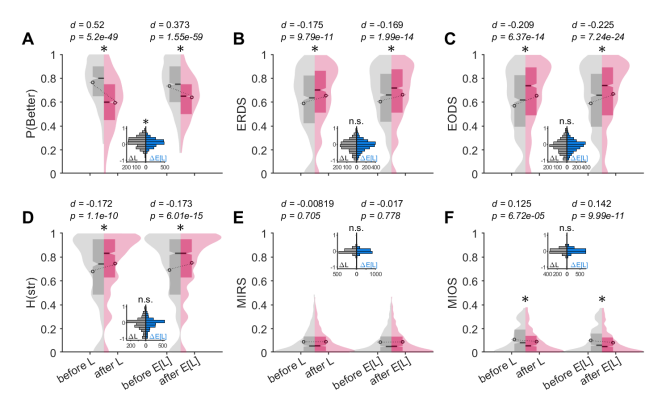


Figure S6. Actual and expected block switches have similar effects on mouse choice behavior. Plotted are distributions of metrics computed from 20 trials around actual block switches (L on the left, containing SE blocks only) and expected block switches (ELI) on the right, containing LE blocks only). Reported values are Cohen's d and p-values based on paired-sample t-test comparing before and after actual and expected block switches (one-sided t-test, with a priori hypothesis that entropy increases and P(Better) (mutual information) decreases after actual and expected block switches). Insets show distributions of paired differences around actual (ΔL) and expected ($\Delta E[L]$) block switches, with asterisks indicating significant difference between two means based on independent-sample t-test (two-sided; n.s. = not significant, p > .05). (A) During 20 trials after the expected block switches point (for LE blocks, right), performance significantly decreased, similar to the behavior after the actual block switches (for SE blocks, left). (B,C) Conditional entropies, ERDS and EODS, similarly increased after both actual and expected block switches. (D) Baseline entropy of strategy, H(str), similarly increased after both actual and expected block switches. (E) MIRS were unaffected by both actual and expected block switches. (F) MIOS significantly decreased after both actual (L, Cohen's d = 0.125, $p = 6.72 \times 10^{-5}$) and expected (E[L], Cohen's d = 0.142, $p = 9.99 \times 10^{-11}$) block switches, indicating a change in the option-dependent strategy.

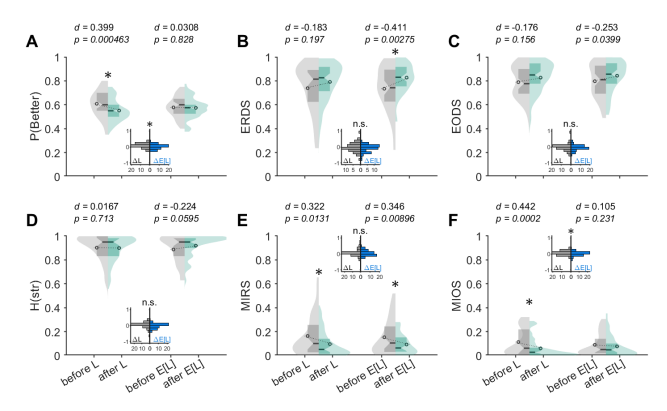


Figure S7. Actual and expected block switches have similar effects on monkey choice behavior. Plotted are distributions of metrics computed from 20 trials around actual block switches (L on the left, containing SE blocks only) and expected block switches (E|L| on right, containing LE blocks only). Reported values are Cohen's d and p-values based on paired-sample t-test comparing before and after actual and expected block switches (one-sided t-test, with a priori hypothesis that entropy increases and P(Better) (mutual information) decreases after actual and expected block switches). Insets show distributions of paired differences around actual (ΔL) and expected ($\Delta E[L]$) block switches, with asterisks indicating significant difference between two means based on independent-sample t-test (two-sided; n.s. = not significant, p > .05). (A) Performance decreased after actual block switches (left), but not significantly after expected block switches (right). (B) ERDS increased after both actual and expected block switches. The increase after actual block switches (left) was moderate but not statistically significant (Cohen's d = -0.183, p = .197). (**C, D**) EODS and H(str) were unaffected by both actual and expected block switches. (E) MIRS significantly decreased after both actual (L, Cohen's d = 0.322, p = .0131) and expected (E[L], Cohen's d = 0.346, p = .00896) block switches, indicating a change in the rewarddependent strategy. (F) MIOS significantly decreased after actual but not expected block switches.

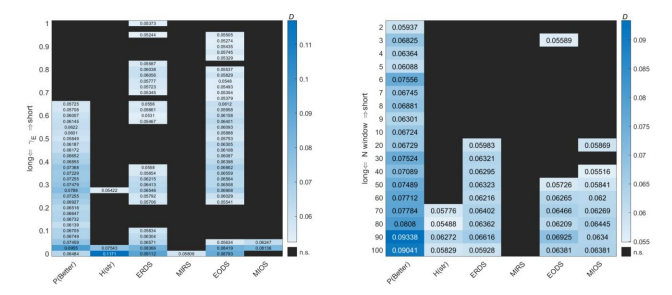


Figure S8. Comparison of different timescales for integration of expectation of block length in mice. Reported are the D statistics of the two-sample Kolmogorov-Smirnov test, with a null hypothesis that the two samples of metrics (from SE and LE blocks) are from the same continuous distribution (two-sided test). n.s. = not significant (p > .025). Left panel: D statistics as a function of γ_E parameter used to compute expected block length E[L] (Eq. 7 in Methods). Larger γ_E values correspond to shorter timescales and smaller values to longer timescales for integration of expected block length. Right panel: D statistics as a function of window size N used to compute expected block length $E[L]_N$ (Eq. 9 in Methods). Smaller N values correspond to shorter timescales, and smaller values to longer timescales for integration of expected block length. For mice, the distinction between SE and LE blocks tended to be more pronounced for longer timescales of E[L].

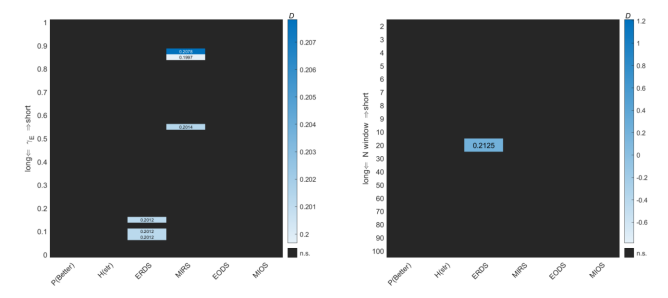


Figure S9. Comparison of different timescales for integration of expectation of block length in monkeys. Reported are the D statistics of the two-sample Kolmogorov-Smirnov test, with a null hypothesis that the two samples of metrics (from SE and LE blocks) are from the same continuous distribution (two-sided test). n.s. = not significant (p > .025). Left panel: D statistics as a function of γ_E used to compute expected block length E[L] (Eq. 7 in Methods). Larger γ_E values correspond to shorter timescales and smaller values to longer timescales for integration of expected block length. Right panel: D statistics as a function of window size N used to compute expected block length $E[L]_N$ (Eq. 9 in Methods). Smaller N values correspond to shorter timescales, and smaller values to longer timescales for integration of expected block length. For monkeys, the distinction between ERDS among SE and LE blocks tended to be significant only for relatively longer timescales of E[L] ($\gamma_E = -0.1$ and N = 20).

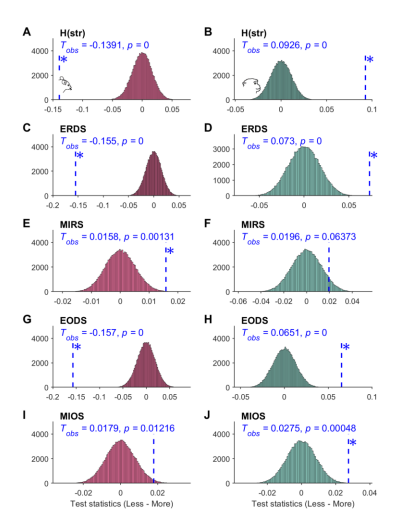


Figure S10. Effects of reward uncertainty on mouse and monkey behavior shown **by permutation test.** Plots show distributions of test statistics from N = 100.000sampled permutations in mice (column 1) and monkeys (column 2), under the null hypothesis that less and more uncertain conditions have the same entropy measures. Vertical dotted lines in blue indicate the actual observed test statistics (T_{obs}), which is the difference between a given entropy metric for the two block types. P-values indicate the probability of observing test statistics as extreme as the observed value (one-tailed). Asterisks indicate p < 0.01. For each permutation, we randomly assign blocks into two groups with sample size n_1 or n_2 , corresponding to the total number of blocks in less or more uncertain conditions, respectively. A single entropy measure is computed for each type of blocks by concatenating all trial vectors from the sampled permutation. Test statistic for each sample is then computed by taking the difference in metrics between the first and the second group. (A,C,E,G,I) Mouse data in red. More uncertain blocks had significantly larger H(str), ERDS, and EODS, indicating an overall increase in randomness of choice strategy. MIRS was also significantly smaller, indicating a decreased link between reward outcome and subsequent strategy. (B,D,F,H,J) Monkey data in teal. More uncertain blocks had significantly smaller H(Str), ERDS, EODS, indicating a more consistent strategy. MIOS was also significantly smaller, indicating a decreased link between previous choice option and strategy.

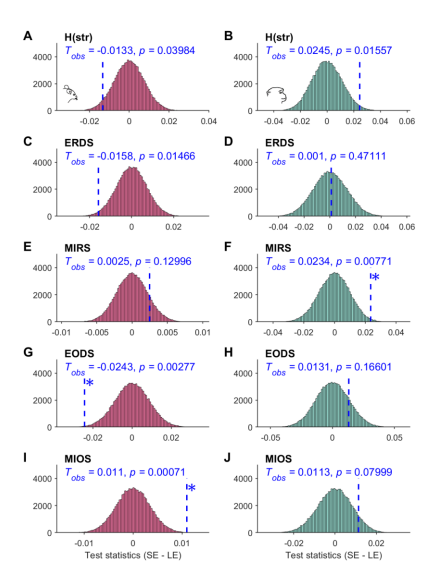


Figure S11. Effects of block switch expectation on mouse and monkey behavior **shown by permutation test**. Plots show distributions of test statistics from N = 100,000sampled permutations in mice (column 1) and monkeys (column 2), under the null hypothesis that SE and LE blocks have the same entropy measures. Vertical dotted lines in blue indicate the actual observed test statistics (Tobs), which is the difference between a given entropy metric for the two block types. P-values indicate the probability of observing test statistics as extreme as the observed value (one-tailed). Asterisks indicate p < 0.01. For each permutation, we randomly assign blocks into two groups with sample size n_1 or n_2 , corresponding to the total number of blocks in SE or LE blocks, respectively. A single entropy measure is computed for each type of blocks by concatenating all trial vectors from the sampled permutation. Test statistic for each sample is then computed by taking the difference in metrics between the first and the second group. (A,C,E,G,I) Mouse data in red. LE blocks had significantly larger EODS $(T_{obs} = -0.0243, p = .00241)$ and smaller MIOS $(T_{obs} = 0.011, p = .00061)$, indicating a change in option-dependent strategy. (B,D,F,H,J) Monkey data in teal. LE blocks had significantly smaller MIRS ($T_{obs} = 0.0131$, p = .00762), indicating a change in rewarddependent strategy.